NOVEL ANALYTICAL STRATEGIES FOR TRACING THE ORGANIC **CARBON CYCLE IN MARINE AND RIVERINE PARTICLES**

By

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S.B., Brown University, 2011

Submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy at the MASSACHUSETTS INSTITUTE OF TECHNOLOGY and the

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Novel analytical strategies for tracing the organic carbon cycle in marine and riverine particles

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Sarah Zhou Rosengard

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Abstract

Particulate organic carbon (POC) in the ocean and mobilized by rivers on land transfers $\sim 0.1\%$ of global primary productivity to the deep ocean sediments. This small fraction regulates the long-term carbon cycle by removing carbon dioxide from the atmosphere for centuries to millennia. This thesis investigates mechanisms of POC transfer to the deep ocean by analyzing particles collected in transit through two globally significant carbon reservoirs: the Southern Ocean and the Amazon River Basin. These endeavors test the hypothesis that organic matter composition controls the recycling and transfer efficiency of POC to the deep ocean, and illustrate new applications for ramped pyrolysis/oxidation (RPO), a growing method of POC characterization by thermal stability. By coupling RPO to stable and radiocarbon isotope analyses of riverine POC, I quantify three thermally distinct soil organic carbon pools mobilized by the Amazon River, and evaluate the degradability and fate of these different pools during transport to the coastal Atlantic Ocean. More directly, RPO analyses of marine samples suggest that POC transfer in the water column is in fact selective. Observations of consistent biomolecular changes that accompany transport of phytoplankton-derived organic matter to depth across the Southern Ocean support the argument for preferential degradation of specific POC pools in the water column. Combining discussions of POC recycling and transfer across both marine and terrestrial systems offer new perspectives of thermal stability as a proxy for diagenetic stability and POC degradation state. The challenges of interpreting RPO data in these two environments set the stage for applying the technique to more controlled experiments that trace POC from source to long-term sink.

Thesis Co-advisors: Valier Galy, Associate Scientist, WHOI Phoebe Lam, Assistant Professor, University of California- Santa Cruz This thesis is dedicated to four important women: Marjorie Rosengard, Barbara Rosengard, my grandmother and Miah. Thank you for your unconditional love and nurturing.

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Chapter 1: Introduction

1 The modern carbon cycle

Fossil fuel emissions in the industrial era have cumulatively released ~545 gigatons (Gt) of carbon dioxide (CO₂), a prominent greenhouse gas, to the atmosphere, resulting in the highest atmospheric CO₂ concentrations in human history. Within this relatively short period, the increase in atmospheric CO₂ concentrations has been largely moderated by carbon fixation into organic matter by land plants and marine phytoplankton, and dissolution of CO₂ into the surface ocean (Sabine et al., 2004). Because these terrestrial and marine reservoirs turn over carbon on relatively short, decadal time scales, their storage size and the rates at which they absorb atmospheric CO₂ have fluctuated throughout the 20th century in response to rising anthropogenic carbon emissions, heat accumulation in the atmosphere and oceans and, in the case of the terrestrial vegetation, land use change (Houghton et al., 1999; Körner, 2003; Behrenfeld et al., 2006).

This research compilation focuses primarily on the small fraction (<1%) of terrestrial and surface ocean primary productivity that is further exported into the deep ocean (Volk and Hoffert, 1985; Martin et al., 1987), where residence times for carbon extend to millennia. Marine particles produced in the surface ocean and terrestrial particles mobilized through rivers are the primary vehicles for export to the deep ocean. Transport of marine particulate organic carbon (POC), or the biological pump, is driven by sinking particles from the euphotic zone to the deep ocean (>2000 m depths) (Riley et al., 2012). The terrestrial analogue to the marine biological pump is active transport of primarily vegetation and soil-derived organic matter (Meybeck, 1982; Hedges et al., 1997) in riverine particles to the coastal ocean and continental shelf. The physical journey of riverine POC depends on the downstream velocity within the river channel and the inland landscape (Burd et al., 2016).

The fraction of marine primary productivity exported out of the surface ocean and terrestrial primary productivity exported from land into rivers equates to ~16 Gt/yr of marine POC (Falkowski et al., 1998) and 0.2 Gt/yr of terrestrial POC (Galy et al., 2015),

which in sum are comparable to the current rate of annual carbon emissions. An even smaller fraction of total primary productivity, <0.5% (Martin et al., 1987; Burdige, 2007), equivalent to 0.1-0.2 Gt/yr, reaches the seafloor sediment surface . These fluxes into the deep ocean are small, owing to rapid heterotrophic remineralization and photo-oxidation in the water column and within rivers (Martin et al., 1987; Amon and Benner, 1996; Benner and Biddanda, 1998; Mayorga et al., 2005; Buesseler and Boyd, 2009; Ward et al., 2013; Giering et al., 2014), but they remove carbon from the atmosphere for centuries to millennia. Over these long time scales, this deep reservoir amasses several orders of magnitude more carbon than do the terrestrial and marine primary producer reservoirs. The storage capacity of the deep ocean reservoir has likely remained unperturbed throughout the industrial era (Sarmiento and Le Quéré, 1996; Sarmiento et al., 1998), and is expected to buffer long-term anthropogenic CO_2 accumulation in the atmosphere (Kwon et al., 2009).

2 Research objectives

Particle transport studies have reported significant geographical and temporal variations in the fraction of marine and terrestrial primary productivity exported (export efficiency) and sequestered (transfer efficiency) in the deep ocean (Hedges and Oades, 1997; Henson et al., 2011; Moreira - Turcq et al., 2013). The following chapters of this dissertation aim to expand mechanistic understanding of this variation in POC export and transfer efficiency by (1) comparing multiple factors that govern the fraction of primary productivity transferred to the deep ocean and (2) assessing different empirical approaches to sampling and analyzing the dynamics of POC exported from the water column and through rivers. The sample data span two globally significant carbon sinks for atmospheric CO₂: the Southern Ocean, which supports 1-4 Gt/year marine primary productivity (Arrigo et al., 1998; Priddle et al., 1998), and the Amazon River Basin, which harbors the largest tropical rain forest and river on the planet in terms of discharge and watershed area. Both regions are likely to shift disproportionately in response to climate change (Gille, 2002; Laufkötter et al., 2015) and landscape transformation across the Amazon River Basin (Davidson et al., 2012; Brienen et al., 2015).

3 Mechanisms of POC export and transfer efficiency

The fraction of both marine and terrestrial primary productivity sequestered in the deep ocean is a balance between two competing but not mutually exclusive forces: the physical processes that drive particle movement, and the geochemical processes that influence rates of respiration. Physical processes include particle size and density (Rouse, 1950; Wilson et al., 2008; Burd and Jackson, 2009; Bouchez et al., 2011; Lupker et al., 2011), mineral-POC associations (Armstrong et al., 2002; Francois et al., 2002; Klaas and Archer, 2002; Mayer et al., 2004; Bouchez et al., 2014), and advection and mixing of water parcels (Dall'Olmo and Mork, 2014). Processes that control remineralization rates include the composition of phytoplankton, zooplankton and particle-associated bacterial communities (Battin et al., 2008; Guidi et al., 2009; Buesseler and Boyd, 2009; Satinsky et al., 2014), and relatedly, the composition and quality of POC exported by different phytoplankton communities (Francois et al., 2002). Abiotic remineralization processes like photo-oxidation are not addressed in this thesis.

Chapter 2 takes advantage of the variable diatom and coccolithophore communities across the Southern Ocean to compare how biomineral ballast mechanisms influence POC transfer efficiency (Armstrong et al., 2002; Francois et al., 2002; Klaas and Archer, 2002). Comparisons among POC, biogenic silica and calcite fluxes in particles through the upper 1000 meters of the water column did not provide consistent evidence for mineral ballasting with depth, leading instead to the hypothesis that phytoplankton community composition and POC quality influence POC export more than mineral ballasting by controlling POC remineralization rates in the water column (Francois et al., 2002; Rosengard et al., 2015).

4 Evaluating POC quality across marine and terrestrial sources

The hypothesis posited at the end of chapter 2 directed me to seek more direct evidence linking POC transfer efficiency to POC quality and primary producer community composition in chapters 3, 4 and 5. The argument that organic matter "quality" controls its degradability by bacteria in the environment and therefore POC transfer efficiency is based on the perspective that POC degradation is selective. Observations that specific molecular classes of organic matter accumulate in the deeper ocean, both terrestrial and marine-derived (Wakeham et al., 1997; Elias et al., 2000; Baldock et al., 2004; Burd et al., 2016), imply that certain pools of organic matter are more diagenetically stable than others. If quantitatively significant, selective remineralization of less stable pools by heterotrophs during POC transport would impact the export and transfer efficiencies of primary productivity into the deep ocean.

Our ability to address the POC quality hypothesis depends on our ability to describe POC composition and the degradability of distinct POC pools. Organic geochemists employ several techniques to probe organic matter quality, from bulk compositional metrics that integrate across the diverse molecular matrix of POC (e.g., Trull and Armand, 2001), to analyses of specific biomolecular tracers (e.g., Elias et al., 2000). Chapters 3-5 utilize a range of these strategies, and further apply a relatively novel technique for assessing organic matter composition by its thermal stability: ramped pyrolysis/oxidation (RPO) (Fig. 1). This method oxidizes or pyrolyzes a sample to CO_2 throughout a constant temperature ramp, generating plots of CO₂ release against temperature, referred to as thermograms, that can be interpreted as the activation energy distribution of POC (Cramer, 2004). Previous applications of RPO have suggested that the activation energy of POC decomposition during ramped oxidation could serve as a proxy for the stability of POC in the environment (Rosenheim et al., 2008; Rosenheim and Galy, 2012; Rosenheim et al., 2013). Chapter 3 uses a controlled sampling experiment in Woods Hole to demonstrate that different sample drying temperatures do not compromise the thermal stability of POC, implying that RPO analyses are appropriate for POC sampled, dried and stored across a range of temperatures.

Collectively, chapters 4 and 5 explore different applications of RPO to probe the relationship between POC composition and export/transfer efficiency. Chapter 4 directly follows up on chapter 2 by comparing the thermal stability and quality of POC produced and exported from different phytoplankton communities across the Southern Ocean. The results offer a unique perspective of compositional transformations in POC that accompany degradation of marine POC. Chapter 5 further demonstrates that RPO is

useful for distinguishing and quantifying input of compositionally distinct sources of terrestrial POC to the Amazon River. Because the river basin drains diverse vegetative landscapes and soil pools spanning different degradation histories, the data provide insights towards the diagenetic stability and proportions of these different sources in riverine POC exported to the coastal Atlantic Ocean.

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Figure 1. Schematic of the ramped pyrolysis/oxidation (RPO) unit at the National Ocean Sciences Accelerator Mass Spectrometery facility at WHOI. Adapted from (Fornace, 2016).

Chapter 2:

Carbon export and transfer to depth across the Southern Ocean Great Calcite Belt

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Abstract

Sequestration of carbon by the marine biological pump depends on the processes that alter, remineralize and preserve particulate organic carbon (POC) during transit to the deep ocean. Here, we present data collected from the Great Calcite Belt, a calcite-rich band across the Southern Ocean surface, to compare the transformation of POC in the euphotic and mesopelagic zones of the water column. The ²³⁴Th-derived export fluxes and size-fractionated concentrations of POC, particulate inorganic carbon (PIC), and biogenic silica (BSi) were measured from the upper 1000 m of 27 stations across the Atlantic and Indian sectors of the Great Calcite Belt. POC export out of the euphotic zone was correlated with BSi export. PIC export was not, but did correlate positively with POC flux transfer efficiency. Moreover, regions of high BSi concentrations, which corresponded to regions with proportionally larger particles, exhibited higher attenuation of $>51 \mu m$ POC concentrations in the mesopelagic zone. The interplay among POC size partitioning, mineral composition and POC attenuation suggests a more fundamental driver of POC transfer through both depth regimes in the Great Calcite Belt. In particular, we argue that diatom-rich communities produce large and labile POC aggregates, which generate high export fluxes but also drive more remineralization in the mesopelagic zone. We observe the opposite in communities with smaller calcifying phytoplankton, such as coccolithophores. We hypothesize that these differences are influenced by inherent differences in the lability of POC exported by different phytoplankton communities.

1 Introduction

The biological pump sequesters atmospheric carbon dioxide (CO₂) in the ocean (Volk and Hoffert, 1985) by way of phytoplankton-driven CO₂ fixation, followed by the sinking of this fixed particulate organic carbon (POC) as aggregates and fecal pellets down the water column (Riley et al., 2012). The quantity per unit area and time of POC exiting the base of the euphotic zone defines the export flux, while export efficiency represents the fraction of bulk primary production comprising this flux (Buesseler, 1998). In the mesopelagic zone (from the base of the euphotic zone to ~1000 m), export flux attenuates due to remineralization mediated by zooplankton grazing and bacteria

(Buesseler and Boyd, 2009; Giering et al., 2014; Martin et al., 1987). The flux of this processed organic carbon leaving the mesopelagic zone, only $\leq 10\%$ of export flux, directly scales with the quantity of atmospheric CO₂ sequestered by the marine biological pump over hundreds to thousands of years (Kwon et al., 2009).

On average, only ~1% of the organic matter produced by phytoplankton in the surface reaches the deep sea (Martin et al., 1987). However, export and sequestration flux vary widely by region, as do export efficiencies and attenuation of export flux (Martin et al., 1987; Buesseler et al., 2007; Thomalla et al., 2008; Buesseler and Boyd, 2009; Henson et al., 2012b; Henson et al., 2011). Such variations may drive observed differences in the weight percent of organic carbon deposited at the sediment surface (Hedges and Oades, 1997), suggesting that the overall strength of the biological pump as a carbon sink is not globally uniform. These geographical differences have spurred decades of research into how mechanisms in the shallower ocean – the euphotic and mesopelagic zones – alter sinking particulate organic matter during vertical transit.

As an example, Armstrong et al. (2002), Klaas and Archer (2002) and (Francois et al. (2002) posited that mineral associations with sinking organic carbon could explain these variations. Their ballast hypothesis model suggested that minerals enhanced the biological pump (1) by increasing the density, and consequently, the sinking speed of particulate organic matter and (2) by inhibiting organic carbon remineralization down the water column. Expediting vertical transit decreases the time for remineralization to act on sinking particulate organic matter, increasing its chances of reaching the deep sea. The authors observed that calcite flux in the bathypelagic zone (>1000 m) explains roughly half of the variation in the magnitude of POC flux reaching the deep sea (Klaas and Archer, 2002), and may also account for some of the observed geographical variation in POC flux attenuation with depth (Francois et al., 2002).

In its conception and infancy, the ballast hypothesis was based upon observed correlations between mineral and organic carbon fluxes in the deep (>1000 m) sea. Yet, evidence for the ballast mechanism in the euphotic and mesopelagic zones remains equivocal, as deeper correlations are scarcely matched by shallower ocean observations (Le Moigne et al., 2012). Several surface regions do not exhibit ballast correlations between mineral flux and POC flux (e.g., Thomalla et al., 2008; Henson et al., 2012b). In

the Atlantic and Southern Oceans, (Le Moigne et al., 2012) found a significant fraction of POC export flux to remain unassociated with minerals altogether. Moreover, tank incubations simulating POC and mineral suspensions yield conflicting results: some have observed mineral associations to increase aggregate sinking rates (Engel et al., 2009), while others find no such effect (Passow and De la Rocha, 2006). De La Rocha et al. (2008) even suggest that sticky polymers from POC might ballast sinking minerals, rather than vice-versa.

The scarcity of evidence supporting a shallow ocean ballast mechanism suggests that the transit of particulate organic carbon in the surface, mesopelagic and deeper ocean is mechanistically de-coupled (Lomas et al., 2010; Lam et al., 2011). Indeed, the debate surrounding the ballast hypothesis arises from a deeper issue of whether the mechanisms that influence carbon export from the euphotic zone are the same as those that control its remineralization in the mesopelagic zone, and/or its transfer beyond the mesopelagic zone into the deep sea.

The following report compares the export of organic carbon from the euphotic zone with its transfer through the mesopelagic zone across the region of the Great Calcite Belt (Balch et al., 2011a; Balch et al., 2014; Fig. 1). Spanning across the Southern Ocean, particularly between the Subtropical and Polar Fronts, the Great Calcite Belt defines a highly reflective band observed from space during each austral spring and summer. Its high reflectivity is caused by calcite-rich surface waters produced by coccolithophore blooms in the Southern Ocean. In this zone, coccolithophores are more abundant than in regions north and south of the Belt. South of the Polar Front, coccolithophore abundances decline dramatically as dissolved silica concentrations increase and diatoms flourish (Balch et al., 2011a).

Spanning a large range in surface mineral concentrations, primary productivity, and phytoplankton community composition (Balch et al., 2011a), the Great Calcite Belt provides an excellent opportunity to assess the processes controlling organic carbon export, export efficiency, and attenuation of POC concentration ([POC]) with depth. Here, we report estimates of ²³⁴Th-derived POC fluxes and [POC] through both the euphotic and mesopelagic zones within the Atlantic and Indian sectors of the Great Calcite Belt. We focus on the upper 1000 m of the Great Calcite Belt because the

attenuation of POC flux and concentration is most dramatic within this depth interval (Martin et al., 1987; Lam et al., 2011). As the following discussion illuminates, this study additionally weighs the ballast hypothesis against other mechanisms hypothesized to control the transfer of organic carbon through the water column, and ultimately into the deep sea, where carbon residence time modulates atmospheric pCO_2 and climate over hundreds to thousands of years (Kwon et al., 2009).

2 Methods

2.1 Field site

Samples from the Great Calcite Belt were collected during two research cruises, GB1 and GB2, which transited the Atlantic and Indian sectors of the Great Calcite Belt during the austral summer of 2011 and 2012, respectively (Fig. 1), concurrent with the putative coccolithophore bloom (Balch et al., 2011a). In 2011, for cruise GB1 (MV1101), the *R/V Melville* crossed the Atlantic sector from Punta Arenas, Chile to Cape Town, South Africa, sampling between 39°S and 59°S. One year later, for cruise GB2 (RR1202), the *R/V Revelle* crossed the Indian sector from Durban, South Africa to Perth, Australia, sampling between 37°S and 60°S (Table 1). Both cruise tracks crossed the Subtropical, Subantarctic and Polar fronts, which are approximately located at 40°S, 45°S and 52°S (e.g., Belkin and Gordon, 1996; Sokolov and Rintoul, 2009), respectively, defining observed shifts in temperature and nutrient characteristics of the surface ocean.

Each day during GB1 and GB2, 30-L Niskin samples were collected pre-dawn for measuring primary production. A Biospherical Instruments (San Diego, CA) sensor was mounted on the CTD/rosette and referenced to a deck sensor mounted on the ship's superstructure to measure Photosynthetically Available Radiation (PAR) during the casts. Water was then sampled at fixed light depths relative to surface irradiance to match light levels in deck-board incubators: 36.5%, 21.1%, 11.7%, 3.55%, 1.93% and 0.28%. The light depths were calculated two ways: (a) between 10:00 and 14:00 h local time (during daylight hours), percentages of surface irradiance were derived directly from the downcast PAR profile immediately preceding bottle firing, or (b) at all other times, the

light levels were back-calculated from the previously-determined relationship between beam transmittance and diffuse attenuation of PAR (Balch et al., 2011b). From these casts, primary production rates were measured using the ¹⁴C microdiffusion technique (Paasche and Brubak, 1994) with modifications by Balch et al. (2000) (see also Fabry and Balch, 2010).

2.2 Size-fractionated particle collection

We report measurements of total and particulate ²³⁴Th activity and sizefractionated particle composition from 27 stations (Fig. 1; Table 1).

Size-fractionated particles were collected at eight depths in the upper 1000 m of fourteen stations from GB1 and thirteen stations from GB2, using modified battery operated in-situ pumps (McLane WTS-LV). The modified pumps directed seawater through two flow paths (Lam et al., 2015), each of which passed through a "mini-MULVFS" filter holder designed to retain large particles (Bishop et al., 2012). Seawater first passed through 51 µm polyester pre-filters in both filter holders for collection of large (>51 µm) size-fraction particles, and then through paired 0.8 µm polyethersulfone (SuporTM) filters in one flow path and paired 1 um quartz fiber (WhatmanTM OMA) filters in the other flow path, both of which collected small (<51 µm) size-fraction particles. An average of 200 L and 500 L of seawater passed through the Supor and OMA flow paths over 1-2.5 hours, respectively. Immediately after collection, half to all of the >51 µm size-fraction particles from one flow path were rinsed off of the polyester prefilters and onto 25 mm 1 µm Sterlitech silver filters using 0.2 µm-filtered seawater, and dried at 50°C for subsequent analysis of particulate ²³⁴Th, particulate organic carbon (POC), and particulate inorganic carbon (PIC, or calcium carbonate). Subsamples of QMA filters were likewise dried at 50°C for 234 Th and POC analysis in the <51 μ m sizefraction. Finally, the polyester pre-filters from the other flow path and Supor filters were dried in a laminar flow hood at room temperature.

In the euphotic zone, where most POC is produced, these operationally defined size fractions allude primarily to the structure of phytoplankton communities producing POC (e.g., large diatoms would be found in $>51 \,\mu m$ size-fraction particles). In the

mesopelagic zone, which extends from the base of the euphotic zone to 1000 m in depth, $>51 \mu m$ POC is predominantly comprised of phytoplankton and bacterial biomass that has been repackaged into aggregates and fecal pellets. The $>51 \mu m$ particles collected at station GB1- 85 illustrate these different size-fraction interpretations by depth. Shallower particles collected at 25 m and 73 m, the base of the euphotic zone, are mainly comprised of intact phytoplankton cells (Figs. 2a, 2b). By contrast, deeper particles collected at 173 m exhibit the features of particulate aggregates and fecal pellets (Fig. 2c).

2.3 Particle composition

Bulk concentrations of POC, PIC, biogenic silica (BSi), and particulate ²³⁴Th activity were measured in both <51 μ m and >51 μ m fractions of particles collected at each station. POC concentrations were measured at all depths of the profiles, while [PIC] and [BSi] were mainly measured at select depths above 200 m and at the deepest depth (800-1000 m) of the profile. Particulate ²³⁴Th activities in all sub-fractions of >51 μ m (25 mm silver filters) and <51 μ m (25 mm QMA filters) samples were measured using low level Risø beta counters immediately on the ship and in the lab at least six ²³⁴Th half-lives post-collection for background activity.

After counting for ²³⁴Th background activity, ~25% of the silver filter (~ 115 L equivalent) was fumed overnight (12-17 hours) with concentrated hydrochloric acid to remove inorganic carbon, before measuring >51 μ m [POC] using an elemental CHN analyzer. A similar protocol was followed to measure <51 μ m [POC] from one 12 mm-diameter subsample of each QMA filter, representing ~1% of the entire sample (~5 L equivalent). Vertical profiles of >51 μ m and <51 μ m [POC] between the base of the euphotic zone and the deepest measurement at 800 – 1000 m were fitted to a power-law function to describe the attenuation of [POC] with depth, based on a function first applied to POC flux by Martin et al. (1987) and then analogously to POC concentration by Lam and Bishop (2007),

$$[POC]_z = [POC]_0 \left(\frac{z}{z_{PAR}}\right)^{-b}$$
(1)

where, at most stations, z_{PAR} represents the depth of 0.3% photosynthetically available radiation (see Section 2.4). The exponent b represents the attenuation coefficient, with

higher attenuation coefficients (more negative exponents) for profiles with greater attenuation of >51 μ m [POC] with depth. We focus our discussion on the attenuation of >51 μ m [POC], because we assume that they contribute disproportionately to sinking fluxes compared to the <51 μ m size fraction (McCave, 1975; Lam and Bishop, 2007; Lam et al., 2011). Figure 3 and Table 2 show all significant (p<0.05) power law fits for >51 μ m [POC] profiles.

PIC in the samples was assumed to be biomineral calcium carbonate (CaCO₃), and was derived from particulate calcium (Ca) corrected for salt Ca using a seawater 0.0382 Ca:Na (g:g) ratio (Lam and Bishop, 2007; Pilson, 2012). In the in-situ pump samples, salt-derived Ca typically accounted for ~60% of total Ca. The >51 µm PIC sizefraction concentrations were measured mainly in subsamples of remaining pre-filter material and occasionally in sub-fractions of the silver filters, if the former were unavailable. The <51 µm size fraction [PIC] was measured in three 12mm circular QMA subsamples, representing ~15 L or ~3% of the sample. Subsamples were leached in 0.6 N ultrapure Sea-StarTM Baseline hydrochloric acid (HCl) at 60°C for 12-16 hours. The leachate was subsequently filtered through a 0.4 µm polycarbonate membrane filter, diluted to 0.12 N HCl, and spiked with 1 ppb of Indium as an internal standard. The spiked leachate solution was then analyzed for Ca, Na and P using an Element 2 sectorfield inductively-coupled plasma mass spectrometer (ICP-MS) in medium and high resolution. Counts per second were converted to concentration using external mixed element standard curves.

For measuring >51 μ m and <51 μ m [BSi], prefilter or Supor subsamples, respectively, were leached in 0.2 N sodium hydroxide at 85°C for one hour, and analyzed by standard spectrophotometric detection of the blue silico-molybdate complex in each leachate within 24 hours of the leach (Strickland and Parsons, 1968; Brzezinski and Nelson, 1989). Absorbance through each sample was converted to concentration using an external Si standard curve.

2.4 ²³⁴Th-derived flux estimates

Particle fluxes were estimated at each station by measuring the water-column disequilibrium between ²³⁴Th and ²³⁸U in the upper 350 m of the water-column (Savoye et al., 2006). ²³⁴Th is the radioactive daughter of ²³⁸U with a short enough half-life (24.1 days) relative to ²³⁸U such that it is assumed to be in secular equilibrium with its parent isotope in the absence of particle scavenging (i.e., ²³⁴Th activity = ²³⁸U activity). Disequilibria between the two isotope activities in the water column are attributed to the scavenging of ²³⁴Th by sinking particles (Savoye et al., 2006). Integrating the deficit in ²³⁴Th relative to ²³⁸U provides a measure of particle flux down the water column (Buesseler et al., 2006). Because of the short half-life of ²³⁴Th, deviation from secular equilibrium exists only in regions of high particle flux. Thus, ²³⁴Th-based flux estimates are most frequently applied in the euphotic zone of the ocean where particle export is maximal.

 234 Th- 238 U deficits were determined by measuring total water-column activities of both isotopes. 238 U activity (A_{U-238}) profiles were calculated from salinity by the following relationship (Owens et al., 2011):

 $A_{\rm II-238}$ (dpm L⁻¹) = (0.0786 × Salinity) – 0.315 (2)Total water-column²³⁴Th activity (A_{Th-234}) profiles were determined from 4 L seawater samples collected by CTD casts down to 300-350 m at each station (Pike et al., 2005). Shortly after collection, each 4 L seawater sample was acidified to pH 2 using concentrated nitric acid (HNO₃), spiked with 1 g of 230 Th of a known activity (50.06 dpm g^{-1}) as a yield monitor, equilibrated for 8 hours, and finally brought up to pH 8.5 using ammonium hydroxide (NH₄OH) (van der Loeff et al., 2006). Manganese chloride $(MnCl_2)$ and potassium permanganate (KMnO₄) were added to the neutralized seawater to form a manganese oxide (MnO₂) precipitate, which efficiently scavenges both natural ²³⁴Th and added ²³⁰Th. After 12 hours, the precipitate was filtered onto a quartz fiber filter, dried at 50°C, and then mounted beneath a sheet of Mylar and aluminum foil. ²³⁴Th activity in the precipitate was measured on board by low level Risø beta counters and post-cruise after at least six ²³⁴Th half-lives for background activity. The ²³⁰Th spike was recovered by fully dissolving the MnO₂ precipitate, adding a 1 g spike of ²²⁹Th of a known activity (69.74 dpm g⁻¹), and measuring ²²⁹Th:²³⁰Th ratios on an Element 2 sectorfield ICP-MS in low resolution. Recovery of ²³⁰Th spike was derived from this ratio, and

used to correct for inefficiencies in the scavenging of total seawater ²³⁴Th by MnO₂ precipitation.

To calibrate beta counting efficiency for each cruise, total deep water (i.e., below 2000 m)²³⁴Th activities were compared to total deep water ²³⁸U activities, as measured in 4-5 replicate samples from 2-3 deep water CTD casts during each cruise (at 5000 m during GB1, and at 2500 m during GB2). Beta counting efficiencies were adjusted such that ²³⁴Th and ²³⁸U activities were equal in these deep measurements, as secular equilibrium would be expected at such depths. We only report upper water-column activities (<350 m) after correcting for experimental efficiencies in both the seawater collection process and beta detector counting. Uncertainties in the total ²³⁴Th activity profiles averaged 4.5% and were propagated from errors associated with counting statistics, recoveries, and beta-counting efficiency.

To calculate ²³⁴Th export flux, ²³⁴Th activity deficits were integrated down to the base of the euphotic zone (z_{PAR}) (Buesseler et al., 2008; Thomalla et al., 2008):

²³⁴Th Flux (dpm m⁻²d⁻¹) =
$$\int_0^{z_{PAR}} (A_{U-238} - A_{Th-234}) dz$$
 (3)

At most stations, the export depth, z_{PAR} , was chosen to be the depth where light levels were 0.3% of surface-level PAR. The exception was station GB2-27, which did not include a PAR measurement profile. For this station, the z_{PAR} value of 105 m was defined as the depth where the transmissometry-based particle concentration decreased. These export depths were compared to one additional metric describing particle concentration in seawater: the depths where ²³⁴Th and ²³⁸U activities re-established secular equilibrium, or $z_{Th/U}$. We explore the sensitivity of ²³⁴Th flux estimates to choice of z_{PAR} in Sects. 3 and 4.1.

 234 Th flux estimates were converted to POC, PIC and BSi fluxes by multiplication with ratios of >51 µm POC, PIC, and BSi concentrations to particulate 234 Th activity in samples at z_{PAR} (Thomalla et al., 2008; Sanders et al., 2010):

POC Flux (μ mol m⁻²d⁻¹) = [POC]: $A_{Th-234} \times {}^{234}$ Th Flux (dpm m⁻²d⁻¹) (4)

PIC Flux (
$$\mu$$
mol m⁻²d⁻¹) = [PIC]: $A_{Th-234} \times {}^{234}$ Th Flux (dpm m⁻²d⁻¹) (5)

Si Flux (μ mol m⁻²d⁻¹) = [BSi]: $A_{Th-234} \times {}^{234}$ Th Flux (dpm m⁻²d⁻¹) (6)

2.5 Interpolation of data

In all cases where ²³⁴Th activity, >51 μ m and <51 μ m [POC] and mineral concentrations, and >51 μ m particulate ²³⁴Th measurements were unavailable at z_{PAR}, linear interpolations between the sampling depths above and below z_{PAR} were used to estimate a value at the export depth (Table 1). The >51 μ m and <51 μ m size-fraction POC concentrations were interpolated by the power law attenuation function when fits were significant (p<0.05), or linearly when these power-law fits were not significant or inconsistent with the broader shape of the [POC] profile at that particular station. In general, corresponding POC:²³⁴Th, BSi:²³⁴Th, and PIC:²³⁴Th ratios are quotients of these interpolated values except as noted in Tables 2 and 3.

3 Results

²³⁴Th activity profiles were measured over the upper 300 – 350 m at the 27 stations of cruises GB1 and GB2 (Fig. 4; Appendix A, Table S1). Each activity profile is associated with two metrics that have been used in previous studies to define the export depth (see Section 2.4): the base of the euphotic zone (z_{PAR}), which we define at 0.3% surface photosynthetically available radiation (PAR) (e.g., Buesseler and Boyd (2009), and $z_{Th/U}$, where ²³⁴Th and ²³⁸U activities re-establish secular equilibrium (Table 1). In most stations, profiles exhibited ²³⁴Th activity deficits over a range from surface to 75 – 170 m in depth, below which ²³⁴Th activity generally returned to secular equilibrium with ²³⁸U activity, within error. The notable exceptions were profiles at stations GB1-6, and GB1-16, which did not return to secular equilibrium by 170 m in depth. Considering that stations GB1-6 and GB1-16 are closest to shore, their sustained ²³⁴Th deficits may have been influenced by lateral advection of particles from the continental shelf. At these stations, $z_{Th/U}$ depths were approximated by the depth below which ²³⁴Th activities remain constant with depth. For example, at station GB1-6, $z_{Th/U} = 130$ m because below this depth ²³⁴Th activities remained relatively constant.

In the Atlantic sector, sampled in January – February 2011, all observed z_{PAR} depths were significantly shallower than $z_{Th/U}$ depths (Student's t-test p<0.05); on average, z_{PAR} was 66 ± 44 % shallower than $z_{Th/U}$. By contrast, in the Indian sector,

sampled roughly a year later in February – March 2012, z_{PAR} was not significantly different from $z_{Th/U}$ (p>0.05), and the average relative difference was -6 ± 29%. In general, when water-column ²³⁴Th activity is at steady-state, the euphotic zone should correspond to the region of ²³⁴Th deficit relative to ²³⁸U (Buesseler et al., 2008; Buesseler and Boyd, 2009), i.e., z_{PAR} should equal $z_{Th/U}$.

Using integrated activity deficits, export fluxes of ²³⁴Th, POC, PIC, and BSi at z_{PAR} were estimated at the 27 sites (Figs. 5, 6; Table 3). Overall mean ²³⁴Th fluxes at z_{PAR} were 1,413 ± 432 dpm m⁻² d⁻¹ (mean ± 1 s.d.), and ranged from 717 to 2,437 dpm m⁻² d⁻¹ at stations GB2-112 and GB1-6, respectively. Mean derived POC fluxes at z_{PAR} were 4.5 ± 3.9 mmol m⁻² d⁻¹, ranging from 0.97 to 20 mmol m⁻² d⁻¹ at stations GB2-112 and GB1-85, respectively. Mean PIC fluxes were 1.2 ± 1.7 mmol m⁻² d⁻¹, and ranged from 0.067 to 6.2 mmol m⁻² d⁻¹ at stations GB2-73 and GB1-59, respectively. Finally, mean BSi fluxes at z_{PAR} were 3.8 ± 5.8 mmol m⁻² d⁻¹, ranging from 0.17 to 28 mmol m⁻² d⁻¹ at stations GB2-46 and GB1-85, respectively. Higher POC export stations frequently corresponded with higher BSi export stations (e.g., station GB1-85), but less so with higher PIC export stations.

The highest and lowest measured biomineral (PIC and BSi) fluxes at z_{PAR} were in GB1 and GB2, respectively, but mean values were not significantly different between ocean basins because of high variability within each basin (Fig. 6). However, mean POC fluxes at z_{PAR} were significantly higher in GB1 (mean ± 1 s.d.= 6.0 ± 4.9 mmol m⁻² d⁻¹) than in GB2 (3.0 ± 1.7 mmol m⁻² d⁻¹) (Student's t-test p>0.05). Because POC:²³⁴Th values did not differ between GB1 and GB2 (p<0.05), we attribute this inter-basin difference in POC fluxes primarily to significantly higher ²³⁴Th fluxes in GB1 (1,574 \pm 463 dpm m⁻² d⁻¹) relative to fluxes in GB2 (1,240 \pm 330 dpm m⁻² d⁻¹).

Further, there were significant latitudinal differences among export fluxes and particulate composition ratios in three temperature/nutrient regimes across both sectors (Fig.1; Table 4): (1) north of 45° S, the approximate location of the Subantarctic front, where temperatures exceeded ~10°C; (2) south of 52 °S, the approximate location of the Polar Front (e.g., (Belkin and Gordon, 1996; Sokolov and Rintoul, 2009), where temperatures remained below ~5°C; and (3) between 45° S and 52 °S, where temperatures ranged from ~5-10 °C. The >51 µm size-fraction POC:²³⁴Th values at z_{PAR}

were significantly lower in the most equatorward zone north of 45 °S, where average ratios were $1.9 \pm 0.9 \ \mu\text{mol dpm}^{-1}$. The highest average ratios, south of 52°S, were $5.4 \pm 3.0 \ \mu\text{mol dpm}^{-1}$, illustrating the wide variation in POC:²³⁴Th ratios with ecosystem type (Buesseler et al., 2006; Jacquet et al., 2011). Likewise, zonally averaged POC export fluxes in the most equatorward zone $(2.7 \pm 2.3 \ \text{mmol m}^{-2} \ \text{d}^{-1})$ were significantly lower than average fluxes in the most poleward zone $(8.0 \pm 6.3 \ \text{mmol m}^{-2} \ \text{d}^{-1})$. BSi:²³⁴Th values were significantly different in all three zones, with highest average ratios south of 52°S ($7.1 \pm 4.1 \ \mu\text{mol dpm}^{-1}$) and smallest ratios north of 45°S ($0.3 \pm 0.1 \ \mu\text{mol dpm}^{-1}$). Similarly, average BSi export fluxes were also significantly different from each other in all three zones, with the greatest average values south of 52°S ($10 \pm 8.7 \ \text{mmol m}^{-2} \ \text{d}^{-1}$), and lowest values north of 45°S ($0.35 \pm 0.16 \ \text{mmol m}^{-2} \ \text{d}^{-1}$). Finally, PIC:²³⁴Th ratios, which averaged $0.72 \pm 0.85 \ \mu\text{mol dpm}^{-1}$ across all zones, and PIC export fluxes were not significantly different from each other in any zone defined by these latitudinal bands.

These fluxes are sensitive to the choice of export depth (z_{PAR} or $z_{Th/U}$), not only because the export depth determines the magnitude of ²³⁴Th flux by influencing the integrated ²³⁴Th deficit, but also because the export depth determines which POC:²³⁴Th ratio best describes particles sinking from the chosen depth (Appendix A, Fig. S1). Across stations, the depth metrics z_{PAR} and $z_{Th/U}$ differed from each other to varying extents (Fig. 4; Table 1). As exemplified by stations GB1-92, GB1-16, and GB2-100, POC fluxes changed significantly between z_{PAR} and $z_{Th/U}$ (Figs. 5b, 5c; Appendix A, Table S2). At station GB1-92, where z_{PAR} was 40 m shallower than $z_{Th/U}$, POC flux decreased from 8.0 mmol m⁻² d⁻¹ at z_{PAR} to 5.1 mmol m⁻² d⁻¹ at $z_{Th/U}$. In contrast, at station GB1-16, where z_{PAR} was 80 m shallower than $z_{Th/U}$, POC fluxes increased from 5.9 mmol $m^{-2} d^{-1}$ to 6.6 mmol $m^{-2} d^{-1}$. At station GB2-100, one of few stations where z_{PAR} was deeper than $z_{Th/U}$, POC fluxes decreased from 3.3 to 1.5 mmol m⁻² d⁻¹ going deeper. At this station, the POC:²³⁴Th ratio at $z_{Th/U}$ was 102% greater than ratios at z_{PAR} , while ²³⁴Th fluxes at $z_{Th/U}$ were 6% greater than fluxes at z_{PAR} , demonstrating that changes in particle composition disproportionately contributed to the observed difference in POC export at z_{PAR} and $z_{Th/U}$. By contrast, at station GB1-16, the relative change in ²³⁴Th fluxes from z_{PAR} to $z_{Th/U}$ (+29%) contributed more to the increase in POC flux with depth than the relative change in POC:²³⁴Th ratio (-13%). Finally, for station GB1-92, the relative

change in ²³⁴Th flux with depth (-19%) was similar to the relative change in POC:²³⁴Th with depth (-21%), demonstrating that the export flux estimate was equally sensitive to changes in both parameters.

4 Discussion

The following discusses these flux measurements in the context of other Southern Ocean observations, and hypotheses surrounding the transformation of sinking organic carbon within the euphotic and mesopelagic zones of the water column.

4.1 Choice of export depth

The two possible depths we use to calculate export flux, z_{PAR} and $z_{Th/U}$, are significantly different in the Atlantic sector, which influences the magnitude of flux estimated (see Section 3). We offer here two possible and not mutually exclusive explanations for why $z_{Th/U}$ depths were on average deeper than z_{PAR} depths at GB1 stations.

One hypothesis is that the ²³⁴Th - ²³⁸U profiles used to calculate export fluxes may not have been at steady-state during the time of sampling on the GB1 cruise. Non-steady state conditions in the ²³⁸U-²³⁴Th system do occur during phytoplankton blooms, particularly during their decline and ascent (Savoye et al., 2006; Buesseler et al., 2009). For example, a recent and rapid increase in the near-surface particle concentration could decrease the depth of light penetration faster than the ²³⁸U-²³⁴Th system can adjust, leading to a z_{PAR} measured on station that is shallower than the $z_{Th/U}$, which reflects conditions prior to the rapid increase. Since the GB1 cruise in the Atlantic sector took place a month earlier in the growing season (January-February 2011) than the GB2 cruise in the Indian sector (February-March 2012), the two sectors may have been sampled at different stages of the seasonal bloom, contributing to differences in agreement between z_{PAR} and $z_{Th/U}$. Satellite chlorophyll time-series, if well-resolved, can shed light on how dynamic primary production was around the time of sampling at each station of GB1 and GB2, whether rapid (i.e., within three weeks) changes in particle production and sinking
fluxes from a bloom could have decoupled ²³⁴Th-²³⁸U deficits from light profiles into the surface ocean of the Great Calcite Belt. Eight-day composites of chlorophyll imagery from December 2010 to February 2011 were required to overcome spatial patchiness in the data due to clouds, and indicate that the changes leading up to sampling during GB1 were not consistent across all stations where $z_{PAR} < z_{Th/U}$. At several stations, chlorophyll concentrations declined towards the sampling date; at others, chlorophyll did not change or increased towards the sampling date. Moreover, out of the three stations where $z_{PAR} = z_{Th/U}$, only one exhibited relatively constant chlorophyll concentrations in the month preceding sampling. In GB2, where the differences between z_{PAR} and $z_{Th/U}$ were not significant, chlorophyll tended to be constant preceding more sampling stations. Nonetheless, as in GB1, several locations still experienced increasing or decreasing chlorophyll concentrations in the weeks before sampling, despite having a similar z_{PAR} and $z_{Th/U}$.

The inability of the chlorophyll time-series to unequivocally resolve the differences between z_{PAR} and $z_{Th/U}$ points to other possible mechanisms underlying the discrepancy. One other mechanism, which does not necessarily preclude non-steady state in the ²³⁴Th system, is sinking particle production below the euphotic zone z_{PAR} (Trull et al., 2008). Physical aggregation and fecal pellet production by zooplankton grazing in the region below z_{PAR} (i.e., the upper mesopelagic zone) can increase the speed and total abundance of sinking of particles by transforming phytoplankton biomass exiting the euphotic zone, thereby contributing to sustained ²³⁴Th deficits below z_{PAR} (Steinberg et al., 2008; Wilson et al., 2008; Abramson et al., 2010). Why this occurs only in GB1 and not GB2 is not known.

For example, the ~70m difference in z_{PAR} and $z_{Th/U}$ at a station like GB1-85 (Table 1) may be attributed to additional production or repackaging of sinking particles in the upper mesopelagic zone, causing ²³⁴Th deficits to persist beyond the euphotic zone of primary productivity, and a deeper $z_{Th/U}$. Images of >51 µm particles from this station highlight the changing nature of >51 µm particles with depth (Fig. 2), from primarily large phytoplankton in the euphotic zone to predominantly fecal pellets in the mesopelagic zone. The difference in POC fluxes measured at both depths may arise from the evolution of these particles during vertical transit, from predominantly intact and

relative buoyant diatoms at z_{PAR} to degraded, sinking fecal pellets produced between z_{PAR} and $z_{Th/U}$.

Going forward, it is most important to keep in mind how the choice of export depth impacts flux estimates. For this study, all export fluxes are defined by z_{PAR} so that they can be compared with integrated primary production measurements (Buesseler and Boyd, 2009). Non-steady-state effects of ²³⁴Th profiles on export fluxes will not be considered further because we do not have Lagrangian observations at multiple time points necessary to detect such effects (Buesseler et al., 2003; Resplandy et al., 2012).

4.2 Comparison of export fluxes to previous studies

The ²³⁴Th fluxes we report (mean \pm S.D.= 1.413 \pm 432 dpm m⁻² d⁻¹) are generally within range of measurements from other Southern Ocean studies $(1.615 \pm 1.050 \text{ dpm m}^{-1})$ ²d⁻¹) (compilation by Le Moigne et al., 2013: Shimmield et al., 1995; Rutgers Van Der Loeff et al., 1997; Buesseler, 1998; Cochran et al., 2000; Buesseler et al., 2001; Friedrich and van der Loeff, 2002; Buesseler et al., 2003; Coppola et al., 2005; Morris et al., 2007; Thomalla et al., 2008; Savoye et al., 2008; Rodriguez y Baena et al., 2008; Jacquet et al., 2011; Rutgers van der Loeff et al., 2011; Zhou et al., 2012; Planchon et al., 2013). By contrast, the POC fluxes we report $(4.5 \pm 3.9 \text{ mmol m}^{-2} \text{ d}^{-1})$ are on average three times lower than fluxes from other studies $(12.6 \pm 13.3 \text{ mmol m}^{-2} \text{ d}^{-1})$ due to lower POC:²³⁴Th ratios measured in >51 µm particles. In general, POC:²³⁴Th ratios can vary widely as a function of season, ecosystem composition, size-fraction, depth, and particle sampling methodology (Coppola et al., 2005; Buesseler et al., 2006; Santschi et al., 2006; Jacquet et al., 2011). In GB1 and GB2, an ecosystem effect likely accounts for the 14-fold difference in POC:²³⁴Th between oligotrophic waters (e.g. 0.8 µmol dpm⁻¹ at GB2-106) and polar waters (e.g., 10.8 µmol dpm⁻¹ at GB1-85) (Table 3). The Le Moigne et al. (2013) dataset may include more studies from diatom-rich ecosystems with high POC:²³⁴Th organic particles, such as observed by (Buesseler, 1998; not included in Le Moigne et al. 2013), driving some of the discrepancy between our observations and POC fluxes reported by (Le Moigne et al., 2012).

Other potential reasons for POC:²³⁴Th differences are the choice of export depth (see Section 4.1) and different sampling methodologies in the previous studies. For instance, in-situ pump filter holders with a small-diameter central intake and thus higher intake velocities have been observed to sample more zooplankton, which typically have higher POC:²³⁴Th ratios, than filter holders with diffuse intakes (Bishop et al., 2012). This is because swimming zooplankton can avoid the gentle intake velocities of filter holders with diffuse intakes but not the higher velocities of small diameter intakes. This would be expected to affect estimates of ²³⁴Th-derived POC flux more than ²³⁴Th-derived biomineral fluxes.

There have been far fewer estimates of ²³⁴Th-derived biomineral export fluxes (Thomalla et al., 2008; Sanders et al., 2010; Le Moigne et al., 2012; Le Moigne et al., 2013). BSi and PIC fluxes observed during GB1 and GB2 are within the range previously observed during the *Crozex* study by the Crozet islands (Le Moigne et al., 2012), the site of station GB2-27. Thomalla et al. (2008) also reported biomineral fluxes from the Atlantic Meridional Transect (AMT), north of the Subantarctic Front. While AMT PIC export fluxes were only two times smaller than our mean PIC fluxes in the Great Calcite Belt region, AMT BSi fluxes were ten times smaller. The disparity in BSi fluxes is unsurprising, since the AMT cruise track was through waters with low abundance of silicifiers. We also find that the PIC and BSi fluxes from our Great Calcite Belt study are 4 and 10 times larger than biomineral fluxes estimated by Henson et al. (2012b), respectively, who used a steady-state model of nutrient uptake against nutrient export (Sarmiento et al., 2002; Sarmiento et al., 2004). The Henson et al. method used annual climatologies of nutrient concentration profiles for their estimates, whereas the ²³⁴Th-derived export method used here integrates over several weeks in the growing season. This difference in timescales of integration likely accounts for the smaller biomineral fluxes in Henson et al. (2012b).

4.3 Export efficiency

We found no significant relationship observed between integrated primary productivity and POC flux at z_{PAR} , highlighting the variable export efficiency across GB1

and GB2. Export efficiencies, or "Ez-ratios" (Buesseler and Boyd, 2009), were calculated as the ratio of POC flux at z_{PAR} to total integrated primary production in the euphotic zone (Fig. 7b; Table 3). Mean export efficiencies were 0.26 ± 0.19 , and ranged from 0.04 to 0.77 at stations GB1-16 and GB2-63, respectively. The lack of association between primary productivity and POC export flux confirms previously observed decoupling between the factors that drive export and those that modulate primary productivity (Buesseler et al., 2001; Coppola et al., 2005; Maiti et al., 2010).

4.4 Vertical attenuation of POC flux and concentration

At most stations, both POC flux and >51 μ m [POC] decline with depth below z_{PAR} as a result of remineralization. In the following, we use two metrics to describe POC transfer in the mesopelagic zone: (1) the attenuation of >51 um [POC] in the mesopelagic zone, expressed as the attenuation coefficients extracted from power-law fits of mesopelagic >51 μ m [POC] (exponent from Eq. 1) and (2) the POC flux transfer efficiency (T₁₀₀), defined as the fraction of ²³⁴Th-based POC flux that survives remineralization and is transferred 100 m below z_{PAR} (Buesseler and Boyd, 2009). The first metric describes the disappearance of POC concentration, and applies to the entire mesopelagic zone; the second metric describes the survival of POC flux, and applies to the upper mesopelagic zone.

The mean T_{100} was 0.71 ± 0.38, ranging from 0.20 to 1.8 at stations GB2-119 and GB1-25, respectively (Fig. 7c; Table 2), generally falling within the spread of values observed globally as well as specifically in the Southern Ocean (Buesseler and Boyd, 2009). At stations GB1-6, GB1-16, GB1-25, GB1-59, and GB2-106, T_{100} values are greater than 1.0 and reflect an increase in POC flux with depth between z_{PAR} and 100 m below z_{PAR} (Figs. 5b, 5d). Transfer efficiencies greater than 1 can occur during a declining bloom (Henson et al., 2015), but examination of satellite chlorophyll time-series does not indicate that these stations were sampled at such conditions. At GB1-6, GB1-16 and GB1-59, the ²³⁴Th-²³⁸U disequilibrium extends relatively deep (>200m) into the water column, thus leading to continually increasing ²³⁴Th flux with depth, suggesting that either renewed particle production at depth or lateral advection of particles away

from these coastal stations could sustain the ²³⁴Th deficit below z_{PAR} . Moreover, because z_{PAR} depths are significantly shallower than $z_{Th/U}$ in most GB1 stations, including GB1-6, GB1-16 and GB1-59, the transfer efficiency calculation at these stations in GB1 captures an increase in ²³⁴Th flux between z_{PAR} and 100 m below z_{PAR} . Thus, for the following discussion, it is important to view transfer efficiency values with the caveat that GB1 and GB2 stations display different ²³⁴Th-²³⁸U disequilibria profiles with respect to z_{PAR} and $z_{Th/U}$, and this difference impacts all calculations that use a ²³⁴Th flux component.

At the two other stations for which $T_{100}>1$, GB1-25 and GB2-106, the increases in POC flux below z_{PAR} arise primarily from increasing POC:²³⁴Th ratios rather than increasing ²³⁴Th flux with depth (Appendix A, Figs. S1a, S1d). The increase in these ratios results from a faster decrease in particulate ²³⁴Th activity compared to changes in >51 µm [POC] with depth. This is unexpected and at all other stations, >51 µm [POC] decreases more quickly than particulate ²³⁴Th activity due to organic carbon remineralization. We suspect that poor >51 µm particle distribution on filters from GB2-106 may have led to anomalously low POC around z_{PAR} , but do not have an explanation for the consistent increase in POC:Th with depth at GB1-25 (Appendix A, Figs. S1a). We proceed by excluding the T_{100} transfer efficiencies from these two stations from statistical tests, but identify them for completeness (Figs. 7, 9).

The general decline in POC flux with depth at most stations is mirrored by a decrease in >51 μ m [POC], both of which are a result of remineralization. Attenuation coefficients from power-law fits of mesopelagic >51 μ m [POC] at 22 stations describe this transformation from z_{PAR} to the lower mesopelagic zone, where >51 μ m [POC] between 800-1000 m was 1.5 to 137 times lower than >51 μ m [POC] at z_{PAR} (Figs. 8b, 8c; Table 2). We discount the attenuation value at station GB2-93 from discussion because it had an anomalously low >51 μ m [POC] at 800m, likely due to incomplete rinsing of particles from the prefilter. This drove the power law fit to yield an anomalously high attenuation coefficient, an outlier, as approximated by Chauvenet's Theorem (Glover et al., 2011). Attenuation coefficients were 1.1 ± 0.50 on average, and varied from 0.4 to 1.9 at stations GB1-25 and GB2-43, respectively (Fig. 8c; Table 2), which spans the global range compiled by (Lam et al., 2011).

The >51 μ m [POC] at z_{PAR} is not correlated with >51 μ m [POC] at lower mesopelagic depths, suggesting that processes controlling >51 μ m [POC] at the top of the mesopelagic differ from those controlling >51 μ m [POC] at the base of the mesopelagic zone. This is supported by the great variation in attenuation coefficients and transfer efficiencies, and suggests that POC concentrations at z_{PAR} are decoupled from [POC] at z >800m, as has also been noted in other POC flux and concentration observations (Lomas et al., 2010; Lam et al., 2011; Henson et al., 2012b). There are some exceptions, such as at GB1-85, which exhibited the highest >51 μ m [POC] both at z_{PAR} and below 800 m, but there is no overall relationship across the dataset. The remaining discussion aims to tease apart the processes that control POC flux and >51 μ m [POC] in each depth regime.

4.5 Biomineral-POC flux correlations at z_{PAR}

We compared POC fluxes to mineral fluxes at z_{PAR} (Figs. 9a, 9b) to test the hypothesis that mineral ballasting facilitates POC export out of the euphotic zone, as has been observed in deeper flux datasets >1000 m (Klaas and Archer, 2002; Armstrong et al., 2002; Francois et al., 2002). Because we use ²³⁴Th activity deficits and the same particulate ²³⁴Th activities to derive all fluxes (Eq. (4-6), comparing export fluxes is equivalent to comparing concentrations of >51 µm POC, BSi and PIC at z_{PAR} . In this dataset, minor differences between flux versus concentration comparisons (not shown) arise from differences in interpolation methods for POC:²³⁴Th, BSi:²³⁴Th, and PIC:²³⁴Th ratios at z_{PAR} (Table 3).

Pearson correlation tests between shallow POC export and the two biomineral fluxes revealed a significantly positive correlation between POC and BSi fluxes (p<<0.001, r^2 =0.77). By contrast, there was no significant relationship between shallow POC and PIC fluxes (p=0.24, r^2 =0.06). Both BSi and POC export fluxes tend to increase poleward from the region north of the Subtropical/Subantarctic fronts to the inter-frontal zone to the region south of the Polar front (Figs. 5b, 6a, 6b). Station GB1-85, which sits just south of the Polar Front (~52°S), is a high BSi and POC flux outlier. When removed, the BSi flux vs. POC flux correlation remains significant, though weaker (r^2 = 0.43),

suggesting that although this correlation is strongly influenced by station GB1-85, the shallow BSi ballast association still remains valid for the rest of the dataset.

We also compared POC export fluxes to both PIC and BSi export fluxes simultaneously by multiple linear regression:

POC Flux = $(m_{BSi} \times BSi Flux) + (m_{PIC} \times PIC Flux) + constant$ (7) The multiple linear regression only explains an additional 5% of the variance in POC flux at z_{PAR} ($r^2=0.82$, p<< 0.001), affirming that BSi flux explains most of the variation in POC export fluxes at z_{PAR} across the Atlantic and Indian sectors of the Great Calcite Belt region.

The per-mole carrying capacities of BSi and PIC for POC, or the slopes m_{BSi} and m_{PIC} in the multiple linear regression Eq. (7), are 0.60 and 0.50, respectively. The perweight carrying capacities of BSi and PIC for POC are 0.23 and 0.13, respectively, assuming 12 x 2.199 g mol⁻¹ POC, 67.3 g SiO₂0.4H₂O mol⁻¹ BSi and 100.1 g CaCO₃ mol⁻¹ PIC (Klaas and Archer 2002). The unassociated POC flux, the constant in Eq. (7), is 1.7 mmol POC m⁻² d⁻¹, or 44 mg POC m⁻² d⁻¹. These carrying capacities for POC are 2-10 times higher than global biomineral carrying capacities of deeper (>2000m) flux data (m_{BSi} =0.025-0.026, m_{PIC} =0.070-0.074; Klaas and Archer, 2002), reflecting how POC remineralization with depth consistently reduces apparent mineral carrying capacities between the base of the euphotic zone and the deep sea.

These upper ocean carrying capacities, especially m_{PIC} , are considerably different than corresponding per-weight carrying capacities reported in the *Crozex* study in the Indian sector of the Southern Ocean (m_{BSi} = 0.16, m_{PIC} = -0.11, constant=105 mg POC m⁻² d⁻¹) (Le Moigne et al., 2012). But, as the *Crozex* study was carried out several months earlier in the growing season than our sampling of the same area within the Great Calcite Belt, seasonal changes in the phytoplankton communities and their associated food webs could account for the differences in upper ocean carrying capacities. The Le Moigne et al. (2012) study also highlighted that variable ecosystem composition contributed to regional variations in upper ocean carrying capacities (Le Moigne et al. 2014), echoing a contemporaneous study that showed that even the deep (>1500 m) flux carrying capacities have statistically significant spatial variability (Wilson et al., 2012). It is worth noting that Le Moigne et al. (2012) included lithogenic minerals in their multiple linear regressions. We did not measure lithogenic minerals on GB1 and GB2, as we assumed lithogenic fluxes to be small in the Southern Ocean due to low terrestrial dust inputs (e,g., Honjo et al., 2000). While omitting this lithogenic component from the multiple linear regression could potentially impact derived m_{BSi} and m_{PIC} values, lithogenic material is nonetheless unlikely to be an important carrier of POC flux because of its low flux in the Southern Ocean. Indeed, regional studies have found that the lithogenic carrying capacity (Wilson et al., 2012) and the lithogenic-associated POC fluxes (Le Moigne et al., 2012) are very low in the Southern Ocean.

4.6 Mineral-POC flux correlations in the mesopelagic zone

To directly test whether minerals facilitate POC transfer through the upper mesopelagic zone of the water column as well, we compared flux transfer efficiencies 100 m below the base of the euphotic zone (T_{100}) with BSi and PIC fluxes at z_{PAR} (Figs. 9c, 9d). If the mineral ballast model were to apply to the upper mesopelagic zone, one would expect greater transfer efficiencies (i.e., lower attenuation of POC flux) in regions of higher mineral export. The data highlight several key differences between the role of minerals in the euphotic and upper mesopelagic zones. For one, the correlation between PIC flux and T_{100} , excluding values at GB1-25 and GB2-106, is significantly positive (p<0.001, r²=0.39). The relationship remains even when assessing data from each cruise individually (for GB1, p=0.047, r²= 0.34; for GB2, p= 0.009, r²=0.52), lending further support to a potential role for PIC in POC transfer through the upper mesopelagic zone.

Further, there was no significant correlation, with or without GB1-25 and GB2-106 T₁₀₀ values, between BSi export fluxes in GB2 and T₁₀₀. However, higher particulate biogenic silica concentrations (>51 μ m [BSi]) at z_{PAR} did correspond with greater attenuation of >51 μ m [POC] below z_{PAR} (p=0.004, r²=0.35; Fig. 10a), suggesting that in contrast to its role in the euphotic zone, BSi is associated with greater degradation in the mesopelagic zone of the water column.

4.7 Other controls on POC transfer

The correlation between the attenuation of >51 µm [POC] and the size fractionation of POC (% >51 µm [POC]) at z_{PAR} is even stronger than with >51 µm [BSi] (p<<0.001, r²=0.63; Fig. 10b). GB1-85 appears to be an outlier for both relationships in Fig. 10, but especially for the relationship between >51 µm [POC] attenuation and >51 µm [BSi] (Fig. 10a). The correlation remains significant when the high [BSi] value from station GB1-85 is removed. Notably, the power law fit at GB1-85 is not very good in the upper mesopelagic; fitting >51 µm [POC] between z_{PAR} and 500 m yields a better fit (higher r²; see Fig. 3) with a higher attenuation coefficient of 2.35 (compared to 1.7 for the entire mesopelagic zone). This modified upper mesopelagic attenuation at GB1-85 improves the overall correlations between the attenuation coefficient and both >51 µm [BSi] (p<<0.001, r²=0.60) and % >51 µm [POC] (p<<0.001 ; r²=0.78), further strengthening the argument that >51 µm [BSi] and % >51 µm [POC] at z_{PAR} are important factors in POC transfer in the upper mesopelagic zone.

The relationships between the attenuation of >51 μ m [POC] and >51 μ m [BSi] and particle size fractionation may arise from a more fundamental feature shared by both high-[BSi] and large-particle stations of the Great Calcite Belt: diatom-rich phytoplankton communities. Indeed, we also observe a strong correlation between >51 μ m [BSi] and % >51 μ m [POC] at z_{PAR} (p<<0.001, r²= 0.65; not shown). This is a consistent feature across diatom-rich populations, which produce large, BSi-rich organic aggregates that sink rapidly out of the euphotic zone (Michaels and Silver, 1988; Buesseler, 1998; Thomalla et al.). Indeed, euphotic zone diatom abundances enumerated with a FlowCam® are significantly correlated with >51 μ m [BSi] at zPAR at corresponding stations in GB1 and GB2 (Appendix A, Fig. S2a). Thus, characteristics describing ecosystem structure may underlie the correlation between BSi export and POC export in the Great Calcite Belt (Francois et al., 2002; Thomalla et al., 2008; Henson et al., 2012a; Henson et al., 2012b).

However, ecosystem composition does not directly explain why larger particles exported into the mesopelagic zone are remineralized more vigorously hundreds of meters below (Fig. 10b). It is paradoxical that the same large particles that sink quickly out of the euphotic zone would then remineralize faster, as well. This association between attenuation coefficient and particle size suggests that these particles sink more slowly than expected in the mesopelagic zone given their size (for example, as a result of high porosity and low excess density), and/or that they are subject to faster remineralization compared to regions with more POC in the small size fraction. Francois et al. (2002) noted a negative relationship between bathypelagic transfer efficiency and opal flux, and attributed this to increased lability in large diatom aggregates. Though we do not observe any negative correlation between upper mesopelagic transfer efficiency (T₁₀₀) and BSi fluxes at z_{PAR} , we suggest that potentially higher degradability of POC produced by diatom-rich communities may similarly explain the relationship between particle size and >51 µm [POC] attenuation in the upper mesopelagic zone.

The view of POC quality as a driving factor behind POC transfer argues for a deterministic role of euphotic zone community structure in POC transfer below the euphotic zone. It supports the conventional perspective that diatom-dominated communities are strong exporters of large, sinking POC particles out of the euphotic zone (Buesseler, 1998; Guidi et al., 2009), but also adds to the growing view that these communities have poor transfer efficiency and high attenuation through the mesopelagic zone (Francois et al., 2002; Guidi et al., 2009; Henson et al., 2012b; Henson et al., 2012a).

For instance, station GB1-85, with over half of [POC] in the >51 μ m size class fraction in the euphotic zone (Fig. 6c; Table 3), has a low >51 μ m [PIC]:[BSi] ratio of 0.035 at z_{PAR} (indicated in log-scale in Figs. 10a and 10b), which indicates relatively high diatom populations producing large BSi-rich aggregates (Appendix A, Figs. 2, S2, S3). Station GB1-85 exhibits a high export efficiency (Ez-ratio= 0.38, within the upper quartile of the data set), and the highest >51 μ m [POC] and export fluxes at z_{PAR} (Figs. 5b, 7b, 8a; Table 3). Notably, >51 μ m [POC] values in the lower mesopelagic zone are also the highest at GB1-85, despite attenuating greatly below zPAR (attenuation coefficient = 1.7) (Figs. 3, 8b, 8c; Table 2). But, because of high attenuation, proportionally less organic carbon transfers to the deep sea at GB1-85. The same diatomrich communities that vigorously export POC ultimately may not sequester as much organic carbon in the deep ocean or draw down as much atmospheric CO₂ (Kwon et al., 2009) as would be expected considering the magnitude of export alone. In contrast to a model diatom community like station GB1-85, station GB1-25 is BSi-deplete, with a >51 μ m [PIC]:[BSi] ratio of 1.4 at z_{PAR} (indicated in log-scale in Fig. 10a), indicating relatively more coccolithophores in the community (Appendix A, Figs. S2, S3). With proportionally less POC in the >51 μ m size-fraction (only 3.2%) (Figs. 6c, 10b; Table 3), >51 μ m [POC] at GB1-25 attenuates little through the mesopelagic zone (attenuation coefficient = 0.4, the lowest of the data set) such that a third of the >51 μ m [POC] at z_{PAR} remains at 1000m, compared to only 1.4% at station GB1-85 (Fig. 3). At GB1-25, export efficiency is very low (Ez-ratio=0.04), suggesting that the particles exiting the euphotic zone here have been recycled vigorously in the euphotic zone prior to export, which may explain their low >51 μ m [POC] and high proportion in the <51 μ m size-fraction at z_{PAR}. In the mesopelagic zone, these particles are not very reactive and thus remineralize very little, perhaps sequestering a higher proportion of the CO₂ fixed in the euphotic zone.

Several other stations with proportionally more small particles and weaker >51 μ m [POC] attenuation in the mesopelagic zone exhibit higher >51 μ m [PIC] than >51 μ m [BSi] at z_{PAR} (labeled in the lower left quadrant of Fig. 10b), suggesting that export regimes characterized by high relative abundance of coccolithophores consistently transfer less reactive POC to the mesopelagic zone. Artificial roller tank experiments have demonstrated that coccolithophore cultures can produce smaller, more compact aggregates than diatom cultures, partly because of smaller cell sizes (Iversen and Ploug, 2010). However, smaller size does not necessarily mean slower sinking velocities (e.g., (McDonnell and Buesseler, 2010). Iversen and Ploug (2010) showed that the higher excess density of these small aggregates generated faster sinking speeds than similarly sized pure diatom aggregates. Another roller tank study that compared aggregate formation by calcifying versus non-calcifying coccolithophores observed that aggregates formed from calcifying coccolithophores were smaller but faster sinking (Engel et al., 2009). In regions like the Great Calcite Belt, dense coccolithophore populations may similarly export small, highly degraded and compact particles out of the euphotic zone. As a result, these communities would efficiently transfer POC towards the base of the mesopelagic zone, even if the magnitude of exported POC is not as high as in diatom-rich regions (Thomalla et al., 2008; Guidi et al., 2009; Henson et al., 2012b). This may

explain why higher PIC export fluxes are associated with higher transfer efficiencies but not higher POC flux at z_{PAR} (Fig. 9), and also why the ballast association between PIC and POC fluxes appears only at greater depths (Francois et al., 2002; Klaas and Archer, 2002).

Attenuation coefficients for >51 µm [POC] across diatom-rich regions exhibit a great spread (standard deviation= 0.47), ranging from 0.47 to 1.88. Not all diatom-rich stations (i.e., >51 µm [PIC]:[BSi]<1 at z_{PAR}) have proportionally larger particles or higher b-values (e.g., stations GB1-70, GB1-77 and GB2-87; Fig. 10b). In contrast, attenuation coefficients across coccolithophore-rich regions (i.e., >51 µm [PIC]:[BSi] \geq 1 at z_{PAR}) exhibit a lower standard deviation (0.31) and a smaller range, 0.35 to 1.12. The greater variance in attenuation across BSi-rich regions may result from sampling the diatom populations at different seasons of the bloom cycle (Lam et al., 2011), and implies that there may be less seasonality in POC transfer to depth in coccolithophore-rich regions. Indeed, massive diatom export events with high transfer efficiency through the mesopelagic zone have been observed (Martin et al., 2011; Smetacek et al. 2012), so there are clearly conditions that can lead to efficient mesopelagic POC transfer from diatom blooms.

It is worth noting that >51 μ m [PIC]:[BSi] ratios did increase with depth at most stations of the Great Calcite Belt, as might be expected because BSi is undersaturated in seawater. The possibility that BSi dissolves faster than PIC in particles sinking through the mesopelagic zone would complicate the connections we draw between diatom-rich communities in the euphotic zone and the attenuation of >51 μ m [POC]. But, there are no associations between the magnitude of [PIC]:[BSi] increase and >51 μ m [BSi] at z_{PAR}, >51 μ m [PIC] at z_{PAR} or >51 μ m [POC] attenuation with depth, suggesting that the issue of differential dissolution should not significantly impact our earlier interpretations. In the future, directly evaluating the degradability of sinking POC using organic characterization techniques (e.g., ramped pyrolysis or biomarker isolation) (e.g., (Wakeham et al., 2002; Rosenheim et al., 2008; Rosenheim and Galy, 2012; Rosenheim et al., 2013) would greatly improve our ability to track the transformation of POC produced by different ecosystem assemblages across the Great Calcite Belt.

5 Conclusion

In summary, we argue here that phytoplankton assemblages play a fundamental role (Francois et al., 2002; Thomalla et al., 2008; Henson et al., 2012b; Henson et al., 2012a) in determining the fate of POC export through the Great Calcite Belt region, the effect of which sometimes, but not always, appears as a mineral ballast mechanism in the euphotic zone (Lam et al., 2011; Henson et al., 2012a; Lima et al., 2014). Though shallow BSi export fluxes were strongly correlated with POC export fluxes, they are also associated with diatom communities that produce larger particles that attenuate more quickly through the mesopelagic zone, such that proportionally less POC reaches the lower mesopelagic zone, and proportionally more is returned to the water column as remineralized carbon (dissolved inorganic and organic carbon).

Author contributions:

S.Z. Rosengard, the primary author, participated in the GB2 field work, sample analysis in lab and writing. P.J. Lam contributed to field work during GB2, and participated in both data interpretation and editing the manuscript. W.M. Balch, supplied the primary productivity and light profile data included here, as well as coccolithophore and diatom cell count data in the supplement, and provided valuable feedback during writing. M.E. Auro and S. Pike participated in field work and sample analysis during and after GB1. D. Drapeau and B. Bowler contributed to field work during both GB1 and GB2, as well as sample analysis, particularly with respect to primary productivity and light profile data.

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Table 1. Locations and times of sampling of total 234Th and size-fractionated particles on cruises GB1 and GB2. Two export depths are indicated: z_{PAR} (depth of 0.3% of surface photosynthetically available radiation) and $z_{Th/U}$ (depth where ²³⁴Th and ²³⁸U activities return to secular equilibrium below surface deficits).

Cruise	Station	Date	Lat.	Long.	ZPAR	Z _{Th/U}
-	-	dd-mm-yy	deg. N	deg. E	т	m
GB1	6	14 Jan 2011	-51.79	-56.11	79	130
GB1	16	17 Jan 2011	-46.26	-59.83	62	141
GB1	25	20 Jan 2011	-45.67	-48.95	62	115
GB1	32	22 Jan 2011	-40.95	-46.00	69	171
GB1	38	24 Jan 2011	-36.52	-43.38	121	121
GB1	46	26 Jan 2011	-42.21	-41.21	63	100
GB1	59	29 Jan 2011	-51.36	-37.85	60	95
GB1	70	1 Feb 2011	-59.25	-33.15	100	100
GB1	77	3 Feb 2011	-57.28	-25.98	98	100
GB1	85	5 Feb 2011	-53.65	-17.75	73	140
GB1	92	7 Feb 2011	-50.40	-10.80	59	100
GB1	101	9 Feb 2011	-46.31	-3.21	81	140
GB1	109	11 Feb 2011	-42.63	3.34	76	130
GB1	117	13 Feb 2011	-38.97	9.49	62	110
GB2	5	21 Feb 2012	-36.94	39.60	78	90
GB2	27	26 Feb 2012	-45.82	51.05	105	105
GB2	36	28 Feb 2012	-46.84	58.25	90	90
GB2	43	1 Mar 2012	-47.53	64.01	108	125
GB2	53	3 Mar 2012	-49.30	71.32	81	100
GB2	63	5 Mar 2012	-54.40	74.54	109	130
GB2	73	7 Mar 2012	-59.71	77.73	93	75
GB2	87	10 Mar 2012	-54.23	88.22	107	100
GB2	93	11 Mar 2012	-49.81	94.13	113	130
GB2	100	14 Mar 2012	-44.62	100.50	113	90
GB2	106	16 Mar 2012	-40.10	105.34	102	95
GB2	112	17 Mar 2012	-40.26	109.63	76	105
GB2	119	20 Mar 2012	-42.08	113.40	92	90

Table 2. POC fluxes, concentrations, and attenuation of >51 μ m [POC] in the mesopelagic zone. Attenuation coefficient is the exponent from significant power law fits to >51 μ m [POC]. z_{PAR} +100m is 100 m below z_{PAR} , as defined in the Table 1 caption. Transfer efficiency is POC flux at z_{PAR} +100m divided by POC flux at z_{PAR} . Deep >51 μ m [POC] was measured at 1000 m and 800 m for GB1 and GB2, respectively. POC flux errors are propagated from ²³⁴Th flux, and POC:²³⁴Th errors.

Cruise	Station	Depth	>51 µm [POC] tenuation	³⁴ Th Flux at z _{PAR} + 100m	OC:Th at ZPAR + 100m	POC Flux at z _{PAR} + 100m	Transfer Efficiency	>51 µm РОС] (≥800m)
			At	~	Δ .			
-	-	т	unitless	$dpm m^{-2} d^{-1}$	µmol dpm ⁻¹	$mmol m^{-2}d^{-1}$	unitless	μM
GB1	6	179	0.8	$3,319 \pm 128^{\circ}$	1.7	5.7 ± 0.31	1.00	0.030
GB1	16	162	1.1	$2,567 \pm 116^{\circ}$	2.4	6.1 ± 0.30	1.04	No data
GB1	25	162	0.4	$1,074 \pm 125$	2.5	2.7 ± 0.37	1.76	0.013
GB1	32	169	0.9	$1,581 \pm 186$	1.3	2.0 ± 0.25	0.86	0.006
GB1	38	221	No fit	911 ± 206	1.6	1.5 ± 0.35	0.70	0.026
GB1	46	163	1.0	$1,937 \pm 146$	1.6	$\textbf{3.1} \pm \textbf{0.27}$	0.4	0.009
GB1	59	160	0.6	$2,582 \pm 126^{\circ}$	3.7	9.5 ± 0.56	1.29	0.014
GB1	70	200	0.6	$1,414 \pm 248$	3.5	5.0 ± 0.90	0.90	0.024
GB1	77	198	0.5	$1,903 \pm 162$	2.1	4.0 ± 0.41	0.44	0.012
GB1	85	173	1.7 ^a	$2,076 \pm 207$	3.9	8.1 ± 0.83	0.41	0.035
GB1	92	159	1.1	1,339 ± 170	3.7	4.9 ± 0.64	0.61	0.019
GB1	101	181	0.8	1,774 ± 135	1.7	3.0 ± 0.24	0.83	0.019
GB1	109	176	1.0	$1,719\pm97$	1.1	1.9 ± 0.13	0.87	0.006
GB1	117	162	1.1	$1,\!258\pm86$	1.2	1.5 ± 0.13	0.87	0.005
GB2	5	178	0.5	$1,402 \pm 3,706^{\circ}$	1.1	1.5 ± 6.1	0.5	No data
GB2	27	205	No fit	$2,063 \pm 205$	1.2	2.5 ± 0.30	0.71	No data
GB2	36	190	1.5	$1,077 \pm 194$	0.9	0.93 ± 0.18	0.48	0.011
GB2	43	208	1.9	$1,247 \pm 200$	2.2	2.7 ± 0.45	0.54	0.005
GB2	53	181	No fit	$1,013 \pm 220$	2.0	2.0 ± 0.45	0.49	No data
GB2	63	209	1.8	$1,292 \pm 262$	1.7	2.1 ± 0.46	0.31	0.014
GB2	73	193	1.5	807 ± 189	1.9	1.6 ± 0.37	0.48	0.008
GB2	87	207	0.7	$1,213 \pm 196$	1.6	1.9 ± 0.34	0.60	0.013
GB2	93	213	2.3^{b}	469 ± 249	1.6	0.77 ± 0.42	0.53	0.001
GB2	100	213	0.8	$1,132 \pm 190$	0.7	0.80 ± 0.15	0.52	0.014
GB2	106	202	0.9	$1,405 \pm 186$	1.3	1.8 ± 0.26	1.63	0.017
GB2	112	176	1.3	270 ± 186	0.9	0.23 ± 0.21	0.24	0.007
GB2	119	192	No fit	756 ± 218	0.8	0.57 ± 0.17	0.20	0.013

^a attenuation coefficient is 2.35 when only fitting > 51 μ m [POC] measurements at depths <500 m (Fig. 3). ^b outlier approximated by Chauvenet's Theorem (Glover, et al., 2011).

^cvalues were estimated by linear interpolation of values at upper and lower depths around z_{PAR}+100m.

Cruise	Station	ZPAR	²³⁴ Th Flux	>51um [POC]	>51um [Bsi]	>51um [PIC]	>51um Th activity	POC:Th	POC Flux	Bsi:Th	BSi Flux	PIC:Th	PIC Flux	Primary Productivity	Ez- Ratio	% >51 um [POC]
		m	dpm m²d ¹	Mu	Wn	Wn	dpm L ⁻¹	μmol dpm ⁻¹	mmol m²d'	µmol dpm ⁻¹	mmol m²d¹	µmol dpm ⁻¹	mmol m²d'	mmol m ⁻² d ⁻¹	unitless	%
GB1	6	79	$2,437 \pm 100$	0.23 ^b	0.03 ^a	0.124 ^a	0.07 ^a	2.3 ª	5.7 ± 0.26	0.4	0.9 ± 0.04	1.8	4.3 ± 0.20	42	0.14	8.8%
GB1	16	62	$1,933 \pm 71$	0.38	0.08	0.390	0.12	3.0	5.9 ± 0.68	0.6	1.2 ± 0.14	3.1	6.1 ± 0.70	165	0.04	17.7%
GB1	25	62	862 ± 46 ^a	0.04	0.005 ^a	0.015 ^a	0.02	1.8	1.6 ± 0.11	0.2	0.2 ± 0.02	0.7	0.6 ± 0.04	35	0.04	3.2%
GB1	32	69	$1,304 \pm 116$	0.07	0.01	0.027	0.04	1.8	2.3 ± 0.21	0.3	$\textbf{0.3} \pm \textbf{0.03}$	0.7	$\textbf{0.9} \pm \textbf{0.08}$	11	0.21	3.9%
GB1	38	121	809 ± 126	0.04	0.003	0.017	0.01	2.7	2.2 ± 0.35	0.2	0.2 ± 0.03	1.2	0.9 ± 0.15	21	0.10	8.4%
GB1	46	63	$2,123 \pm 69$	0.23	0.005	0.059	0.06	4.1	$\textbf{8.8} \pm \textbf{0.38}$	0.1	0.2 ± 0.02	1.1	2.2 ± 0.10	13	0.67	5.3%
GB1	59	60	$1,844 \pm 102$	0.09	0.10	0.072	0.02	4.0	7.3 ± 0.52	4.6	$\textbf{8.6} \pm \textbf{0.53}$	3.4	6.2 ± 0.39	26	0.28	5.3%
GB1	70	100	1,280 ± 94	0.11	0.06 ^a	0.001 ^a	0.02	4.3	5.5 ± 0.44	3.5 ^a	$\textbf{4.5} \pm \textbf{0.35}$	0.1 ^a	0.1 ± 0.09	10	0.53	10.6%
GB1	77	98	$1,\!485\pm105$	0.03	0.03	0.002	0.01	6.0	9.0 ± 1.3	5.6	$\textbf{8.3} \pm \textbf{0.98}$	0.4	0.7 ± 0.23	57	0.16	3.6%
GB1	85	73	$1,858\pm94$	2.50	3.44	0.124	0.23	10.8	20 ± 1.1	14.9	28 ± 1.5	0.5	1.0 ± 0.05	53	0.38	52.0%
GB1	92	59	$1,639 \pm 77$	0.40	0.46	0.020	0.08	4.9	8.0 ± 0.40	5.6	9.3 ± 0.46	0.2	0.4 ± 0.02	26	0.31	11.3%
GB1	101	81	$1,763 \pm 82$	0.19	0.05	0.013	0.09	2.0	3.6 ± 0.18	0.5	0.9 ± 0.04	0.1	0.2 ± 0.01	22	0.17	12.5%
GB1	109	76	$1,524 \pm 76$	0.19ª	0.05^{a}	0.027^{a}	0.14 ^a	1.4	2.1 ± 0.11	0.4	0.6 ± 0.03	0.2	0.3 ± 0.02	14	0.16	21.0%
GB1	117	62	$1,177 \pm 50$	0.21	0.02	0.032	0.15	1.4	1.7 ± 0.07	0.1	0.2 ± 0.01	0.2	0.3 ± 0.01	18	0.09	6.6%
GB2	5	78	$\textbf{1,889} \pm 5207$	0.08^{b}	0.01 ^a	0.048 ^a	0.05 ^a	1.6	$\textbf{3.0} \pm \textbf{8.8}$	0.2	0.4 ± 1.2	1.0	1.9 ± 5.2	8.2	0.37	7.6%

Table 3. POC, biomineral, and ²³⁴Th concentrations and fluxes at z_{PAR} . Ez- ratio is ²³⁴Th-derived POC flux at z_{PAR} divided by integrated primary productivity. The % >51 μ m [POC] metric is the fraction of total [POC] in the >51 μ m size fraction. POC and biomineral flux errors are propagated from ²³⁴Th flux, and POC:²³⁴Th errors.

GB2	27	105	1,869 ± 160	0.08 ^a	0.10 ^a	0.060 ^a	0.04 ^a	1.9	3.5 ± 0.32	2.2	$\textbf{4.0} \pm \textbf{0.35}$	1.3	2.5 ± 0.22	8.0	0.44	6.7%
GB2	36	90	988 ± 89	0.43	0.28	0.074	0.22	2.0	2.0 ± 0.18	1.3	1.3 ± 0.12	0.3	0.3 ± 0.03	12	0.16	15.6%
GB2	43	108	1,221 ± 153	0.74 ^a	0.62 ^a	0.041 ^a	0.18 ^a	4.1	5.0 ± 0.63	3.4	$\textbf{4.2} \pm \textbf{0.53}$	0.2	0.3 ± 0.04	12	0.43	37.8%
GB2	53	81	1058 ± 100^{a}	0.54 ^a	0.80 ^a	0.081 ^a	0.14 ^a	3.9	4.1 ± 0.40	5.7	6.1 ± 0.59	0.6	0.6 ± 0.07	16	0.25	22.5%
GB2	63	109	$1,229 \pm 138$	0.71 ^a	1.04 ^a	0.028 ^a	0.13 ^a	5.6 ^a	6.9 ± 0.78	8.1	9.9 ± 1.1	0.2	0.3 ± 0.03	9.0	0.77	33.2%
GB2	73	93	977 ± 108	0.21 ^b	1.13 ^a	0.014 ^a	0.20 ^a	3.3 ^a	3.2 ± 0.36	5.6	$\textbf{5.4} \pm \textbf{0.60}$	0.1	0.1 ± 0.01	8.8	0.36	17.6%
GB2	87	107	$1,299 \pm 115$	0.06 ^b	0.30 ^a	0.041^{a}	0.06 ^a	2.5 ^a	$\textbf{3.2} \pm \textbf{0.40}$	4.7	6.1 ± 0.55	0.6	0.8 ± 0.14	11	0.29	3.4%
GB2	93	113	$1,142 \pm 137$	0.07^{a}	0.01 ^a	0.023 ^a	0.05 ^a	1.3	1.5 ± 0.25	0.2	0.3 ± 0.06	0.4	0.5 ± 0.14	12	0.12	4.3%
GB2	100	113	$1,112 \pm 130$	0.08	0.02	0.006	0.06	1.4	1.5 ± 0.19	0.3	$\textbf{0.3} \pm \textbf{0.04}$	0.1	0.1 ± 0.02	14	0.11	12.8%
GB2	106	102	1394 ± 82^{a}	0.09 ^b	0.04 ^a	0.024 ^a	0.12 ^a	0.8	1.1 ± 0.86	0.3 ^a	0.4 ± 0.02	0.2	0.3 ± 0.02	22	0.05	12.2%
GB2	112	76	717 ± 97	0.22 ^b	0.17 ^a	0.087 ^a	0.36 ^a	1.4 ^a	1.0 ± 0.13	0.5	0.3 ± 0.05	0.2	0.2 ± 0.02	no data	no data	13.3%
GB2	119	92	$1,223 \pm 124$	0.51 ^a	0.12 ^a	0.048 ^a	0.22 ^a	2.3	$\textbf{2.8} \pm \textbf{0.29}$	0.5	0.7 ± 0.07	0.2	0.3 ± 0.03	17	0.17	21.5%

^a values at z_{PAR} estimated by linear interpolation of values at upper and lower depths around z_{PAR} . ^b >51 µm [POC] values interpolated by significant power-law fits (Fig. 3). "no data": not enough depths were sampled and analyzed to interpolate at z_{PAR} .

Table 4. Mean \pm standard deviations of ²³⁴Th fluxes, POC:²³⁴Th, BSi: ²³⁴Th, PIC: ²³⁴Th, POC fluxes, and biomineral fluxes at z_{PAR} , divided by three latitude zones. 45 °S marks the approximate latitude of the Subantarctic front, while 52 °S marks the approximate latitude of the Polar front (Belkin and Gordon, 1996; Sokolov and Rintoul, 2009).

Lat. zone	²³⁴ Th Flux at ^{ZPAR}	POC:Th at z _{PAR}	POC Flux at z _{PAR}	BSi:Th at z _{PAR}	BSi Flux at z _{PAR}	PIC:Th at z _{PAR}	PIC Flux at Z _{PAR}	# stn
°S	$dpm m^{-2} d^{-1}$	$\mu mol dpm^{-1}$	mmol $m^{-2}d^{-1}$	$\mu mol dpm^{-1}$	mmol $m^{-2}d^{-1}$	$\mu mol dpm^{-1}$	mmol $m^{-2}d^{-1}$	-
36 - 45	1.3 ± 0.44	1.9 ± 0.9	2.7 ± 2.3	0.3 ± 0.1	0.33 ± 0.17	0.5 ± 0.4	0.73 ± 0.76	10
45 - 52	1.5 ± 0.50	2.8 ± 1.2	4.4 ± 2.2	2.3 ± 2.2	3.4 ± 3.3	1.1 ± 1.2	2.0 ± 2.4	11
52 ->60	1.4 ± 0.30	5.4 ± 3.0	8.0 ± 6.3	7.1 ± 4.1	10 ± 8.7	0.3 ± 0.2	0.49 ± 0.4	6



Figure 1. Cruise tracks across the Atlantic (cruise GB1) and Indian (cruise GB2) sectors of the Great Calcite Belt showing sea surface temperature along the two transects. Station numbers where ²³⁴Th and size-fractionated particles were sampled are indicated by crosses. The two horizontal dashed lines at 45 °S and 52 °S represent the approximate locations of the Subantarctic and Polar fronts, respectively (Belkin and Gordon, 1996; Sokolov and Rintoul, 2009).



Figure 2. Digital images of >51 μ m filters from station GB1-85 (refer to Fig. 1 for station location). >51 μ m particles are from (a) 25m in the euphotic zone, (b) 73m, which corresponds to z_{PAR} , as defined in Table 1, and (c) at 173m, below both metrics of export depth, z_{PAR} and $z_{Th/U}$ (Table 1). >51 μ m particles in the euphotic zone appear as dense sheets of intact cells packed onto the filters (a, b) and as more sparsely arranged cylindrical fecal pellets on filters collected below z_{PAR} (c).



Figure 3. Significant power law fits of >51 μ m [POC] below z_{PAR} , according to Eq. (1). Only the 22 significant fits are shown as lines. Three stations are highlighted to show the range in >51 μ m [POC] attenuation across GB1 and GB2 profiles (symbols represent measurements): GB1-85 had the highest POC concentration through the water column and an attenuation coefficient of 1.7; GB1-25 had the lowest attenuation coefficient (0.4); GB2-43 had the highest attenuation coefficient (1.9) (Table 2). Fitting GB1-85 >51 μ m [POC] measurements between z_{PAR} and 500 m yields a higher attenuation coefficient of 2.35. Refer to Fig. 1 for station locations.





Figure 4. Total ²³⁴Th and ²³⁸U activity profiles measured at 14 stations of GB1 and 13 stations of GB2 (note different x-axis for station GB2-5) (Appendix A, Table S1). Error bars for ²³⁴Th activity are propagated errors. ²³⁸U is calculated from salinity. All ²³⁴Th activity profiles exhibit a deficit relative to ²³⁸U activity at the surface, and mostly return to equilibrium with ²³⁴U within error at depth of $z_{Th/U}$ (Table 1). Refer to Fig. 1 for station locations.



Figure 5. Distribution of ²³⁴Th flux and ²³⁴Th-derived POC flux at 27 stations along GB1 and GB2 (circle area scales with flux magnitude). (a) ²³⁴Th fluxes at z_{PAR} range from 717 dpm m⁻² d⁻¹ to 2,437 dpm m⁻² d⁻¹ at stations GB2-112 and GB1-6, respectively. (b) POC fluxes at z_{PAR} range from 0.97 mmol m⁻² d⁻¹ to 20 mmol m⁻² d⁻¹ at stations GB2-112 and GB1-85, respectively. (c) POC fluxes at $z_{Th/U}$ range from 0.57 to 12 mmol m⁻² d⁻¹ at stations GB2-112 and GB1-85, respectively (Appendix A, Table S2). (d) POC fluxes at 100m below z_{PAR} range from 0.23 to 9.5 mmol m⁻² d⁻¹ at stations GB2-112 and GB1-59, respectively. A few station numbers discussed in the text are indicated. Red outlines distinguish stations where fluxes are greater at the specified depth than at z_{PAR} . The two horizontal dashed lines at 45 °S and 52 °S represent the approximate locations of the Subantarctic and Polar fronts, respectively (Belkin and Gordon, 1996; Sokolov and Rintoul, 2009). Refer to Fig. 1 for other station locations. z_{PAR} and $z_{Th/U}$ are defined as in Table 1.



Figure 6. Distribution of BSi flux, PIC flux, and $\% >51 \ \mu m$ [POC], the percent of total [POC] in the >51 \ \mum m size class, at z_{PAR} (Table 1) along GB1 and GB2 (circle area scales with magnitude). (a) BSi fluxes range from 0.17 mmol m⁻² d⁻¹ to 28 mmol m⁻² d⁻¹ at stations GB1-46 and GB1-85, respectively. (b) PIC fluxes range from 0.067 to 6.2 mmol m⁻² d⁻¹ at stations GB2-73 and GB2-59, respectively. (c) The proportion of [POC] in the >51 \ \mum m size-fraction at z_{PAR} ranges from 3.3% to 52% at stations GB1-25 and GB1-85, respectively. A few station numbers discussed in the text are indicated. The two horizontal dashed lines at 45 °S and 52 °S represent the approximate locations of the Subantarctic and Polar fronts, respectively (Belkin and Gordon, 1996; Sokolov and Rintoul, 2009). Refer to Fig. 1 for other station locations.



Figure 7. Distribution of primary productivity, export efficiency, and transfer efficiency along GB1 and GB2 (circle area scales with magnitude). (a) Primary productivity integrated through the euphotic zone ranges from 8.0 to 165 mmol m⁻² d⁻¹ at stations GB2-27 and GB1-16, respectively. (b) Export efficiency (Ez-ratio) at z_{PAR} (Table 1), which is the ratio of ²³⁴Th-derived POC flux at z_{PAR} to primary productivity integrated to z_{PAR} , ranges from 0.04 to 0.77 at stations GB1-16 and GB2-63, respectively. (c) Transfer efficiency at z_{PAR} , which is the ratio of POC flux 100 m below z_{PAR} to POC flux at z_{PAR} , ranges from 0.20 to 1.8 at stations GB1-119 and GB1-25, respectively. A few station numbers discussed in the text are indicated. The two horizontal dashed lines at 45 °S and 52 °S represent the approximate locations of the Subantarctic and Polar fronts, respectively (Belkin and Gordon, 1996; Sokolov and Rintoul, 2009). Refer to Fig. 1 for other station locations.



Figure 8. Distribution of [POC] and vertical attenuation coefficient of >51 μ m [POC] (circle area scales with magnitude). (a) >51 μ m POC concentrations at z_{PAR} (Table 1) range from 0.03 μ M to 2.5 μ M at stations GB1-77 and GB1-85, respectively. (b) >51 μ m [POC] at the deepest pump depth in the lower mesopelagic zone (800 m-1000 m). Concentrations range from 0.001 μ M to 0.035 μ M at stations GB2-93 and GB1-85, respectively. (c) Attenuation coefficient from significant power-law fits of 22 >51 μ m [POC] profiles, excluding GB2-93 (see Section 4.4). A few station numbers discussed in the text are indicated. The two horizontal dashed lines at 45 °S and 52 °S represent the approximate locations of the Subantarctic and Polar fronts, respectively (Belkin and Gordon, 1996; Sokolov and Rintoul, 2009). Refer to Fig. 1 for other station locations.



Figure 9. ²³⁴Th-derived POC flux as a function of (a) PIC flux and (b) BSi flux at z_{PAR} . POC flux transfer efficiency between z_{PAR} and $z_{PAR}+100$ m (T_{100} , defined in Section 4.4) as a function of (c) PIC flux and (d) BSi flux at z_{PAR} . Significant linear relationships are plotted as a solid blue line. T_{100} values at GB1-25 and GB2-106 were excluded from all correlations (Section 4.4). The color bar indicates the longitude of GB1 and GB2 stations. Refer to Fig. 1 for more specific station locations.



Figure 10. Attenuation coefficient as a function of (a) >51 μ m [BSi] at z_{PAR} and (b) the proportion of [POC] in the >51 μ m size-fraction at z_{PAR} . The open circle indicates where GB1-85 would plot with a higher attenuation coefficient of 2.35, derived from fitting >51 μ m [POC] at depths between z_{PAR} and 500 m. Significant linear relationships using the lower and higher attenuation coefficient values for GB1-85 are shown as solid and dashed lines, respectively; p and r² values are provided for the solid lines. The color bar is the natural logarithm of the ratio of >51 μ m PIC:BSi at z_{PAR} . We interpret all warm colors >0 to indicate stations with a high relative abundance of diatoms (Appendix A, Figs. S2, S3). A few station numbers discussed in the text are indicated. Refer to Fig. 1 for station location.
Chapter 3:

The effect of sample drying temperature on marine particulate organic carbon composition

Abstract

Compositional changes in marine particulate organic carbon (POC) trace important carbon cycle processes that control the biological pump's efficiency. Each particle sampling effort in the ocean water column is labor-intensive, but also a valuable opportunity to glean more information on the organic composition of POC during transfer to the deep ocean. At the same time, not all particle samples are processed and stored at standards accepted by organic geochemists. For example, particles are often air-dried in an oven or at room temperature, rather than immediately frozen at -80°C and subsequently freeze-dried. This study investigates the effects of different drying methods on three aspects of POC composition: bulk, compound-specific and thermal stability. The last metric derives from ramped pyrolysis/oxidation (RPO), a relatively novel method of organic matter characterization that uses a continuous temperature ramp to differentiate components of organic carbon by their decomposition temperature, i.e., thermal stability. Particle samples for the study were collected in June 2014 off Woods Hole, MA and immediately dried at 56°C, at room temperature, or stored at -80°C until freeze-dried for analysis. Results show that oven- and air-drying did not shift the bulk or thermal composition of POC in the samples relative to freeze-drying, but did compromise lipid abundances. The effect on lipid abundances depended on compound size and compound type, suggesting that physical mechanisms such as compound volatilization likely accounted for the drying effects. The data suggest that particle samples collected for intentions other than organic geochemical analyses are still appropriate for assessing bulk POC composition and thermal stability. But, different drying and storage temperatures would likely bias analyses of specific lipid classes, which represent a small fraction of the bulk organic carbon in the sample.

1 Introduction

Particulate organic carbon (POC) in the ocean's water column is the primary vehicle for the biological pump, which sequesters ~ 0.2 Gt carbon/year on the seafloor (~ 0.1 % marine primary productivity) for hundreds to thousands of years (Burdige,

2007). These particles harbor thousands of distinct organic biomolecules produced by primary producers and heterotrophs (Repeta, 2014). Changes in this biomolecular matrix with depth allude to important processes that govern the transport and recycling of marine POC (Burd et al., 2016). Approaches to describing POC composition range from measuring bulk characteristics that integrate over biomolecules (e.g., the stable isotope composition of total POC, reported as δ^{13} C) to quantifying trace abundances of specific biomolecules (e.g., the abundance of specific biomarkers and their δ^{13} C values) (Wakeham and Volkman, 1991; Trull and Armand, 2001; Cavagna et al., 2013).

To accurately interpret the composition of POC in the water column, its organic matrix must be preserved at the time of collection. Prior studies have demonstrated that sample storage temperatures post-collection influences the degree of organic matter preservation. Continual microbial degradation following sample collection and storage at room temperature can significantly alter the distribution of lipid abundances after one month (Grimalt et al., 1988). But, even longer-term sample storage at cooler temperatures can still shift the lipid composition of organic matter (Wakeham and Volkman, 1991). For instance, Ohman (1996) observed that storing copepod biomass at -20°C to -15°C increases the extraction efficiency of free fatty acids by rupturing cell membranes, while also depressing polar lipid abundances by concentrating enzymes in solution that decompose lipids. Cumulatively, these effects amounted to negligible change in total lipid abundances, but did compromise the observed abundances within specific compound classes. To avoid these effects, environmental samples are frozen immediately after collection and stored at low enough temperatures, typically -80°C, to prohibit heterotrophic remineralization and alteration of organic matter composition prior to analysis.

POC samples intended for bulk elemental analysis are often dried in a heated oven, and samples intended for inorganic analysis are sometimes air-dried, and stored at room temperature before analysis (e.g., Rosengard et al., 2015), all conditions unfavorable to traditional organic analyses (Wakeham and Volkman, 1991). The tremendous labor and financial cost of sampling marine particles across the ocean warrants an answer to how much POC integrity is sacrificed when these distinct drying methods do not meet optimal standards of organic geochemical analysis (i.e., frozen immediately post-collection). Adding to the earlier studies that explored issues associated with storage temperatures of wet samples, this study explores the effects of different sample drying methods on three aspects of organic composition. First, we compare the influence of air-drying samples at room-temperature, oven-drying at elevated temperatures and freeze-drying on bulk organic matter composition (C/N ratio, δ^{13} C and total organic carbon content). Then, we assess the effects of different drying treatments on the biomolecular composition of POC, specifically the abundances of specific lipid compounds in the organic matrix. These comparisons differ from the studies described above, as they relate to the effects of different drying methods, instead

Finally, this report further evaluates the effects of different sample drying methods on thermal stability. Here, we define thermal stability as the distribution of temperatures over which a POC sample decomposes to CO₂ during ramped pyrolysis/oxidation (RPO). RPO is a novel but increasingly utilized technique for characterizing the composition of organic matter in the environment, and has been seldom applied to POC from the marine water column. By pyrolyzing or oxidizing POC throughout a controlled temperature ramp from ~100°C to 800°C, and monitoring the release of CO_2 throughout the ramp, this technique differentiates distinct pools of bulk organic carbon by their thermal stability, alluding to the activation energy of decomposition/oxidation of specific POC pools in the sample (Cramer, 2004). RPO has been applied over a range of organic matter samples from complex depositional environments, from suspended riverine POC to ocean sediments (Rosenheim et al., 2008; Rosenheim and Galy, 2012; Rosenheim et al., 2013; Subt et al., 2016). Observations of increasing radiocarbon age of POC with oxidation temperature in these studies have led to the hypothesis that more degraded, recalcitrant pools of organic carbon are generally more thermally stable, with a higher activation energy associated with thermal decomposition. Thus, RPO offers a unique opportunity to relate POC composition to its biological reactivity in the environment, its fate during transport in the water column and, more relatedly, the strength of the biological pump (Francois et al., 2002; Burd et al., 2016).

We hypothesized that air-drying and oven-drying of a sample post-sampling would negligibly influence the bulk composition of POC (e.g., Kaehler and Pakhomov, 2001). These drying techniques would also impose little changes in thermal stability, as the energy imposed by air-drying and oven-drying relative to freeze-drying is too small to shift the bonding energies (Wagner et al., 1994), and therefore the activation energies of thermal decomposition of the biomolecular matrix (Cramer, 2004). However, we expected that compound-specific abundances would be compromised by the different drying techniques. The results of this report have several implications towards the opportunistic use of marine organic carbon samples for organic analysis, even when not collected and stored according to ideal organic geochemical standards.

2 Methods

2.1 Sample Collection

Samples of marine POC were collected in June 2014 off the dock of the Woods Hole Oceanographic Institution (WHOI), at 41.524°N, 70.672°W. The waters near the WHOI reach bloom-level primary productivity in June.

A battery operated *in situ* pump (McLane WTS-LV) was deployed ~10 meters below surface and pumped seawater through two flow paths (A and B) (Lam et al., 2015). Each flow path directed seawater through a 51 µm pore-size polyester pre-filter screen followed by two pre-combusted 1 µm pore-size quartz fiber filters (WhatmanTM QMA), both mounted onto "mini-MULVFS" filter holders (Bishop et al., 2012). The active collection area of these filters was 125 cm². The pre-filter retained >51 µm size-fraction particles, while the QMAs collected 1-51 µm size-fraction particles, most of which settled on the topmost QMA filter relative to the seawater influx. After ~45 minutes, equivalent to 40-100 L seawater filtered, the filters clogged and the pump stopped. The pump was deployed twice, allowing for collection of three pre-filters, three QMA filters (Fig. 1a, c) and two deployment blank QMA filters (Fig. 1b). The second pump deployment immediately followed the first. Depth control was poor, so the two deployments likely occurred at slightly different depths. Two sample filters (QMA1 and QMA2), one filter per flow path, were collected in the first deployment, while one filter from a single flow (QMA3, flow path B) was collected in the second deployment. One blank filter was collected per deployment by submerging a filter in between two 51 μ m pre-filters in a perforated plastic Tupperware, which was externally attached to the McLane pump frame.

Within hours following collection, each sample and blank QMA filter was divided into thirds, one of which was immediately frozen in a freezer, and then transferred to - 80°C two weeks later (Fig. 1, Table 1). Another third was placed in an oven for drying at 56°C for 17-19 hours. Finally, one third was dried under a clean laminar flow hood at room temperature for 17-19 hours. The filters were apportioned as such to account for environmental heterogeneity associated with particle sampling in POC-rich waters (Sections 2.3, 2.4). All oven-dried and air-dried filters were stored in Whirl-Pak bags or combusted glassware at room temperature before analysis. Frozen filters were freeze-dried for 12 hours before analysis. All blanks and sample particles collected onto the topmost filter relative to the filtration flow path were analyzed within one year of sample collection.

Images of QMA filters were developed at various magnifications for a freezedried portion of deployment blank and sample filters using a scanning electron microscope (Fig. 1b,c). No coccoliths or other calcium carbonate shells were observed in the sample filter. Thus, we assumed that the particulate inorganic carbon content (PIC) in the particle samples was negligible, and would not interfere with any of the POC analyses described in the following sections. The stable isotope data reported in Section 3 will affirm this assumption.

2.2 Bulk Composition

The bulk composition of particles collected onto the QMA filters (particulate organic carbon concentration ([POC]), particulate nitrogen concentration ([PN]), δ^{13} C and δ^{15} N) was analyzed using a *Fisons Instruments Carlo / Erba 1108* elemental analyzer interfaced via a *Finnigan MAT Conflo II* to a *Thermo Finnigan Delta-Plus* stable isotope ratio mass spectrometer (IRMS). Analysis of each drying treatment was conducted on a sub-sample from a different filter replicate prior to combining the thirds from each filter (i.e., the freeze-dried sub-sample came from filter QMA2, the air-dried sub-sample came

from filter QMA3, and the oven-dried sample came from filter QMA1) (Table 1). One deployment blank sub-sample from a freeze-dried filter portion was analyzed. Each subsample represented ~0.9% of the active area of a whole QMA filter (125 cm^2).

2.3 Ramped Oxidation Measurements

The ramped pyrolysis/oxidation (RPO) system at the National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) facility converts sample organic carbon to carbon dioxide (CO₂) gas through a continuous temperature ramp, either by pyrolysis or oxidation. For each drying treatment, the same size subsample across the three QMA filters or two deployment blank filters from the three deployments were combined prior to analysis. When combined, the equivalent to 3-5% of active sample filter area and ~9% of deployment blank filter area were inserted into a quartz reactor inside a furnace programmed to heat at 5°C/minute from room temperature to 1000°C. Furthermore, ~5% of a pre-combusted (to 450°C) but non-deployed QMA filter was also analyzed by ramped oxidation.

RPO analyses were conducted following protocol described in Rosenheim et al. (2008) and Hemingway et al. (accepted). During the temperature ramp, a roughly 35 mL/min. mixture of 92% Ultra-High Purity helium and 8% oxygen gas flowed through the quartz insert from the programmed furnace, carrying oxidation products to an 800°C furnace equipped with a copper, platinum and nickel catalyst that fully converts the products into CO_2 gas. Downstream of the catalyst, the CO_2 and carrier gas passed through a *Sable Systems CA-10* infrared gas analyzer, calibrated at 0 ppm and >400 ppm, which measured the outgoing CO_2 flux and concentration (ppm). After the gas analyzer, the gas mixture flowed into a cryogenic Pyrex coil that was coupled to a vacuum line, where the CO_2 gas was cryogenically trapped, and released into a vacuum line within user-specified temperature intervals. Maintaining consistent flow rates, He: O_2 , furnace catalyst material, furnace insulation, and plumbing minimizes instrumental variation across RPO analyses (e.g., Appendix B, Fig. S3).

For each drying treatment, dock test samples were analyzed three times by ramped oxidation. During each analysis, at least one CO₂ fraction was flame-sealed into

Pyrex tubes in the vacuum line with 50 mg copper oxide and 10 mg silver. For one analysis per drying treatment, three fractions were collected, two to three of which were then baked at 525°C for 1 hour to ensure the full oxidation of any residual incompletely oxidized carbon compounds from the ramped oxidation run, and then analyzed for its stable isotope composition (reported as δ^{13} C) using a *VG Isotech Prism II* IRMS. The precision of δ^{13} C measurements from this IRMS was 0.1 ‰. δ^{13} C values were referenced to Vienna Pee Dee Belemnite (VPDB), using an internal standard calibrated against NBS-19 calcite.

2.4 Compound-Specific Measurements

Compound-specific abundances offer the highest resolution comparisons of organic matter composition across drying treatments in this report. We limit analysis to the abundances of fatty acids, sterols and alcohols, which are commonly applied to study marine POC dynamics in the water column (e.g., Wakeham and Canuel, 1988; Cavagna et al., 2013).

About 5% of the 125 cm² sample QMA filter active area and ~7-15% of the deployment blank filter area were extracted in 15-20 mL of 9:1 dichloromethane: methanol (DCM: MeOH) at 100°C for 20 minutes using a Microwave Accelerated Reaction System (MARS, CEMS Corp.). Each sample extracted was subsampled from the combined sub-fractions of the three sample filter deployments or the two blank filter deployments (Fig. 1). After extraction, the total lipid extract was saponified in 0.5 M potassium hydroxide in MeOH for 2 hours at 70°C. Following saponification, liquid-liquid extractions separated the basic phase from the acidic phase, each of which was flushed through aminopropyl silica gel columns to separate compounds into five compound classes based on their polarity. Fatty acids of both acid and base phases were recombined and methylated for 12 hours at 70°C, and purified through another silica gel column prior to analysis on a flame ionization detector coupled to a *Hewlett Packard 5890 Series II* Gas Chromatograph (GC-FID). Sterols and alcohols of acid and base fractions were acetylated separately in acetic anhydride and pyridine (1:1) for 2 hours at 70°C prior to analysis on the GC-FID.

All sample and deployment blank analyses were accompanied by synthetic standards containing a suite of fatty acids, sterols and alcohols with known concentrations and retention times. Retention times were used to identify specific compounds in the samples and deployment blanks. Other compounds identities, particularly the unsaturated and branched FAMES, were separately validated using an Agilent 7890A gas chromatograph interfaced with a Markes/Almsco BenchTOF-Select time of flight mass spectrometer. To evaluate sample drying effects on compound abundances, we directly compared peak areas across samples and blanks after normalizing the areas to the fraction of QMA filter active area extracted. Peak abundances errors were assumed to be ~10%.

3 Results

3.1 Bulk Composition

Table 1 summarizes the bulk characteristics of particulate organic carbon across sample treatments and one deployment blank (freeze-dried only). All carbon and nitrogen quantities measured on QMA filter sub-samples were normalized to the total active area of the filter 125 cm² and volume filtered, yielding total C and total N concentrations, respectively (Table 1). Filter-normalized total carbon concentrations ranged from 8-14 mg carbon/QMA, corresponding to $11.9 - 16.7 \mu$ M. Total nitrogen concentrations were up to an order of magnitude lower in all samples, 1.8μ M in the freeze-dried sample and 2.5μ M in the oven-dried sample. This corresponded to a small range of molar C:N ratios from 6.5 - 6.8. The bulk δ^{13} C values of sample carbon ranged from -23.9 ‰ in the freeze-dried filter to -23.6 ‰ in the air-dried filter, while δ^{15} N values ranged from 7.9 ‰ in the air-dried POC to 8.3 ‰ in freeze-dried sample.

The total carbon and nitrogen concentrations on the freeze-dried deployment blank filters were ~20 times lower than the content on all sample filters analyzed (Table 1). The scanning electron images of the deployment blank filters show that no particles are loaded onto the filter fibers (Fig. 1b), suggesting that the carbon on the deployment blank filters is likely sorbed dissolved organic carbon. The δ^{13} C of organic carbon in the blank carbon was -25.6 ‰, which was more depleted than the δ^{13} C values of all samples. The C/N ratio in the deployment blank was 7.7, which was higher than in the samples. There was not enough organic nitrogen on the deployment blank filters to resolve a blank carbon δ^{15} N value.

3.2 Ramped Oxidation

All <51 µm particle samples and deployment blanks, oven-dried, air-dried and freeze-dried, were analyzed by ramped oxidation between 100°C and 1000°C. For each drying treatment, sub-samples were analyzed three times, resulting in three thermograms per drying treatment that recorded CO₂ evolved per temperature interval (Fig. 2a-c). When the CO₂ concentrations (thermogram y-axis) are normalized to the active area of a QMA filter (125 cm²), the total CO₂ quantified by the gas analyzer serves as a rough measurement of the total sample carbon integrated across three different filter deployments treated by the same drying process. The deployment blank thermograms. TOC values in the deployment blanks, which were run only once, ranged from 0.28 mg C/filter in the freeze-dried deployment blank to 0.40 mg C/filter in the oven-dried blank (Fig. 2d). These values are two to three times greater than the TOC yielded by analyzing a pre-combusted blank QMA, 0.15 mg/filter (Fig. 2e).

Each sample thermogram was corrected for sorption of carbon onto the filter by subtracting the QMA-normalized deployment blank thermograms from the QMA-normalized thermograms of sample carbon dried by the same process. All blank-corrected and deployment blank thermograms show that most of sample and blank carbon oxidizes between 150°C and 600°C (Fig. 2). The average total filter-normalized carbon across three ramped oxidation analyses of the oven-dried filters is 10.4 ± 0.5 mg/filter (mean ± 1 S.D.). The average across three air-dried filter analyses is 10.8 ± 0.4 mg/filter, and the average across three freeze-dried filter analyses is 10.7 ± 0.5 mg/filter (Fig. 2a-c).

For one analysis per drying treatment, the stable isotope composition of two to three CO_2 gas fractions representing ~95% of all the CO_2 evolved during ramped oxidation of that sample were analyzed within distinct temperature intervals (Section 2.3;

Table 2). In the lowest temperature fraction (~125°C - 415°C), δ^{13} C values ranged from -26.9 ‰ in the oven-dried sample thermogram to -25.8 ‰ in the freeze-dried sample thermogram. In the mid-temperature fraction (~415°C - 620°C), values ranged from -24.4 ‰ in the oven-dried sample to -22.7 ‰ in the freeze-dried sample. The third, highest temperature CO₂ fraction was only analyzed in the oven-dried sample run (610°C -904°C), which had a more depleted δ^{13} C value of -27.8 ‰. The three fraction δ^{13} C values from this sample were normalized to the carbon quantity in each fraction, the sum of which was 2.2-2.5 ‰ more depleted than all bulk δ^{13} C measurements (Section 3.1, Table 1). This difference is similar but slightly greater (0.5 ‰) than the average weighted sum vs. bulk δ^{13} C discrepancy expected from kinetic fractionation during ramped oxidation, which has been observed across several other RPO analyses (n=66) of standard and samples from different environments (Hemingway et al., accepted).

3.3 Lipid Abundances

Peak area-based abundances of fatty acids, sterols and alcohols in particle samples and deployment blanks were quantified across a range of molecular sizes. Fatty acids analyzed include saturated straight-chain fatty acids (12 to 24 carbon chain lengths, or C_{12} - C_{24}), straight-chain unsaturated fatty acids (C_{14} - C_{22}), and saturated branched fatty acids of bacterial origin (C_{15} , C_{16} , C_{17}). Specific alcohols and sterols were identified using a suite of seven standard compounds with different molar masses, carbon chain lengths and environmental origins (Table 3). For each sample and deployment blank chromatogram, areas of select compound peaks in the chromatograms were normalized to the area of an entire QMA filter, representing an average of three replicate filters deployed in Woods Hole waters. Then, filter-normalized peak areas of specific compounds across the deployment blank chromatograms were subtracted from filternormalized compound abundances in samples with the same drying treatment. Comparing just blank-corrected abundances allowed us to control for potential differences between drying effects on sorbed blank carbon and drying effects on sample POC (Figs. 3-5). Indeed, fatty acid abundances in the oven-dried and air-dried blank filters always exceeded abundances in the freeze-dried filters, by up to four times for some compounds.

We then compared ratios of each normalized and blank-corrected peak area in the oven-dried ("O") and air-dried ("A") particle samples to the same normalized peak area in the freeze-dried sample ("F") across chain lengths. For the rest of the discussion, we call these ratios $R_{O/F}$ and $R_{A/F}$, respectively. $R_{O/F}$ and $R_{A/F}$ for saturated straight-chain, unsaturated straight-chain and branched fatty acid peak areas (blank-corrected) in oven-dried or air-dried particle samples relative to peaks areas in freeze-dried samples (blank-corrected) were consistently below 1 (Figs. 3-5), indicating that these compounds are less abundant in the oven and air-dried samples relative to the freeze-dried samples. The $R_{O/F}$ values for all fatty acid compounds analyzed were generally lower than the $R_{A/F}$ values for the same compounds, especially among the straight-chain and branched saturated fatty acids (Figs. 3c, 5a-b).

Two alcohol compounds and no sterols were identified in the deployment blanks: 1-hexadecanol and phytol (Table 3). The blanks make up less than 1% of phytol in the sample peaks, but 18-51% of the *n*-alcohol, with the highest relative proportions in the freeze-dried deployment blank and the lowest proportions in the oven-dried deployment blank. At the same time, the normalized peak areas of these compounds were not significantly different across deployment blanks with different drying treatments. $R_{O/F}$ values were lowest (0.57) for gorgost-5-en-3-ol, as high as 1.7 for 1-hexadecanol, and averaged at 0.98 ± 0.08 across all other compounds (Fig. 6a). Similarly, $R_{A/F}$ values were 0.96 ± 0.17 for most compounds, lowest for gorgost-5-en-3-ol (0.68), and highest for 1hexadecanol (1.3) (Fig. 6b).

4 Discussion

The following compares the bulk, compound-specific and thermal composition of marine POC collected from the coastal waters of Woods Hole, MA and treated by ovendrying, air-drying or freeze-drying. To interpret thermal stability, we assume that biomolecules in an organic matrix span a range of activation energies of decomposition (Cramer, 2004; Rosenheim et al., 2008; Rosenheim and Galy, 2012; Rosenheim et al., 2013), which would drive differences in the oxidation temperature of compounds and fragments of compounds during ramped oxidation. Our interpretations of bulk carbon composition and thermal stability rely on the assumption that particulate inorganic carbon (PIC) concentrations in particulate matter are negligible, and that the carbon quantities reported in section 3 correspond predominantly to organic carbon in the samples. The consistently depleted isotope values of the bulk carbon and CO₂ collected across thermograms (Tables 1, 2) corroborate this assumption. This is further reflected by the lack of any significant calcifying organisms observed on images of particle sample QMA filters (Fig. 1c).

4.1 Oven- and air-drying do not affect bulk characteristics or thermal stability

Bulk δ^{13} C, δ^{15} N and C/N values on QMA filter sub-samples suggest that different drying treatments post-sampling do not appreciably affect bulk POC composition (Table 1). Because the bulk measurements were conducted on individual filter replicates before combination (Section 2.2), environmental variability and heterogeneity of particle loads on the QMA filters may drive the observed differences in the normalized [POC] and [PN] values across filters and deployments, in addition to any effects imposed by the different drying treatments. The fact that the POC concentrations measured in two filters from deployment 1 are more similar to each other than to the concentration measured in deployments likely contributed to the concentration differences. In addition, considering that δ^{13} C, δ^{15} N and C:N values are statistically invariant across filters and drying treatments, air-drying and oven-drying marine particle samples do not change the bulk composition of POC relative to freeze-drying. The standard deviation of the three δ^{13} C (±0.1 ‰), δ^{15} N (±0.2 ‰) and C:N (± 0.2 µmol/µmol) values measured across sample filters are comparable or smaller than to the analytical precision of each measurement.

By comparison, the thermograms generated by ramped oxidation of particle samples and blanks do represent the average of three deployed sample replicates or two deployed blank replicates (Section 2.3), and thus integrate influence of environmental heterogeneity associated with different pump deployments and uneven particle loading (Fig. 3a-c). Thus, any differences across thermograms reflect effects of different drying treatments more so than uneven particle distributions on the filters. The small variations of the area-normalized (125 cm²) and deployment-blank corrected total carbon yields (i.e., total CO₂ evolved) across all nine runs of the three drying treatments (average ± 1 S.D. = 10.6 \pm 0.4 mg), demonstrates that these drying treatments do not influence the total carbon and nature of carbon loaded on the filters. This average carbon/filter quantity calculated from the thermograms is comparable to the three average carbon loadings measured in bulk using the elemental analyzer, 11.9 \pm 3.3 mg/filter (Section 3.1, Table 1). Thus, the thermogram data further support the argument that differences in total sample C concentrations across bulk measurements result from heterogeneity of particle loading among sample filters deployments rather than from the differences in the drying treatments.

The deployment blanks are less susceptible to environmental heterogeneity because the sorption of dissolved carbon compounds to QMA filters is likely to be more homogenous within and between filters. Nonetheless, there are greater relative differences in the total carbon yield among the deployment blank thermograms, which is highest in the oven-dried sample thermogram and lowest in the freeze-dried sample thermogram (Fig. 3d). We posit that these differences result from intrinsically higher relative error associated with analyzing very small quantities by ramped oxidation. Indeed, the non-normalized quantity of CO_2 evolved during ramped oxidation of these samples was small, ranging from 25 µg in the freeze-dried blank thermogram to 37 µg in the oven-dried blank thermogram, i.e., only about ten times the estimated blank contribution for the NOSAMS RPO system (Hemingway et al., accepted).

The differences in thermogram shape across blank-corrected sample thermograms with different drying treatments are comparable to and do not exceed the differences among triplicate thermograms of the same drying treatment (Fig. 2a-c), suggesting that air-drying and oven-drying do not significantly change the thermal stability of the POC relative to freeze-drying (Fig. 2). Use of different programmable ovens could account for some of the smaller differences among the thermogram shapes across triplicate ramped oxidation analyses of the same sample, particularly shifts along the temperature axis. Fig. 2f shows the particle sample thermograms obtained via replicate analyses using the same

oven (referred to as Oven A), which line up better in temperature space than repeated analyses of the same sample using different ovens. In the future, running standard compounds with well-defined peaks, such as sodium bicarbonate, periodically in the different sample ovens would help normalize these temperature shifts in thermograms generated by different ovens (Rosenheim et al., 2008).

In addition, different sample ovens could have driven the 1-2 ‰ differences in the δ^{13} C value of CO₂ gas fractions trapped and analyzed within similar temperature intervals across different sample thermograms (Table 2). Within each of the two lower temperature fractions, CO₂ released by ramped oxidation of the freeze-dried sample was more enriched than CO₂ released by ramped oxidation of the oven-dried and air-dried sample. However, as these gas fractions were collected during analyses on different sample ovens, it is possible that differences in oven insulation and unsteady temperature ramp rates on the earlier end of the temperature ramp affected the thermal evolution of isotopically distinct components of POC in the sample to CO₂.

As hypothesized, the bulk measurements and ramped oxidation data suggest that preservation of marine POC from the field by drying does not compromise the bulk isotope composition, bulk C/N and thermal stability of the sample compared to storage and drying at -80°C. Thus, samples dried at temperatures above -80°C can still be characterized using these metrics.

4.2 Oven- and air-drying shift lipid abundances by compound size and structure

Comparisons of filter-normalized and blank-corrected peak areas, a proxy for compound abundance, across samples show that oven-drying and air-drying shift lipid abundances, and that the magnitude of this shift depends on compound class and size. The results generally agree with prior studies demonstrating that warmer sample storage temperatures compromise specific lipid classes more than others (Ohman, 1996).

Comparisons of saturated straight-chain fatty acids across samples show that $R_{O/F}$ and $R_{A/F}$ values (Section 3.3) correlate positively with the carbon chain length of the compound (p<0.05) (Fig. 3). The slope and r² value of the correlation between $R_{A/F}$ and chain length are higher (Fig. 3b) than corresponding values for the correlation between

 $R_{O/F}$ and chain length (Fig. 3a). This suggests that differences in the volatility of different compounds with different chain lengths and molar masses plays a role in the effects of drying treatment on straight-chain saturated fatty acid abundances (Meylan and Howard, 1991; Lide, 1995; Daubert, 1997; Schwarzenbach et al., 2003). Relatedly, $R_{A/F}$ and $R_{O/F}$ values are similar at lower molecular weights and diverge at higher molecular weights as $R_{A/F}$ values approach 1 more than $R_{O/F}$ values. Thus, drying temperatures can influence the distribution of fatty acids in a sample by driving differential loss of compounds with varying molecular weights.

There is no clear relationship between chain-length and $R_{O/F}$ and $R_{A/F}$ values for unsaturated straight-chain and saturated branched fatty acids quantified across samples (i.e., no significant correlation), as they are relatively invariant and below 1 across chain lengths (Fig. 4-5). Among the straight-chain unsaturated fatty acids, ratios tend to decrease with increasing numbers of double bonds, indicating that compounds with higher degrees of unsaturation are less physically stable and therefore more sensitive to the effects of air-drying and oven-drying (Lide, 1995; Schwarzenbach et al., 2003). Because these branched fatty acids derive from bacteria, the lack of enrichment of the three chain-length branched fatty saturated fatty acids in the oven-dried and air-dried samples relative to the freeze-dried samples implies that the different drying treatments did not enhance microbial degradation of the particles prior to analysis (Ohman, 1996; Wakeham et al., 2002).

Peak areas of seven sterols and alcohols quantified across particle samples are less affected by different drying treatments. $R_{O/F}$ and $R_{A/F}$ values for five of the seven compounds ranged from 0.8 to 1.1 (Fig. 6a-b). Only abundances of gorgost-5-en-3-ol were anomalously depressed (<0.7), while abundances of 1-hexadecanol were anomalously elevated (>1.25) in the oven-dried and air-dried samples. The high abundances of 1-hexadecanol in these oven/air-dried samples relative to the freeze-dried particles can be attributed to proportionally larger sorption blank corrections for this compound in the freeze-dried sample particles. In fact, the ratios of compound abundances between oven-dried, air-dried and freeze-dried samples prior to blank correction are much closer to 1.0 (Fig. 6d-e). Regardless of any biases introduced by different blank corrections, different drying treatments affect alcohol and sterol abundances much less than they affect fatty acid abundances.

The fact that specific lipid abundances in oven- and air-drying shift lipid abundances according to their physical properties (e.g., molecular weight and structure), while bulk characteristics and thermal stability remain unaffected across treatments, supports the argument that sample treatment effects are physical rather than biochemical. The oven- and air-drying may impose enough heat and airflow, respectively, to physically remove some of these light-weight, more volatile molecules. The relatively depressed branched fatty acid abundances in the oven-dried and air-dried samples (Fig. 5) indicates that it was likely that the overnight drying period for the oven-dried and airdried samples was short enough to limit any significant alteration by heterotrophic activity observed, which Ohman (1996) had observed after longer periods of wet sample storage.

Typically, lipids make up at most 25% of total biomass in living microalgae (Finkel et al., 2016), which are the primary contributors to POC in the productive coastal waters of Woods Hole in June. In this study, we have monitored only a small fraction of the marine lipid pool (Wakeham and Volkman, 1991), and have found that some compounds are as much as ~90% depleted in non-freeze-dried samples relative to freeze-dried samples. More than 50% of several of these compounds, however, are still present in oven-dried and air-dried filters. Taken together, such shifts in lipid abundances as a result of different drying treatments must be small relative to the bulk organic matter composition, as bulk δ^{13} C, δ^{15} N, C/N and thermal stability do not vary across drying treatments.

Conclusion

The results of this methods study show that bulk C/N, δ^{13} C, δ^{15} N, and thermal stability are not compromised and can be used as tracers of POC composition and cycling in the water column when particle samples are dried rather than frozen following collection and subsequently freeze-dried. Drying treatments like oven-drying or air-drying do shift the lipid distribution in organic matter, however, and possibly the

distribution of other compound classes. But the magnitude of effects of these drying treatments on compound class abundances are dwarfed by the greater proportion of unaltered material in the bulk carbon pool. More work remains to assess how these compound-specific changes from drying treatments fractionate the stable isotope composition of these biomolecules.

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Table 1. The bulk composition of $<51 \mu m$ particulate organic matter loaded onto three sample and two blank $<51 \mu m$ QMA filters (active area $\sim 125 \text{ cm}^2$), collected during two deployments and then dried in an oven, laminar flow hood or freeze-dried. Filters from each flow path or deployment blank were divided into thirds so that each drying treatment could be applied to each blank or sample filter (Section 2.1, Fig. 1a). Sub-samples from individual thirds of different deployments were analyzed for bulk composition. Sample total carbon and nitrogen contents are not blank-corrected.

Sample	Deployment	Flow Path	Volume (L)	Treatment	C/QMA	[C]	[PN]	C/N	<u>δ13</u> C	δ ¹³ N
			L		mg/QMA	μM	μM	µmol/µmol	‰	‰
QMA1	1	А	95.4	oven-dried						
QMA1	1	А	95.4	air-dried	14	12.4	1.9	6.5	-23.6	7.9
QMA1	1	А	95.4	freeze-dried						
QMA2	1	В	93.8	oven-dried						
QMA2	1	В	93.8	air-dried						
QMA2	1	В	93.8	freeze-dried	13	11.9	1.8	6.8	-23.9	8.3
QMA3	2	В	40.5	oven-dried	8	16.7	2.5	6.8	-23.8	8.2
QMA3	2	В	40.5	air-dried						
QMA3	2	В	40.5	freeze-dried						
db1	1	N/A	0	oven-dried						
db1	1	N/A	0	air-dried						
db1	1	N/A	0	freeze-dried	0.7	N/A	N/A	7.7	-25.6	BDL
db2	2	N/A	0	oven-dried						
db2	2	N/A	0	air-dried						
db2	2	N/A	0	freeze-dried						

BDL= below detection limit.

"db"= deployment blank.

Table 2. δ^{13} C values of CO₂ collected in specific temperature intervals during ramped oxidation of the oven-dried, air-dried and freeze-dried dock test samples. Fraction of total carbon released in each temperature interval was calculated using the gas pressures quantified by a pressure transducer (baratron) in the vacuum line downstream of the RPO system.

Treatment	Oven	Fraction	Temperature Interval	Fraction of Total C	$\delta^{13}C$
			$^{\circ}C$	%	%0
Oven-dried	Α	1	130-412	68%	-26.9
Sample	Α	2	412-610	31%	-24.4
	Α	3	610-904	1%	-27.8
Air-dried	В	1	125-415	63%	-26.8
Sample	В	2	415-617	33%	-23.2
	В	3	617-900	5%	no data
Freeze-dried	В	1	130-416	67%	-25.8
Sample	В	2	416-625	28%	-22.7
	В	3	625-910	5%	no data

Table 3. Constituents of the alcohol/sterol standard and their properties.

Standard compound	Class	Molar Mass	# Carbon	Known sources
		g/mol	#	
Phytol	Alcohol	196.539	20	Chlorophyll
1-hexadecanol	Alcohol	242.447	16	Ubiquitous
5α-Androstan-3β-ol	Sterol	276.464	19	
cholesterol	Sterol	386.664	27	Ubiquitous
5α-cholestan-3β-ol	Sterol	388.68	27	Cholesterol derivative in biological matter Unicellular algae,
Brassicasterol	Sterol	398.675	28	some terrestrial plants
stigmasterol	Sterol	412.702	29	Terrestrial vegetation



Figure 1. (a) Schematic of dock sample collection and post-processing before analysis. A similar approach was used to apportion the deployment blank filters by drying treatment. (b-c) Scanning electron microscope images of freeze-dried (b) blank and (c) sample particle filters. Total filter active area is 125 cm², corresponding to a ~12.6 cm diameter. By comparison, the images below are ~0.6 cm wide.



Figure 2. Blank-corrected thermograms of particle sample filters generated by ramped oxidation (a-c). The furnaces used for the temperature ramp are specified as colored lines for oven A vs. gray lines for oven B. Analyses were conducted three times per sample drying treatment. (d) Thermograms of deployment blanks analyzed once per drying treatment using oven A. (e) Thermogram of a pre-combusted non-deployed QMA filter, also analyzed using oven A. (f) All sample particle thermograms analyzed in oven A. The legend in (d) applies to (f).



Figure 3. (a-c) Ratios of filter-normalized and blank-corrected peak areas of straightchain saturated fatty acids in (a) oven-dried particle samples relative to freeze-dried samples ($R_{O/F}$) and in (b) air-dried samples relative to freeze-dried samples ($R_{A/F}$). These ratios correlate significantly with compound carbon chain length (p<0.05, r²= 0.61-0.81). (c) $R_{O/F}$ and $R_{A/F}$ plotted on the same axes. The stars are ratios of oven-dried (black) or air-dried (blue) blank compound abundances to freeze-dried blank compound abundances. (d) Absolute normalized and blank-corrected peak areas in just the freezedried samples.



Figure 4. (a-b) Ratios of filter-normalized and blank-corrected peak areas of straightchain unsaturated fatty acids in (a) oven-dried particle samples relative to freeze-dried samples ($R_{O/F}$) and in (b) air-dried samples relative to freeze-dried samples ($R_{A/F}$). The numbers on the panels indicate the degrees of unsaturation for each chain length (i.e., number of double bonds). (c) $R_{O/F}$ and $R_{A/F}$ plotted on the same axes. The stars are ratios of oven-dried (black) or air-dried (blue) blank compound abundances to freeze-dried blank compound abundances. (d) Absolute normalized and blank-corrected peak areas in just the freeze-dried samples.



Figure 5. (a-b) Ratios of filter-normalized and blank-corrected peak areas of three branched saturated fatty acids in (a) oven-dried particle samples relative to freeze-dried samples ($R_{O/F}$) and in (b) air-dried samples relative to freeze-dried samples ($R_{A/F}$). The two clusters of data points for chain lengths 15 and 17 represent isomers of the same size molecule. (c) $R_{O/F}$ and $R_{A/F}$ plotted on the same axes. The stars are ratios of oven-dried (black) or air-dried (blue) blank compound abundances to freeze-dried blank compound abundances. (d) Absolute normalized and blank-corrected peak areas in just the freeze-dried samples.



Figure 6. (a-b) Ratios of filter-normalized and blank-corrected peak areas of seven alcohols/sterols in (a) oven-dried particle samples relative to freeze-dried samples ($R_{O/F}$) and in (b) air-dried samples relative to freeze-dried samples ($R_{A/F}$). The x-axis is the compound ID number, corresponding to the legend in panel a. Refer to Table 3 for compound properties. (c) Absolute normalized blank-corrected peak areas of alcohols/sterols in the freeze-dried sample. Quantities in panels d-f are analogous to quantities in a-c, but are not blank-corrected.

Chapter 4:

Using ramped oxidation to track changes in euphotic and mesopelagic zone particulate organic carbon composition

Abstract

The euphotic and mesopelagic zones of the water column are the most dynamic depths intervals for particulate organic carbon composition (POC). Ramped oxidation offers a unique perspective of how POC composition changes in these intervals by decomposing a particle sample over a constant temperature, differentiating POC into components by their thermal stability. Here, we report fifteen thermograms, or plots of CO_2 released against temperature, generated by ramped oxidation of POC collected within three depth profiles (euphotic zone to 1000 m) from three sites of the Southern Ocean Great Calcite Belt region that span a range in primary productivity, phytoplankton community composition and export efficiency. The mixture of organic carbon and calcite in particles at these stations produce sample matrix effects during ramped oxidation that influence the thermal stability distribution of inorganic carbon, which must be accounted for prior to comparing samples by their POC composition alone. The three thermogram profiles demonstrate that the thermal stability range of POC narrows with depth, leading to the accumulation of POC that oxidizes over a mid-temperature range. Further, these transitions occur regardless of location and euphotic zone conditions, supporting the argument for selective preservation of biomolecules in the water column. Changes in the stable isotope values of CO_2 released within specific intervals during ramped oxidation of these samples suggest that different biomolecular pools are selectively preserved across stations, owing to differences in POC source and conversion of POC export to heterotrophic biomass.

1 Introduction

The biological pump moderates the long-term carbon cycle and climate by exporting ca. 16 gigatons/year of sinking particulate organic carbon (POC) from the euphotic zone (Volk and Hoffert, 1985; Eppley and Peterson, 1979; Behrenfeld and Falkowski, 1997; Falkowski et al., 1998; Kwon et al., 2009). The quantity sequestered in the deep ocean comprises $\leq 1\%$ of the carbon fixed by primary production because zooplankton and bacterial heterotrophs rapidly consume and remineralize organic carbon in the euphotic and mesopelagic zones of the ocean (Martin et al., 1987; Buesseler and Boyd, 2009; Giering et al., 2014). At the same time, global compilations of POC flux from the surface ocean to 2000 m (Henson et al., 2011) suggest that the efficiency of the biological pump varies from region to region, as a result of strong latitudinal gradients in both the fraction of primary productivity exported from the euphotic zone and the fraction of POC export transferred to 2000 m. Many interrelated factors have been proposed to drive these variations in POC transport efficiency, from physical processes like advection and the water column temperature gradient (Dall'Olmo and Mork, 2014; Marsay et al., 2015; Laufkötter et al., 2016) to biochemical processes like particle formation and size (Benner et al., 1997; Burd and Jackson, 2009), surface phytoplankton community structure (Francois et al., 2002; Thomalla et al., 2008; Henson et al., 2012a; Henson et al., 2012b), and particle composition (Armstrong et al. 2002; Klaas and Archer, 2002).

This chapter explores the extent to which organic matter quality in particles controls the efficiency of POC transfer in the euphotic and mesopelagic zones of the water column. Defining organic matter quality has challenged marine geochemists for decades, but offers the potential to relate biomolecular reactivity to the biological pump (Burd et al., 2016). Scientific understanding of POC quality has largely evolved with the methods available for characterizing the components of marine POC (Hedges et al., 2000). Yet, different approaches, from bulk carbon isotope measurements to compound-specific biomolecule abundances, and from field observations to controlled culture experiments, have consistently illustrated that certain biomolecular groups accumulate in degraded organic matter, therefore more likely to survive in particles transiting the water column (Wakeham et al., 1997; Hedges et al., 2000; Freeman, 2001). Global differences in POC quality could further bolster the observed relationships between phytoplankton community composition and POC transfer efficiency if different phytoplankton produce and transfer compositionally distinct pools of POC (Francois et al., 2002; Rosengard et al., 2015).

Ramped pyrolysis/oxidation (RPO) offers a novel approach to characterize marine organic matter composition, and an opportunity to differentiate marine POC by its quality or degradability (Rosenheim et al., 2008; Rosenheim and Galy, 2012; Rosenheim et al., 2013). The technique oxidizes or pyrolyzes carbon at a steady, controlled temperature

ramp from 100°C to up to 1000°C, and monitors the release of CO₂, an oxidation/pyrolysis product, throughout the ramp. Plots of CO₂ released against temperature, thereafter referred to as thermograms, separate organic carbon by its thermal stability, or activation energy (Cramer, 2004), offering a perspective of organic matter composition that is finer than bulk measurements, and more quantitative and less laborintensive than compound-specific assessments of POC quality. The reaction can be coupled to a vacuum line to trap and analyze the stable and radiocarbon isotope composition of CO₂ gas produced within specific temperature intervals. Applications of RPO coupled to isotope measurements in the last decade have helped identify and parse diverse carbon sources in different river sediments and complex depositional environments, both marine and terrestrial (e.g., Rosenheim et al., 2008; Rosenheim and Galy, 2012; Subt et al., 2016). These earlier RPO data frequently illustrate increases in radiocarbon age with pools of carbon that oxidize at higher temperatures, supporting the hypothesis that thermochemical stability during pyrolysis/oxidation is a proxy for diagenetic stability in nature (Rosenheim et al., 2008; Rosenheim and Galy, 2012; Rosenheim et al., 2013).

Here, we report and interpret the first set of thermograms generated by ramped oxidation of marine POC from the euphotic and mesopelagic zones of the water column. These POC samples originate from different phytoplankton communities across the Atlantic and Indian sectors of the Southern Ocean, including the Great Calcite Belt, a particulate inorganic carbon (PIC) rich feature that has been attributed to a massive coccolithophore bloom (Balch et al., 2011a; Balch et al., 2016). In this sampling region, the highest diatom abundances occur south of the Polar Front and the highest coccolithophore abundances are found north of the Polar Front in the southwest Atlantic. These changes in community composition are accompanied by variations in carbon transfer regimes through the euphotic zone vary twenty-fold, from 0.97 to 20 mmol/m²/d, representing 4 to 77% of total primary productivity at each respective site. Transfer efficiencies, or the fraction of POC flux transferred to 100 meters below the euphotic zone, ranged from 20% to over 100% in zones with particle production or particle advection below the euphotic zone.

We focus on three depth profiles at stations GB1-117, GB2-43 and GB2-100, which differ in their surface community composition, biomineral composition (i.e., concentrations of biogenic silica, [BSi], and particulate inorganic carbon, [PIC]), POC export efficiency, and the attenuation of POC concentrations in the mesopelagic zone (Fig. 1, Table 1; Rosengard et al., 2015). Thus, analyzing euphotic and mesopelagic zone POC from these three station profiles provides contrasts in primary producer community structure, particle composition, and bulk POC transfer. Coupled to the unique perspective offered by ramped oxidation, we address four questions: (1) What are the compositional differences among POC, in terms of thermal stability and stable carbon isotope ratios, produced by distinct phytoplankton community assemblages in the euphotic zone? (2) How does POC composition change in the mesopelagic zone? (3) How do changes in POC composition with depth relate to the composition of POC produced at the surface? (4) How does thermal stability relate to the diagenetic stability of POC that has been recently produced and thus should exhibit little variation in radiocarbon age?

2 Methods

2.1 Field work

In January – February 2011 and February – March 2012, the *R/V Melville* and *R/V Revelle* traversed the Atlantic and Indian sectors of the Southern Ocean, from Punta Arenas, Chile to Cape Town, South Africa (GB1) and then Durban, South Africa to Perth Australia (GB2), respectively. GB1 and GB2 were timed to coincide with the annual coccolithophore bloom of the austral summer and autumn. Together, both cruise tracks spanned a latitudinal range of 39°S to 59°S, crossing the Subtropical, Subantarctic and Polar Fronts (Fig. 1; Table 1; Balch et al., 2016). Modified dual-flow McLane in-situ pumps (WTS-LV), equipped with 51 µm pore-size polyester pre-filters upstream of paired 0.8 µm polyethersulfone (SuporTM) filters in one flow path, and 51 µm prefilters upstream of paired, pre-combusted 1 µm quartz fiber (WhatmanTM QMA) filters in a second flow path, were deployed at 27 stations across GB1 and GB2 to collect two size classes of particles (>51 µm and 1-51 µm diameter) from the euphotic zone to 1000

meters, as described in Rosengard et al. (2015). All filters had an active particle collection area of 125 cm². Euphotic zone depth, z_{PAR} , was defined as the depth at 0.3% of surface photosynthetically available radiation (Balch et al., 2011b). At each pump cast, complete filter sets sandwiched within a 1 μ m mesh filter were externally attached to the frame of the deepest pump. As these filters passed through each depth of the profile and remained at the deepest depth throughout the entire pump time, they served as deployment blanks to correct for any sorption of dissolved species to the filters during the cast.

2.2 Particle composition

The >51 μ m POC concentrations in all >51 μ m particle samples and deployment blank pre-filters, as well as >51 μ m biomineral (PIC, BSi) concentrations at z_{PAR}, were reported in Rosengard et al. (2015). Analyses focus on the more POC-rich <51 μ m size class from profiles at stations GB1-117, GB2-43 and GB2-100 because resolving stable carbon isotope measurements within fractions of an RPO run requires ≥0.3 μ mol carbon per fraction. At the base of the euphotic zone, the >51 μ m size-fraction particles comprised 14.1 ± 11.6% (mean ± S.D.) of the total particulate organic carbon (Rosengard et al., 2015). This fraction generally decreased towards the deeper mesopelagic zone (800- 1000 m), owing to the longer residence time of <51 μ m particles (months to years) relative to that of >51 μ m particles (days to weeks) in the mesopelagic zone (Riley et al., 2012; McDonnell and Buesseler, 2010).

The <51 μ m BSi, organic carbon and nitrogen concentrations were measured following methods described in Rosengard et al. (2015). <51 μ m [BSi] values were only measured in particles collected onto SuporTM filters at z_{PAR} and in the deployment blank filters at each station. Organic carbon and nitrogen concentrations were measured in QMA filters from 7-8 depths and one deployment blank per station using a *Flash 1112* Elemental Analyzer at Woods Hole Oceanographic Institution (WHOI), following acid fumigation to remove PIC. Only the two shallowest samples from the GB1-117 cast were not fumigated. For these two samples, an independent <51 μ m [PIC] measurement from acid hydrolysis (see below) was subtracted from the total carbon concentration obtained
via elemental analysis (*Flash 1112 EA*) to calculate [POC]. Molar C:N ratios were calculated from these quantities. Attenuation of $<51 \mu m$ [POC] in the mesopelagic zone (Buesseler and Boyd, 2009; Giering et al., 2014) was measured by fitting POC concentration profiles to the following equation (Martin et al., 1987; Lam and Bishop, 2007):

$$[POC]_{z} = [POC]_{zPAR} \left(\frac{z}{z_{PAR}}\right)^{-b}$$
(1)

where z is the sample depth. Profiles with greater remineralization and [POC] attenuation have higher b values, or attenuation coefficients.

Finally, <51 µm PIC concentrations in each profile sample and deployment blanks from GB1-117 and GB2-100 were measured by hydrolyzing QMA sub-samples and deployment blanks in ~1.9 M phosphoric acid in a sealed glass vessel following Galy et al. (2007). After ~16 hours of reaction, well beyond the minimum reaction time observed in a sample time series test (Appendix B, Fig. S1), the CO₂ gas product of hydrolysis was quantified manometrically in a vacuum line. The averaged blank PIC quantities were subtracted from the sample PIC quantities. The CO₂ gas was cryogenically trapped and its stable isotope composition was measured using a *VG Isotech Prism II* isotope ratio mass spectrometer (IRMS). The CO₂ quantities released during hydrolysis of each blank filter were combined prior to δ^{13} C analysis because individual gas quantities from each blank filter were too small for δ^{13} C analysis. All δ^{13} C values in this report are referenced to Vienna Pee Dee Belemnite (VPDB), using an internal standard calibrated against NBS-19 calcite.

Bulk δ^{13} C of sample and blank carbon were also measured by analyzing the total sample CO₂ released from closed tube combustion or elemental analysis, using a *Prism II* IRMS. In closed tube combustion, raw QMA sub-samples were flame sealed in pre-baked evacuated quartz tubes with 2 g of copper oxide and 100 mg of granulated silver at 850°C for six hours (McNichol et al., 1995). During elemental analysis, QMA sub-samples were combusted at 1025°C and quantified as CO₂ using a *Costech Instruments Elemental Combustion System 4010*, before the gas was trapped in a vacuum line and analyzed for $\delta^{13}C_{bulk}$. Section 2.6 describes how the $\delta^{13}C$ value of sample POC was calculated using the measured total sample and blank carbon concentrations, PIC concentrations, $\delta^{13}C_{PIC}$, $\delta^{13}C_{bulk}$ and $\delta^{13}C_{blank}$, as described here. Section 2.6 also compares different ways to measure total [POC] concentration directly and implicitly using the bulk and PIC measurements.

2.3 Sample PIC treatment prior to RPO

Fifteen samples of $<51 \ \mu m$ particles from five depths per three profiles were analyzed by ramped oxidation without removal of PIC from the sample matrix. Only two additional $>51 \ \mu m$ pre-filters from GB2-43 at 20 m and 80 m were selected for comparison with the $<51 \ \mu m$ POC at the same depth because $>51 \ \mu m$ [POC] was high and the $>51 \ \mu m$ [PIC] and the deployment blank [DOC] contributions to total sample carbon were relatively small, <2% (Rosengard et al., 2015). Because pre-filters are made of polyester, the particles were sonicated, flushed off the filters and freeze-dried before being loaded into the RPO reactor. Sub-samples of three deployment blank QMA filters from each profile were also analyzed via ramped oxidation.

The presence of particulate inorganic carbon in particles from the Great Calcite Belt region presents a challenge to interpreting organic matter composition via ramped oxidation, especially if the thermal stability of inorganic carbon overlaps with that of organic carbon. One common approach to separating these two carbon components is to remove the inorganic carbon completely through acid fumigation (Section 2.2). However, residual CaCl₂ in the fumigated sample matrix decomposes during the run, releasing Cl that reacts with the catalyst wire plating in the RPO system (Section 2.5), forming precipitates that obstruct the flow of CO₂ and carrier gas to the gas analyzer downstream. Thus, acid-treated samples must be rinsed in Milli-Q water to remove CaCl₂ before the RPO analysis, potentially altering the composition and mass-balance of the organic carbon pool. A less chemically-disruptive strategy for separating inorganic from organic carbon is to analyze untreated samples in the RPO system, and mathematically subtract the PIC contribution from the raw sample thermogram.

The first approach requires knowledge of whether and how fumigation and rinsing influence the thermal stability of organic carbon, beyond its "intrinsic" thermal stability, by re-arranging organic bonds, affecting the sample matrix, and directly hydrolyzing, dissolving and flushing away components of the organic carbon (Plante et al., 2013). The second approach requires precise knowledge of the thermal stability of inorganic carbon, and is ideal in cases of minimal overlap between the thermal stability of organic carbon and that of inorganic carbon in the sample. We applied both approaches to analyze the marine POC from the Great Calcite Belt region via ramped oxidation. In our discussion, we comment on the applicability of both PIC removal strategies.

To assess the fumigation approach, three $<51 \mu m$ samples of varying PIC concentration (GB2-43 at 25 m, and GB2-100 at 20 m and 800 m) were fumigated for \sim 14.5 hours in 12 N hydrochloric acid vapor, dried for an additional 13 hours at 60°C, and flushed with \sim 75 mL Milli-Q water (Table 1). These filters were then freeze-dried before ramped oxidation analysis. The three sample thermograms were compared to the non-fumigated sample thermograms to evaluate the efficacy of PIC removal relative to the effect of fumigation/rinsing on the thermal stability of organic carbon.

2.4 End-member preparation for RPO

To supplement the no-fumigation approach to PIC removal, we prepared several end-member samples with >93% PIC and 100% POC for RPO analysis. These end-members provide insight into the range and overlap of thermal stabilities between marine PIC and POC.

Two PIC end-members were selected from an Icelandic spar (abiotic, 100% terrestrial calcite) and a Pacific Southern Ocean surface (0-0.5 cm) sediment sample collected by the Joint Global Ocean Flux Study (JGOFS) at 60°S, 170°W (Table 2). The JGOFS sediment contains roughly 62.4% BSi, 31% calcium carbonate, 0.24% POC, and 6.2% lithogenic material by weight, according to prior analyses of average surface sediment composition (0-5 cm) of this core (Sayles et al., 2001). This value was re-affirmed coulometrically following heated acid hydrolysis of the sediment using a *UIC CM140 Total Inorganic Carbon Analyzer* at University of California- Santa Cruz, which yielded a 32% by weight carbonate content in the sediment, equating to a 94%:6% PIC:POC ratio. The $\delta^{13}C_{PIC}$ in the JGOFS sediment was also analyzed following acid hydrolysis in 1.9 M phosphoric acid (Section 2.2). The Icelandic spar was homogenized

and ground before analysis, while the JGOFS sediment, which had been archived at - 80°C, was freeze-dried, homogenized and ground before analysis. Two different quantities of each end-member were analyzed by ramped oxidation to test the sensitivity of PIC thermal stability to the quantity of PIC in the RPO reactor.

Biomass filtered from axenic cultures of *Emiliani huxleyi* served as the POC endmember. 417 mL of f/50 filtered seawater media were autoclaved and then inoculated with two strains of *E. huxleyi*: a non-calcifying strain (CCMP373) and a calcifying strain (DHB624) (Guillard, 1975; Harvey et al., 2015). Cultures were grown in triplicate for 6 days alongside non-inoculated f/50media as a blank batch at 18°C on a 12 hour alternating light/dark cycle in a *Percival Intellus Control System*. Cells were counted immediately after inoculation, and at three additional time points over six days (Appendix B, Fig. S2), using a *Guava easyCtye HT* flow cytometer. On the sixth day, when the cell numbers reached a plateau (i.e., negligible growth rate), the cultures were filtered onto pre-combusted 4.7 cm diameter QMA filters with an active area of ~9.6 cm². Three volumes, 120 mL each, were filtered onto three filters from each replicate of each strain. The filters were freeze-dried before analysis. To verify the assumption that CCMP373 did not calcify, a subsample of a filter from one of the replicates was hydrolyzed in ~1.9 M phosphoric acid in a sealed glass vessel (Section 2.2). A negligible quantity of CO₂ gas was recovered and quantified after ~16 hours of hydrolysis.

2.5 Ramped oxidation

The thermal stability distribution of carbon in each end-member, particle sample (both fumigated and untreated), and sorption blank, was analyzed using the RPO system at the National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) facility at Woods Hole Oceanographic Institution, following Rosenheim et al. (2008) and Hemingway et al. (accepted). For each analysis, sub-samples were loaded into a pre-baked (850°C, 6 hours) quartz reactor within a sample furnace programmed to heat from 100°C to 1000°C at 5°C/minute (Chapter 1, Fig. 1). 35 ± 3 mL/minute of mixed ultrahigh purity helium and oxygen gas ($7 \pm 3 \% O_2$) flowed through the reactor, from the head of the sample furnace, picking up oxidation products generated throughout the

temperature ramp, to the constant 800°C combustion furnace, and then downstream to an infrared gas analyzer and vacuum line. In the constant 800°C combustion oven, which was equipped with copper/platinum/nickel catalyst wire, incomplete oxidation products were oxidized to CO₂ gas. The CO₂ concentration (ppm) in the gas exiting the combustion furnace was measured using a *Sable Systems*© *CA-10* infrared gas analyzer, calibrated at 0 ppm and >400 ppm CO₂, yielding thermograms plotting CO₂ concentration throughout the entire temperature ramp. All RPO analyses in this report were conducted using one of two sample furnaces with similar temperature ramp conditions, which limited instrument-related variation across thermograms (e.g., Appendix B, Fig. S3). The use of two different sample furnaces rather than one, however, was unavoidable and may have introduced some variability in thermograms along the temperature axis.

Downstream of the gas analyzer, the gas mixture flowed into a pyrex coil submerged in liquid nitrogen (N₂), which cryogenically trapped CO₂ while allowing the He and other residual gases to vent. At distinct temperatures, the CO₂ trapped within a user-specified temperature interval was released into a vacuum line for manometric quantification using a pressure transducer (baratron) and cryogenic recollection into a pre-baked, evacuated pyrex tube with ~50 mg of copper oxide and ~10 mg of granulated silver. These tubes were flame-sealed, and baked at 525°C for 1 hour to fully oxidize any incompletely oxidized carbon compounds and scavenge any sulfur- and halogen-containing compounds before re-quantification and stable carbon isotope analysis. Re-quantification of sample fraction mass after the 525°C bake is essential to accurately assess carbon mass balance during the RPO runs because the mass quantified pre-bake includes the non-carbon gas impurities.

2.6 Isotope corrections

The sums of post-bake gas quantities from the untreated $<51 \ \mu m$ POC, $>51 \ \mu m$ POC and deployment blank runs, once normalized to the total filter area, were interpreted as total sample carbon and used to calculate POC content within each filter (Equation 2). In the few cases where the Pyrex tubes were not baked prior to quantification, an average offset between the carbon quantities calculated before and quantities calculated after the 525° C bake was applied to correct the pre-bake quantities before summing all gas fractions. On average, post-bake quantities were 5% lower than the pre-bake CO₂ quantities, but ranged from 39% lower to 9% higher than pre-bake CO₂ quantities. We attribute the relatively lower quantities to excess water and volatile halogen-derived compounds that get cryogenically trapped with CO₂ in the vacuum line coupled to the RPO system, but then stripped away during the additional bake, cryogenic purification and re-quantification step. The relatively elevated quantities could result from products of incomplete oxidation, which still get trapped in the vacuum line, but are not fully oxidized and accurately quantified until after the additional bake step.

Filter area-normalized quantities of PIC in each sample (Section 2.2) and blank carbon, C_{blank} , were subtracted from the total carbon quantified from RPO analyses of each sample, C_{bulk} , as follows:

$$POC = C_{Bulk} - C_{blank} - PIC$$
 (2)

Concentrations of these individual components were subsequently calculated by normalizing the OMA filter quantities to the volume of water filtered. Compared to the other measured sample [POC] values (Section 2.2), [POC] values calculated from Equation 2 are on average 2% greater than concentrations measured by closed tube combustion or elemental analysis on the Costech Instruments Elemental Combustion System, and 12% lower than concentrations measured via elemental analysis on the Flash 1112 Elemental Analyzer (Table 3). The deviations are larger among the deployment blank values, which have relatively low organic carbon loadings, and deeper mesopelagic particles (300-1000 m), where the blank and PIC proportions are high (Table 1). Heterogeneity in the particle load across the OMA filters (Bishop et al., 2012) could also account for such discrepancies. When we discuss POC/filter or concentration quantities in the following sections, we report RPO-based quantities to remain consistent with subsequent calculations that aid in the data interpretation in Section 4. We consider the standard deviations among the two or three POC/QMA values measured for each sample an error for the reported RPO-based quantity that accounts for both instrumental differences and particle heterogeneity upon the filters.

Similarly, the δ^{13} C value of sample POC was calculated by subtracting the PIC and blank contributions to the δ^{13} C_{bulk} values measured via IRMS (*Prism II*) (Section 2.2), using the relative proportions of PIC, POC and sorption blank carbon:

$$\delta^{13}C_{POC} = (\delta^{13}C_{Bulk} - \delta^{13}C_{PIC}f_{PIC} - \delta^{13}C_{blank}f_{C,blank}) \times (\frac{1}{f_{POC}})$$
(3)

In Equation 3, f_{PIC} , f_{blank} and f_{POC} are the fractions of the total carbon that are PIC (e.g., PIC/Total C), blank organic carbon and POC, respectively. To maintain methodological consistency, the RPO-based [OC]_{blank} and [POC] quantities were also used to calculate $f_{OC,blank}$ and f_{POC} Equation 3, respectively. Further, PIC measured separately in the blank filters was subtracted from the total blank carbon measurements to yield $\delta^{13}C_{OC,blank}$ and $f_{OC,blank}$.

Further, the bulk δ^{13} C measurements measured by IRMS, δ^{13} C_{bulk}, were compared to the weighted sums of δ^{13} C values across thermogram gas fractions from each untreated <51 µm and deployment blank run, as well as from one of each PIC end-member run (Table 3). In some sample analyses, where one CO₂ fraction was lost during transfer in the vacuum line or during isotope analysis, the isotope value of the missing gas fraction was calculated by mass balance using bulk δ^{13} C_{bulk} and mass-weighted sum values, δ^{13} C_{sum}:

$$\delta^{13}C_{\text{missing}} = \left(\left[\delta^{13}C_{\text{Bulk}} \times \operatorname{avg}\left(\frac{\delta^{13}C_{\text{sum}}}{\delta^{13}C_{\text{bulk}}}\right) \right] - \sum_{i}^{i} f_{i} \delta^{13}C_{i} \right) \times \left(\frac{1}{f_{\text{missing}}}\right)$$
(4)

Here, *i* is the CO₂ gas fraction, f_i and $f_{missing}$ are the relative mass proportions of CO₂ in each measured and lost fraction, respectively. The second term in the first bracket represents the average of all $\delta^{13}C_{bulk}$ and $\delta^{13}C_{sum}$ values for sample thermograms for which no fractions were lost.

2.7 Error propagation

Errors associated with carbon quantities and δ^{13} C values of different components of sample carbon derive from various instrumental errors, heterogeneity of particle loads onto QMA filters, and blank or PIC corrections. In the results, the uncertainty in the blank subtraction stems from the standard deviation of blank carbon quantities measured by RPO and two elemental analyzers (Section 2.6). The uncertainty in the PIC correction derives from the standard deviation of the different deployment blank PIC quantities. The error of all directly measured δ^{13} C values is equal to the precision of the IRMS, 0.1‰ for the *Prism II*. The carbon quantities in thermogram fractions have a relative error of ~1%, which is minimal compared to the standard deviation of carbon quantities calculated across the replicate POC measurements for each sample. These different errors are propagated for any calculated concentrations or δ^{13} C values reported in the results and discussion.

3 Results

3.1 Bulk composition in marine samples

Blank-corrected PIC, POC and BSi concentrations measured in <51 µm particle samples were measured in three profiles from the Great Calcite Belt region, GB1-117, GB2-43 and GB2-100 (Table 1). Acid hydrolysis-based [PIC] values range from 0.01 µM at 110 m at station GB2-100 to 0.18 µM in the shallowest sample at GB1-117. BSi concentrations at or near the export depth z_{PAR} of each station range from 0.0074 µM at GB2-100 to 0.17 µM at GB2-43. POC concentrations, calculated from the RPO runs (Equation 2; Section 2.6), range from 0.11 µM in the deepest mesopelagic zone sample at GB2-100 to 5.1 µM in the shallowest euphotic zone sample at GB1-117. The *Flash 1112* EA-based [POC] values, measured at more mesopelagic zone depths than the other [POC] quantification methods, were fit to a power law (Section 2.2, Equation 1) which yielded attenuation coefficients from 0.8 at GB2-43 and 1.1 at GB1-117.

[PIC]/[BSi] ratios at or near z_{PAR} are highest at GB1-117 (8.2) and lowest at GB2-43 (0.1-0.2), in agreement with the relative [PIC]/[BSi] ratios observed in the >51 µm size-fraction particles collected at these three stations (Table 1; Rosengard et al., 2015).

Following Rosengard et al. (2015), we use [PIC]/[BSi] ratios in both size-fractions as an estimate of the relative coccolithophore and diatom abundances at stations GB1-117, GB2-43, and GB2-100.

At all stations, [POC] and [total C] from the untreated, uncorrected ramped oxidation analyses decrease with depth while the proportion of blank carbon and PIC in the total carbon pool increases with depth (Table 1). The quantities of dissolved organic carbon sorbed onto the deployment blank filters, range from 35 µmol to 57 µmol per filter (Table 1). At station GB1-117, the most PIC-rich station, PIC comprises 3-9% and blank POC comprises 3-28% of the total carbon. At station GB2-43, the most BSi-rich and PIC-poor station, PIC comprises 1-4% of the total carbon, while blank POC comprises 9-47% of the total organic carbon. At station GB2-100, which is also PIC-rich, PIC contributes to 2-14% of the total sample carbon, and blank carbon comprises 4-33% of the total organic carbon.

 $\delta^{13}C_{POC}$ values, calculated by Equation 3, are overall most ¹³C-enriched at GB1-117 and most depleted at GB2-43 (Table 1, Fig. 2a-c). They are relatively constant with depth at GB1-117 and GB2-100, but increase from -26.9‰ in the euphotic zone to -24.1‰ in the mesopelagic zone at GB2-43. At all stations, the $\delta^{13}C$ values of the deployment blanks are ≥5‰ lighter than $\delta^{13}C_{POC}$ measured in the profile samples.

Blank-corrected $\delta^{13}C_{PIC}$ values in each sample are very ¹³C-enriched, varying little between -2 and 1.5‰ across stations and depths (Table 1, Fig. 2d-f). The $\delta^{13}C_{PIC}$ averaged across the three deployment blanks is more enriched than the samples (2.9‰). The range in sample $\delta^{13}C_{PIC}$ is comparable to the variation observed in Southern Ocean DIC (Trull and Armand, 2001), in PIC from marine calcifiers (Keigwin et al., 2005), and in the PIC present in core-top sediments from the Pacific sector of the Southern Ocean (Section 3.2).

The following sections expand upon these bulk observations with thermograms generated by oxidizing several sets of samples from 100°C to 800°C: (1) three fumigated samples of <51 μ m POC from the Southern Ocean Great Calcite Belt; (2) 95-100% pure inorganic carbon, terrestrial and marine; (3) pure organic carbon from axenic coccolithophore cultures; and (4) fifteen untreated samples of <51 μ m POC from the Great Calcite Belt. We describe three aspects of thermogram shape across distinct

temperature intervals: (1) the fraction of total CO_2 released, (2) the isotopic composition of CO_2 gas released, and (3) T_{max} within these intervals, or the temperatures of peak maxima.

3.2 Acid fuming results

With significant PIC content in the Great Calcite Belt particles, we explored two methods of correcting for the PIC contribution to total sample carbon in thermograms (Section 2.3). The first method removes PIC prior to ramped oxidation via acid fuming, while the second accounts for PIC by subtracting it from a thermogram of an untreated sample.

To test the first method, we compared three acid-fumigated (and rinsed) and untreated thermograms of three <51 μ m POC samples: one 85 m depth sample from station GB2-43, one 90 m sample from station GB2-100, and one 800 m sample from GB2-100. Comparing thermograms of acidified and untreated 800 m particles, where PIC comprised 14% of the total carbon before acid treatment (Table 1), provided an assessment of the combined effects of fumigation on POC thermal stability and PIC removal in the sample (Fig. 3a). By contrast, comparisons of fumigated and untreated sample thermograms from surface depths, where PIC represents <5% of the total carbon, provided a control for evaluating how fumigation and rinsing influence the thermal stability of particulate carbon dominated by POC alone (Figs. 3b,c).

The data show that the fumigation and rinsing affect PIC-rich sample thermograms just as much as they affect thermograms of PIC-poor samples (Fig. 3). But, the treatment affects the two surface samples differently than it affects the 800 m sample in terms of mass balance and loss. For all samples, the quantity of carbon released from ramped oxidation after fumigation and rinsing decreased by 22-34% in the surface samples and 26% in the deep sample. This difference in carbon yield exceeds the expected heterogeneity in POC loading on QMA filters (Bishop et al., 2012), and is likely a result of carbon lost by fumigation and rinsing. Considering that no more than 10% of the carbon in the surface samples is PIC and sorption blank contribution, the additional carbon lost must be POC mobilized and/or solubilized by the fumigation and rinsing process (Table 1, Figs. 3a-b). These results do not preclude the possibility that the acid treatment and rinsing further shifted the thermal stability of organic carbon in the samples, which also could drive changes in thermogram shape observed in these panels.

By contrast, the carbon lost after treatment and rinsing of the 800 m sample exceeds the PIC content in the sample (14% of total carbon) but not the combined PIC and blank organic carbon content (47% of total carbon) (Table 1, Fig. 3c). Thus, assuming that all the PIC was hydrolyzed, either some but not all of the blank carbon, or a small combination of both POC and blank carbon was removed in the treatment process. Again, it is still possible that fumigation and rinsing could have further shifted the thermal stability of POC in the thermogram.

From these comparisons and our inability to tease apart the effects of fumigation and rinsing on (1) carbon mass balance (POC, blank organic carbon and PIC) and (2) the thermal stability of POC in samples with different PIC:POC, we conclude that such treatments introduce significant methodological artifacts to the thermal stability of POC in a sample. Thus, the remaining discussion focuses on thermograms generated only from untreated samples.

3.3 End-members

Having ruled out acid fumigation, we must consider the approach of subtracting the PIC component from a specified temperature range from each Great Calcite Belt sample thermogram. To improve expectations of these different oxidation temperature ranges for marine POC and PIC, thermograms of PIC and POC end-member samples were generated to assess the overlap between PIC and POC thermal stability. The pure end-member samples analyzed here are a terrestrial CaCO₃ (calcite), marine biogenic CaCO₃, and fresh marine organic carbon (i.e., pure phytoplankton biomass) (Table 2).

The first PIC end-member is an Icelandic spar, pure terrestrial calcite. Two masses of homogenized grains from this sample, equivalent to ~450 μ g and ~250 μ g total inorganic carbon, were analyzed through ramped oxidation to assess whether there is an amount effect on its thermal stability. Both thermograms from these analyses display one peak, skewed towards lower temperatures (Fig. 4a). The T_{max} is constant across both

thermograms, suggesting that the thermal stability of the PIC does not depend on the mass of inorganic carbon in the reactor. δ^{13} C values in this peak, measured in CO₂ gas fractions of the 450 µg analysis, are ¹³C-enriched and relatively invariant, varying between 1 and 3‰ (Fig. 4b), indicating small kinetic isotope fractionation of carbon oxidized during the temperature ramp (Hemingway et al., accepted). The weighted sum of these isotope measurements is nearly identical to the separately measured bulk value of 3.0‰ (Table 3).

The second PIC end-member came from a Southern Ocean core top sediment at ~5 km water depth (60°S, 170°W), collected during the U.S. JGOFS sampling campaign (Sayles et al., 2001). ~94% of the total carbon in this sediment is PIC, according to Sayles et al. (2001) and independent coulometry measurements of inorganic carbon content in the lab. Two quantities, equivalent to ~950 μ g and ~250 μ g total carbon, were analyzed by ramped oxidation. Both thermograms exhibit one major negatively skewed peak (Fig. 4c), the T_{max} of which shifts from 651°C in the 950 μ g C analysis to 630°C in the 250 μ g C thermogram. This shift suggests that the thermal stability and thus the kinetics of PIC decomposition in the JGOFS sediment depends on the carbon quantity available for oxidation and follow a higher than first order kinetic rate.

Compared to the Icelandic spar, the PIC present in this marine sediment oxidizes over a broader distribution of temperatures. Stable carbon isotope measurements in the higher quantity analysis of the JGOFS sediment show that the CO₂ evolving between 436°C and 721°C is similar to the hydrolysis-based δ^{13} C of the PIC in this sample (0.7‰) and thus originates from the PIC component in the sample (Fig. 4d, Table 2). CO₂ in the first fraction, only 4% of the total carbon evolved during ramped oxidation, is comparatively ¹³C-depleted (-20.4‰) and likely derives entirely from organic carbon in the sample. Between 364°C and 436°C, the CO₂ likely derives from a mixture of PIC and organic carbon, yielding an intermediate δ^{13} C value of -10‰. The weighted sum bulk δ^{13} C value from the JGOFS thermogram fractions (-1.1‰) is statistically identical to the independently measured JGOFS δ^{13} C value (-0.9‰) (Table 3).

Figure 5 shows four thermograms of *E. huxleyi* culture biomass, each generated by oxidizing a filter sub-sample from one culture replicate of either the DHB624 or CCMP373 cultures. In general, the culture POC oxidizes from 150 to 600°C, spanning a

wider range of thermal stabilities than the range of thermal stabilities of Icelandic spar and JGOFS sediment-derived PIC. Smaller-scale variations in thermogram shape across sample runs indicate that differences in the distribution of thermal stabilities in biomass from replicates of the same strain (i.e., CCMP373) are comparable to the differences observed across strains (i.e., CCMP373 vs. DHB624). The hydrolysis-derived PIC concentration measurement in one of the strain CCMP373 filters shows that less than 0.5% of the total carbon on the filters is PIC (Table 2; Section 2.4), affirming that the variations across each replicate thermogram reflect differences in POC composition rather than mixing of POC and PIC. Further, strain DHB624 *E. huxleyi*, which can create calcite liths, did not appear to calcify in culture prior to filtration, which is supported by similar light δ^{13} C values of CO₂ fractions across thermograms of both strains (-27.7‰ to -18.5‰). The δ^{13} C of CO₂ fractions from the strain CCMP373 replicate thermograms were not significantly different from isotope values measured within fractions from the one strain DHB624 thermogram (p>0.05, Student's t-test).

Sub-samples from the same *E. huxleyi* strain and culture replicate were not analyzed more than once due to sample limitations. However, RPO analyses of two different masses of the same POC-rich Great Calcite Belt sample (Appendix B, Fig. S3) suggest that the reaction kinetics of organic carbon is mass-independent. Most importantly, Fig. 5 demonstrates that while all of the organic carbon from the culture biomass is less thermally stable than the inorganic carbon from the Iceland spar (i.e., no temperature overlap in the thermograms), there is significant overlap between the temperature range for thermal decomposition of PIC from the JGOFS sediment (Fig. 4c,d) and for oxidation of POC in the culture biomass. In Section 4.2, we discuss the impact of this overlap on interpreting thermograms of samples that comprise a mixture of marine biogenic POC and PIC.

3.4 Blank-corrected thermograms

Fifteen thermograms from three stations and five depths per station were generated by analyzing untreated sub-samples of QMA filter particles through ramped oxidation (Fig. 6). In addition, two sub-samples of $>51 \mu m$ POC collected at 20 m and 85 m from station GB2-43, and three sub-samples of the deployment blank QMA filter from each station profile were similarly analyzed (Fig. 7). In all twenty thermograms, most of the total sample carbon – a mixture of POC and PIC – oxidizes from 200°C to 600°C, similar to the oxidation temperature range observed in the *E. huxleyi* culture thermograms (Fig. 5). At least two prominent peaks are present in thermograms of all samples.

The blank thermograms were directly subtracted from the $<51 \mu m$ POC thermograms to yield thermograms of total suspended particulate carbon (i.e., sample PIC and POC). The $>51 \mu m$ POC thermograms were not blank-corrected because the DOC sorption at the two chosen depths of GB2-43 contribute at most 1% to the total sample carbon (Rosengard et al., 2015).

3.5 δ^{13} C distribution across thermograms

On average, the weighted sums of all δ^{13} C values measured from a thermogram were ~1.2‰ lighter than the independently measured bulk isotopic composition (Table 3). This offset is similar in magnitude to other weighted sum vs. bulk comparisons compiled in Hemingway et al. (accepted), but greater than the offset between the δ^{13} C sums from the PIC end-member thermograms and their bulk δ^{13} C values.

 $δ^{13}$ C values measured in CO₂ trapped throughout the ramped oxidation of the untreated <51 µm POC samples range from -33.1‰ to -17.4‰ (Fig. 8). The presence of ¹³C-enriched PIC (Table 1) and ¹³C-depleted sorbed organic carbon, which ranged from -35.5 to -26.9‰ across the deployment blank thermogram fractions (Fig. 7), contributes to this wide range, especially for the deeper samples, where $f_{OC,blank}$ and f_{PIC} are high (Table 1). But, the average $δ^{13}$ C range of CO₂ fractions from the surface samples, where the combined PIC and blank contributions represent <10% of the total sample carbon, is comparable to the $δ^{13}$ C range across fractions from the *E. huxleyi* thermograms (Fig. 5), and must result from biomolecular differences in the CO₂ evolved through temperature. Isotopes measured across CO₂ fractions from one of the >51 µm POC thermograms (GB2-43, 85 m) are much lighter than the isotopes in the corresponding <51 µm particles, ranging from -35.7‰ to -30‰.

4 Discussion

In the following section, we use two metrics to compare RPO data: thermal stability and isotope composition. We considered comparisons across two spatial dimensions: (1) a vertical dimension that provides a gradient of degradation state within each profile, and a (2) geographic dimension that provides a gradient in POC export efficiency and phytoplankton community composition.

Thermal stability can be inferred from thermogram shape and the distribution of temperatures across which distinct carbon pools oxidize to CO₂. We assume that the spectrum of biomolecules in complex organic mixtures such as marine particles span a range of activation energies of decomposition, causing different biomolecules and subgroups within the larger, more structurally complex biomolecules to oxidize to CO₂ throughout the continuous temperature ramp imposed by RPO analysis (Burnham and Braun, 1999; Cramer, 2004; Rosenheim et al., 2008; Hemingway, 2017 thesis). Thus, CO₂ that evolves at higher temperatures originates from more thermally stable biomolecules that require more energy to decompose. Conversely, CO₂ that evolves at a lower temperature originates from less thermally stable biomolecules with weaker bonds, i.e., lower activation energies. We evaluate the strength of thermal stability and activation energy as a proxy for diagenetic stability, and probe the geochemical information gleaned from coupling thermal stability to stable carbon isotope measurements.

Because the following RPO analyses are limited to $<51 \mu m$ particles from the Great Calcite Belt region, commonly thought to represent the composition of suspended POC rather than that of sinking POC (e.g., Rosengard et al., 2015). In the euphotic zone, marine particles primarily comprise relatively fresh cells that are still living or were recently alive. By comparison, POC in the mesopelagic zone represents more degraded material, with further differences expected between the $<51 \mu m$ and $>51 \mu m$ size fractions because they reside in the mesopelagic zone over significantly different time scales (Sheridan et al. 2002). Assuming that $>51 \mu m$ particles sink 20-200 meters/day (McDonnell and Buesseler, 2010), a $>51 \mu m$ size-fraction is composed of very slowly sinking or suspended small particles that originate from disaggregating large, fast-sinking

particles that can also be subject to re-aggregation (Riley et al., 2012; Lam and Marchal, 2015). Studies using long-lived thorium isotopes and organic biomarkers have shown that aggregation of small to large particles and disaggregation of large to small particles occur throughout the water column (Murnane et al., 1994), but that the level of particle exchange can vary by ecosystem type and season (Abramson et al., 2010).

The mesopelagic <51 μ m size fraction thus integrates POC pools that experience longer and more widely ranging residence times, from a few months to a few years, controlled by the rate of respiration, sinking and mass transfer with larger size-fraction particles (Marchal and Lam, 2012; Lam and Marchal, 2015). Because remineralization in the mesopelagic zone is predominantly driven by bacteria, the cumulative conversion of POC exported from the euphotic zone to heterotrophic biomass may further drive differences between <51 μ m and >51 μ m POC composition (Griffith et al., 2012; Cavagna et al., 2013). The choice to focus analyses on the longer residence time size fraction provides a strong gradient in degradation for evaluating the compositional changes in marine POC through the upper 1000 m of the water column. For the purposes of this report, differences between euphotic zone and mesopelagic zone <51 μ m POC presumably reflect differences in organic matter source and degradation state of organic matter.

4.1 Latitudinal isotopic depletion in Southern Ocean POC

Overall, the calculated $\delta^{13}C_{POC}$ values from station profiles GB1-117, GB2-43 and GB2-100 fall within the range of values observed in other water column particles from the Southern Ocean (Trull and Armand, 2001; Cavagna et al., 2013). The isotopic depletion across the three stations follows the latitudinal gradient observed in other Southern Ocean studies, agreeing well with the relationship between the CO₂ supply and kinetic fractionation associated with photosynthesis. In the Southern Ocean, as euphotic zone waters grow colder and more $CO_{2(aq)}$ -replete with latitude, carbon fixation by phytoplankton discriminates more in favor of ¹²C (Goericke et al., 1994; Popp et al., 1999). In the Great Calcite Belt samples, we see euphotic zone water temperatures

decrease from ~15-20°C to ~6°C (Table 1) from GB1-117 to GB2-43 while surface $\delta^{13}C_{POC}$ values decrease from ~-20‰ to ~-25‰.

The small magnitude of variation in δ^{13} C values of <51 µm POC with depth in these samples is also consistent with other profiles (Cavagna et al., 2013), and supports the notion of non-selective remineralization in POC with depth. However, as the following sections elaborate, the thermogram profiles reveal greater variations in POC composition with depth that are not resolved by these bulk $\delta^{13}C_{POC}$ profiles.

4.2 Sample matrix effects on thermal stability

A major challenge to interpreting the Great Calcite Belt thermograms (Fig. 6) is the presence of PIC in the particles, especially in those from the mesopelagic zone. Considering that the inorganic carbon in both terrestrial and marine calcite thermally decompose at temperatures above 600°C (Fig. 4), it is surprising that little to no CO₂ evolves in any of the Great Calcite Belt sample thermograms above 600°C. We hypothesize that matrix effects in these marine particle samples have lowered the activation energy required to decompose PIC, causing this component to decompose at lower temperatures than expected from Fig. 4. Prior studies have demonstrated matrix effects on sample thermal stability during ramped pyrolysis by showing that thermograms of simple organic compound mixtures deviated from thermograms produced by pyrolyzing the pure compounds separately (Williams et al., 2014). To test the hypothesis that matrix effects also shift PIC thermal stability, five artificial sample matrices were prepared by mixing sub-samples of E. huxleyi culture replicates with one of the two PIC end-members at various proportions (Table 2), and analyzed by ramped oxidation. Temperature ramp conditions and the programmable sample oven were kept constant across analyses (Rosenheim et al., 2008; Hemingway et al., accepted). Only strain CCMP373 filters were used in matrix mixtures because they had negligible PIC concentrations and served as an organic end-member (Section 2.4).

Matrix mixtures I and II were composed of roughly equal proportions of POC from the culture sub-samples and terrestrial or marine PIC. Matrices III, IV and V consisted of much higher proportions of organic carbon relative to inorganic carbon.

Matrices IV and V had essentially the same proportions of culture-derived POC and JGOFS sediment-derived PIC. However, matrix V contained about two times the amount of total carbon than did matrix IV, which enabled us to test for possible effects of massdependent oxidation kinetics on thermogram shape. The PIC quantities were approximated using the known weight % PIC quantity in the Icelandic spar and JGOFS sediment samples (Section 2.4). The E. huxleyi quantities were approximated using the total carbon recovered per unit filter area from individual RPO runs of each culture triplicate (Section 3.3, Fig. 5). Using these C vields, ~24-82% filter active area (~9.6 cm^2) was sub-sampled for each matrix mixture. These matrix proportions prepared before RPO analysis were estimates because either small masses of PIC from very PIC-rich samples are difficult to weigh, or the carbon loading on the E. huxlevi culture filters was uneven and therefore the carbon content was difficult to estimate based on filter area. During ramped oxidation, five fractions of CO_2 gas were trapped at select temperature intervals and analyzed for δ^{13} C composition. The actual proportions of the two endmembers in each mixture, $f_{E,hux,Total}$ and $f_{Calcite,Total}$, were determined after ramped oxidation by isotope mass balance among the mass-weighted sums of δ^{13} C measured across the matrix thermograms (Fig. 9) and the mass-weighted sums of δ^{13} C measured across pure end-member thermograms (Table 3):

 $\delta^{13}C_{matrix,Sum} = \delta^{13}C_{E.hux,Sum}f_{E.hux,Total} + \delta^{13}C_{Calcite,Sum}f_{Calcite,Total}$ (6) The values of $f_{E.hux,Total}$ and $f_{Calcite,Total}$ are reported in Table 2. For matrix mixtures II, IV and V, we assume that the organic component from the JGOFS sample contributes negligibly to the isotope mass balance in Equation 6, as it comprises only 6% of the total carbon in the sediment (Sayles et al., 2001).

To assess the presence of a matrix effect, we compared the expected thermograms that would result from the null hypothesis, a simple linear combination of the endmember thermograms, against measured thermograms from each of the five mixtures (Fig. 9). These "null hypothesis" thermograms were calculated by adding end-member thermograms (Figs. 4, 5) adjusted by their calculated proportions in each matrix sample (Table 2, Equation 6). Because the pure JGOFS thermograms vary by the mass loaded into the RPO reactor (Section 3.3), null thermograms were constructed from the lower quantity RPO analyses in Fig. 4, as similarly low PIC quantities were present in mixtures I-V (Table 2). Deviations of the measured from the reference/null thermograms indicate the presence of a matrix effect.

We further compared the actual contribution of PIC (calcite) and POC (culture)derived CO₂ to each gas fraction of the matrix thermograms with the expected contributions of PIC- and POC-derived CO₂ to each trapped gas fraction from the null hypothesis case. Expected CO₂ from PIC in each gas fraction, $mass_{Calcite,null,i}$ was calculated as:

$$mass_{Calcite,null,i} = \int_{Ti0}^{Tif} Calcite_{norm} dt \times f_{Calcite,Total}$$
(7a)

where the first term is the integral of the PIC end-member thermogram, normalized to total CO₂ evolved, between the start (T_{i0}) to the end (T_{if}) temperatures for fraction *i*. This term was then scaled by the proportion of PIC in the matrix (Table 2). Similarly, expected CO₂ in each gas fraction, $mass_{E,hux,mull,i}$, was calculated as:

$$mass_{E,hux,null,i} = \int_{Ti0}^{Tif} E. hux_{norm} dt \times f_{E,hux,Total}$$
(7b)

where the *E*. *huxleyi* end-member thermogram, normalized to total CO₂, was integrated from the start (T_{i0}) to the end (T_{ij}) temperatures of each fraction *i* and then scaled by the proportion of culture-derived POC in the matrix (Table 2).

The calculated $mass_{Calcite,null,i}$ and $mass_{E,hux,,null,i}$ values were compared to observed quantities of PIC in each matrix fraction, which were constrained by the matrix gas fraction δ^{13} C measurements, or δ^{13} C_{matrix,i}, using the following relationship:

$$\delta^{13}C_{\text{matrix},i} = f_{\text{Calcite},obs,i}\delta^{13}C_{\text{Calcite},i} + f_{\text{E}.\text{hux},obs,i}\delta^{13}C_{\text{E}.\text{hux},i}$$
(8a)

The end-member δ^{13} C values for each gas fraction, δ^{13} C_{Calcite, i} and δ^{13} C_{E.hux.,i}, were measured in CO₂ released within similar temperature intervals during ramped oxidation of each pure end-member (Figs. 4-5). Substituting $f_{E.hux.,i}$ with 1- $f_{calcite,obs,i}$, we solve for the fraction of CO₂ within gas fraction *i* that derives from PIC decomposition:

$$f_{Calcite,obs,i} = \frac{\delta^{13}C_{matrix,i} - \delta^{13}C_{E.hux,i}}{\delta^{13}C_{Calcite,matrix,i} - \delta^{13}C_{E.hux,i}}$$
(8b)

Thus, the observed PIC-derived CO₂ quantity in each gas fraction, $mass_{Calcite,obs,i}$, was then calculated as:

$$mass_{Calcite,obs,i} = C_i \times f_{Calcite,obs,i}$$
(9a)

where C_i is the total CO₂ released within each gas fraction *i* of the matrix thermogram. Similarly, the measured CO₂ from POC for each gas fraction, $mass_{POC,meas,i}$ was calculated as:

$$mass_{E.hux,obs,i} = C_i \times (1 - f_{Calcite,obs,i})$$
(9b)

The Matrix I thermogram, composed of 50% organic carbon and 50% inorganic carbon from the Icelandic spar, differs from the null hypothesis thermogram (Fig. 9a). The most ¹³C-enriched ($\delta^{13}C_{matrix,i} = 3\%$) peak in the thermogram, thus derived purely from PIC, has a T_{max} of 680°C. This Tmax is 14°C cooler than would be expected in the absence of matrix effects, indicating that the sample matrix has lowered the thermal stability of the Icelandic spar in this mixture. Further, the $\delta^{13}C$ enrichment observed in fractions 3 (427-529°C) and 4 (529-599°C) of matrix I relative to the depleted $\delta^{13}C$ value of CO₂ trapped at similar intervals from the *E. huxleyi* thermograms can only be explained by the presence of PIC in these fractions. The quantity of PIC required to produce this ¹³C-enrichment in these two intervals is greater than the theoretical quantity of PIC expected from the linear addition of PIC and POC from each end-member, respectively (Fig. 10a). This explains why less PIC than expected in the null hypothesis thermally decomposes above 600°C.

Matrix III was constructed from a greater proportion of culture-derived POC relative to Icelandic spar PIC (Fig. 9c). With one isotope measurement missing between 531 and 599°C, it is not possible to use Equation 6 to constrain the actual proportions of inorganic and organic carbon in the matrix. But, assuming 15% inorganic carbon and 85% organic carbon, which were the target proportions during sample preparation, there are significant differences in observed vs. null thermogram shape. CO₂ released above 600°C is mostly PIC, with an enriched δ^{13} C value (1.5‰), but again peaks 14°C cooler than the peak in the pure PIC thermogram. The smaller relative area of total CO₂ in this fraction and greater relative area of CO₂ between 525°C and 600°C, compared to the null case, indicates that the matrix effect has shifted the thermal stability of PIC towards a lower temperature range, as well (Fig. 10c).

The carbon in matrix II, 59% organic culture biomass and 41% PIC from the JGOFS sediment, mostly oxidizes between 200°C and 600°C (Fig. 9b). The thermogram displays two prominent peaks at 500°C and 610°C. Again, the results differ from what

would be expected had the thermogram reflected just a linear sum of the pure endmember thermograms. The most ¹³C-enriched peak, mostly derived from PIC, with a T_{max} of 610°C, must correspond to the highest CO₂ concentration peak in the JGOFS thermogram, which also has a T_{max} of 610°C. At the same time, it is clear that the amount of CO₂ that evolves above 600°C in matrix II is lower than the amount released in the null hypothesis case (Figs. 10b). The isotopic enrichment of CO₂ across the matrix II thermogram relative to the CO₂ analyzed the *E. huxleyi* thermograms shows that this difference is compensated by more inorganic carbon decomposing between 426°C and 599°C than expected in the null case.

Finally, both matrix IV and matrix V contain 88% POC from E. huxleyi culture biomass and 12% PIC from the JGOFS sediment, the closest analogue to the Great Calcite Belt samples. The absolute quantity of POC and PIC that were loaded into the RPO reactor differ, with matrix V containing twice the amount of carbon than matrix IV (Table 2). Despite this mass difference, the two thermograms are similar, and have similar deviations from their respective null thermograms. First, the highest temperature peak observed in the pure JGOFS sediment thermogram above 600°C is absent in the two matrix thermograms (Figs. 9d, 9e). The isotopic enrichment of the CO₂ released above 400°C relative to the CO₂ released in similar intervals across the E. huxleyi thermograms indicates that nearly all of the PIC expected to react above 600°C in the null hypothesis case has oxidized between ~420°C and 600°C (Figs. 10d, e). The fact that these shifts in thermal stability occur regardless of the differences in PIC mass in mixtures IV and V implies that these effects are predominantly driven by the matrix rather than by the absolute mass of PIC available for reaction. This result further implies that with high enough POC:PIC proportions in a mixture, matrix effects dominate amount effects on PIC thermal stability.

Overall, the results show that the mixing of POC and PIC can significantly lower the activation energy of PIC. Comparing matrix II, IV and V, the matrix effect on the PIC from the JGOFS sediment is non-linear and sensitive to matrix proportions. It is difficult to discern whether the higher sensitivity of the JGOFS sediment to matrix proportions stems from an intrinsic thermodynamic property of marine biogenic PIC, or from the non-carbon matrix in the sediment itself (i.e., 62% BSi) (Sayles et al., 2001). Using matrix IV and V thermograms as the best analogues to Southern Ocean particles in terms of POC:PIC, the presence of BSi in the sample matrix, and the marine origin of PIC in the particles, we conclude that the PIC in the Great Calcite Belt thermograms most likely reacts between 420°C and 600°C.

In addition, isotope comparisons of CO₂ across each matrix (excluding matrix III) and *E. huxleyi* thermogram allowed for calculation of POC mass in each matrix fraction (Eq. 9b). Comparisons to the null hypothesis POC quantities calculated in the matrix fractions (Eq. 7b) show that the thermal stability of POC has shifted in all matrix mixtures (Fig. 10). For example, the matrix 1 analysis shows that less CO₂ from organic carbon evolves below 427°C and more evolves between 427°C and 530°C (Fig. 10a). In the matrix II thermogram, more POC evolves between 325°C and 530°C, compared to the null case (Fig. 10b). In more POC-rich mixtures IV and V, POC-associated shifts are smaller (Figs. 10d, e). It is likely that a combination of matrix effects and the formation of thermally stable char from incomplete oxidation of POC increases the activation energy of distinct organic carbon pools in these thermograms (Williams et al., 2014). Because marine POC oxidizes over a wide temperature range, it is challenging to distinguish and correct for these possible effects in the Southern Ocean thermograms. The following discussion only takes into account matrix effects on PIC thermal stability, as such effects likely exceed the matrix/charring effects on POC thermal stability.

4.3 Narrowing thermal stability range with depth

All twelve blank-corrected (but not PIC-corrected) <51 μ m POC thermograms of samples collected above 800 m across GB1-117, GB2-43 and GB2-100 highlight striking differences between the thermal stability of euphotic zone POC and that of mesopelagic zone POC that exceed differences among stations (Fig. 6). In the euphotic zone thermograms, where most sample carbon derives from POC (PIC content is \leq 5% of total C), CO₂ is released over a broad, bimodal temperature distribution with a lower temperature peak T_{max} at 355 ± 15°C (average ± 1 standard deviation across seven euphotic zone thermograms) and a higher temperature peak T_{max} at 508 ± 5°C. By contrast, mesopelagic zone POC reacts over a narrower range of oxidation temperatures (Fig. 6), and is distributed more homogenously over this narrower range, rather than distributed bimodally around two peak temperatures. This can be measured by computing the temperature range over which the normalized CO₂ concentrations exceed the entire thermogram half maximum (Fig. 11a-c; Appendix B, Fig. S4). The temperature range decreases in thermograms from z_{PAR} to the deeper mesopelagic zone. A similar pattern appears when considering just the CO₂ released at temperatures below 420°C, where little PIC decomposes (Section 4.2; Fig. 11d-f; Appendix B, Fig. S4). The temperature range over which evolved CO₂ concentrations exceed the half maximum CO₂ concentration below 420°C is significantly lower among the mesopelagic zone thermograms than among the euphotic zone thermograms (Student's t-test, p<0.05).

At the same time, the proportion of total carbon that decomposes to CO₂ within a middle range of temperatures or activation energies increases with depth (Fig. 11g-i; Appendix B, Fig. S4). We define the lower bound of this range as the average T_{max} of the lower temperature peak across the seven euphotic zone thermograms (355°C). The upper bound of this range is operationally defined as the highest temperature at which we expect only sample POC (and no PIC) to decompose to CO₂ due to the matrix effects described in Section 4.2. This fraction is significantly higher in the mesopelagic zone thermograms than in the euphotic zone thermograms (p<0.05). The ingrowth of this midtemperature fraction with depth is balanced by a decrease in the fraction of POC that oxidizes between 100°C and 355°C, and a decrease in the PIC-corrected fraction of POC that oxidizes above 420°C at stations GB2-100 and GB1-117. This highest temperature POC fraction increases with depth at GB2-43.

The transition from a defined POC partitioning between two activation energy distributions in euphotic zone particles to a more homogenous distribution of POC across a narrower, intermediate range of activation energies occurs regardless of the "starting" POC distribution in the euphotic zone thermograms (Fig. 6), and regardless of the composition of phytoplankton communities producing and exporting the POC from the surface (Table 1). The data suggest that the broader distribution of thermal stability exhibited in the euphotic zone thermograms is characteristic of fresh organic matter, recently produced by photosynthesis. This also may explain why the thermograms of pure and freshly produced *E. huxleyi* culture biomass (Fig. 5) resemble the euphotic zone

thermograms more than they resemble deeper mesopelagic zone thermograms, even though they are compositionally different from the complex biomass sources to particles in the open ocean.

Thermograms of the two euphotic zone $>51 \,\mu m$ particle samples analyzed from GB2-43 (Fig. 6), which have a very low PIC and blank component (Rosengard et al., 2015), also overlap closely with blank-corrected thermograms of the <51 µm samples from the same depths and station. Interestingly, the δ^{13} C values of CO₂ collected across the 85 m >51 μ m thermogram are 1-6‰ lighter than the δ^{13} C values across the <51 μ m thermogram. This indicates that, in spite of similarities in thermogram shape, the POC in the two size-fractions is compositionally distinct (Fig. 8), possibly due to POC production by different size classes of phytoplankton (Trull and Armand, 2001; Finkel et al., 2016), and/or minimal exchange of organic carbon between the two size-fractions (Wakeham and Canuel, 1988; Abramson et al., 2010). Thus, the broad bimodal distribution evolved CO₂ in both size fraction thermograms reflects their similar diagenetic state, irrespective of compositional differences. It is possible that dissolution of >51 µm POC to DOC and incomplete flushing of the total POC pool off of the polyester pre-filters during sonication introduced methodological artifacts to the >51 µm POC thermograms and gas fraction δ^{13} C measurements. These processes would remove certain organic carbon subpools (Buesseler et al., 1998) and/or shift the thermal stability of the remaining organic carbon matrix prior to RPO analysis. Nonetheless, the close overlap in thermogram shape between the two POC size-fractions suggests that these artifacts are minimal.

4.4 Increasing diversity of $\delta^{13}C_{POC}$ with depth

The stable isotope composition of POC oxidized throughout thermograms further allows us to explore the biomolecular changes that accompany transitions in thermogram shape (Fig. 11) between fresher euphotic zone samples and more degraded mesopelagic zone samples. Significant variation in the δ^{13} C values of CO₂ released during ramped oxidation of POC-replete euphotic zone samples and *E. huxleyi* culture biomass (Figs. 5, 8) likely reflects the heterogeneity of biomolecular classes produced by different metabolic pathways (Hayes et al., 1990; Freeman, 2001). To compare δ^{13} C values across thermograms of Southern Ocean POC, it is necessary to subtract the PIC component from the 420°C-600°C range of all thermograms. The δ^{13} C measurements across CO₂ fractions of each sample thermogram are not significantly enriched within any narrow temperature range above 420°C (Fig. 8), indicating that the PIC component, which varies between -2‰ and 2‰ (Fig. 2d-f), is spread unevenly throughout the 420°C- 600°C interval, rather than as a predictable peak (e.g., Fig. 4a). As a result, we bisect every bulk carbon and deployment blank thermogram into two components above and below 420°C that comprise fractional proportions of A₁ and A₂ within intervals *T₁* and *T₂*, respectively, and only subtract the acid hydrolysis-based PIC contributions to the bulk isotope composition within T₂. This strategy to compare the δ^{13} C distribution across two components is largely methodologydriven, not an intrinsic property of marine thermal stability.

Towards this end, we first calculate the δ^{13} C value of CO₂ released within intervals T_1 and T_2 of each bulk sample thermogram and deployment blank thermogram by inverting the following matrix (Rosenheim et al., 2008):

$$\begin{bmatrix} \delta^{13}C_1 \\ \vdots \\ \delta^{13}C_i \end{bmatrix} = \begin{bmatrix} f_1, T_1 & f_1, T_2 \\ \vdots & \vdots \\ f_i, T_1 & f_i, T_2 \end{bmatrix} \times [\delta^{13}C_{T_1} \, \delta^{13}C_{T_2}]$$
(10)

This inversion solves for δ^{13} C values of CO₂ released in temperature intervals T₁ and T₂: δ^{13} C_{Bulk,T1}, δ^{13} C_{Bulk,T2}, δ^{13} C_{Blank,T1}, and δ^{13} C_{Blank,T2}. In Equation 10, *i* is the number of CO₂ fractions trapped and analyzed, δ^{13} C_i is the stable isotope value measured in each fraction, and $f_{i,T1}$ and $f_{i,T2}$ are the fractions of A_1 and A_2 in each temperature interval over which fractions were collected. The sum of all fractions in the same row of the matrix equates to 1.

Next, each $\delta^{13}C_{blank,T1}$ and $\delta^{13}C_{blank,T2}$ value calculated from the deployment blank thermograms was subtracted from each $\delta^{13}C_{Bulk,T1}$ and $\delta^{13}C_{Bulk,T2}$ value calculated from the <51 µm sample thermograms, respectively (Equation 11a, 12a). The ¹³C-enriched PIC contribution was subtracted from the $\delta^{13}C_{Bulk,T2}$ values from each sample thermogram (Equation 11b, 12b).

$$\delta^{13}C_{\text{POC},\text{T1}} = \left[\delta^{13}C_{\text{Bulk},\text{T1}} - \delta^{13}C_{\text{Blank},\text{T1}}f_{\text{Blank},\text{T1}}\right]/f_{\text{POC},\text{T1}}$$
(11a)

$$\delta^{13}C_{\text{POC},\text{T2}} = \left[\delta^{13}C_{\text{Bulk},\text{T2}} - \delta^{13}C_{\text{PIC}}f_{\text{PIC},\text{T2}} - \delta^{13}C_{\text{Blank},\text{T2}}f_{\text{Blank},\text{T2}}\right]/f_{\text{POC},\text{T2}}$$
(11b)

$$f_{POC,T1} = 1 - f_{Blank,T1} \tag{12a}$$

$$f_{POC,T2} = 1 - f_{Blank,T2} - f_{PIC}$$
(12b)

The fractions of blank carbon in each component of the bulk sample thermogram, $f_{Blank,T1}$ and $f_{Blank,T2}$, were calculated by dividing the integrated area of the blank thermograms within each temperature interval by the same integrated area within each non blankcorrected sample thermogram. $\delta^{13}C_{PIC}$ was determined by acid hydrolysis (Section 2.2). The fractional contribution of PIC to the bulk sample carbon in T₂, $f_{PIC,T2}$, was calculated by multiplying f_{PIC} (Table 1) by the total carbon in each non blank-corrected sample thermogram, and then dividing this product by the total carbon within T_2 of the sample thermogram.

Figure 12 illustrates $\delta^{13}C_{POC,T1}$ and $\delta^{13}C_{POC,T2}$ profiles across three stations (Table 4). Because of small kinetic fractionation of carbon during ramped oxidation, we only consider isotopic differences >2‰ in magnitude significant (Hemingway et al., accepted). The δ^{13} C values of the deepest samples at GB2-43 and GB2-100 have a very high propagated error (up to ± 17.0 %), owing to relatively high blank and PIC corrections at these depths (Table 4). We assume that the error of sample $\delta^{13}C_{T1}$ and $\delta^{13}C_{T2}$ values calculated in Equation 10 prior to blank and PIC subtraction are similar to the precision of the IRMS, 0.1 ∞ . At GB2-43, the weighted sum of the δ^{13} C values of these two 800 m components is 6.7 % heavier than the bulk $\delta^{13}C_{POC}$ value (Table 4, Table 1). For this reason, we only focus on δ^{13} C values above 800 m at GB2-43. The weighted sum of the 800 m δ^{13} C values at GB2-100 agree more closely with the bulk $\delta^{13}C_{POC}$ value at that depth, but the individual component-specific $\delta^{13}C$ values, -42 % for T₁ POC and -16 ‰ for T₂ POC, exceed the range of previously observed δ^{13} C values for organic matter in the Southern Ocean (Trull and Armand, 2001; Cavagna et al., 2013). Thus, at GB2-100, we cautiously include the 800 m δ^{13} C values in our interpretation. Finally, we are more confident about the GB1-117 δ^{13} C profiles because the weighted δ^{13} C sums of the two components (δ^{13} C_{POC,T1} and δ^{13} C_{POC,T2}) closely match the bulk $\delta^{13}C_{POC}$ values at all depths, and the errors of $\delta^{13}C_{POC,T1}$ and $\delta^{13}C_{POC,T2}$ at 1000 m are smaller.

While absolute values of and changes in $\delta^{13}C_{POC,T1}$ and $\delta^{13}C_{POC,T2}$ through each profile differ by station, the difference between the two values ($\delta^{13}C_{POC,T1} - \delta^{13}C_{POC,T2}$) $\Delta\delta^{13}C_{POC}$ increases with depth at all stations. $\Delta\delta^{13}C_{POC}$ is small (-2 to +2‰) in the euphotic zone and generally increases to >3 ‰ (Fig. 12d-e, Table 4). In all of the mesopelagic zone samples for which we could calculate $\delta^{13}C_{POC,T1}$, the lower thermal stability component is consistently heavier than the higher thermal stability component. In GB2-43, this positive $\Delta\delta^{13}C_{POC}$ is primarily due to an increase in the depth-averaged $\delta^{13}C_{POC,T1}$ from the euphotic zone to the mesopelagic zone, with no change in $\delta^{13}C_{POC,T2}$. In GB2-100, this positive difference is a combination of increasing average $\delta^{13}C_{POC,T1}$ and a decreasing $\delta^{13}C_{POC,T2}$ with depth. In GB1-117, the positive $\Delta\delta^{13}C_{POC,T1}$.

The increase in $\Delta \delta^{13}C_{POC}$ with depth implies increasingly different biomolecular groups are distributed across a narrower range of activation energies. While we cannot determine the identity of the biomolecules that persist in the deeper mesopelagic zone samples, two trends are clear. First, when depth-averaged $\delta^{13}C_{POC,T1}$ values change with depth, as in the GB2-43 and GB2-100 profiles, they become more ¹³C-enriched. Second, when depth-averaged $\delta^{13}C_{POCT2}$ values shift with depth, as in GB2-100 and GB1-117, they become more ¹³C-depleted. Thus, at each station, relatively heavy low thermal stability biomolecules and/or lighter high thermal stability molecules accumulate with depth, as well. The low thermal stability biomolecules may contain proportionally more amino acids and carbohydrates because they are relatively enriched (Deines, 1980; Galimov, 2006). A weak but significantly positive correlation (p<0.05) between $\delta^{13}C_{POC,T1}$ values and molar C:N ratios in particle sub-samples analyzed by elemental analysis (Table 1, Table 4; correlation not shown) suggests that the heavier biomolecules preserved within interval T_1 could be more carbohydrate-derived rather than amino acidderived, because amino acids have relatively low C:N ratios. Meanwhile, the high thermal stability POC pool that grows lighter with depth could be influenced by an increasing proportion of ¹³C-depleted lipids (Hayes et al., 1990). It is important to acknowledge that these interpretations are limited by the possibility that thermograms further reflect the decomposition of not only discrete biomolecules, but sub-groups of larger molecules that are isotopically distinct from the rest of the compound (Wagner et

al., 1994; Westerhout et al., 1997). Thus, the compounds accumulating in T_1 could also be enriched fragments of other structurally complex biomolecules, not necessarily carbohydrates, decomposing to CO_2 within this temperature interval.

4.5 Selective biomolecular preservation

The changes in thermogram shape with depth across station profiles (Section 4.3, Fig. 11) support the argument for selective preservation of POC. At all stations, the proportion of the lowest activation energy POC components in the euphotic zone samples decreases with depth as the fraction of POC oxidizing over a narrow, middle range of activation energies accumulates. It is not surprising that the lowest activation energy compounds are most reactive in the water column, especially if these thermally unstable components are smaller and more readily available to bacterial uptake (Benz and Bauer, 1988). The preferential removal of POC from higher thermal stability ranges with depth in profiles at GB2-100 and GB1-117 is more unexpected. However, it is possible for the most thermally stable POC pools to be equally degradable in the water column because marine heterotrophs use enzymes to break apart structurally complex and large biomolecules before assimilation (Chröst, 1991; Hedges et al., 2000; Sinsabaugh et al., 2009). Thus, the data demonstrate that thermal stability is not a straightforward proxy for diagenetic stability, at least on the higher thermal stability end of the spectrum (Section 1).

Because the range in activation energy distributions of POC in the water column decrease regardless of differences in euphotic zone POC composition, the data support the perspective that a similar suite of degradation/organic matter alteration reactions in the water column transform organic matter regardless of its source composition when exported (Baldock et al., 2004). This trend is similar to observations that the proportion of labile and identifiable organic compounds (amino acids, carbohydrates and lipids) decreases with water column depth as the proportion of stable and molecularly uncharacterizable carbon (MUC) accumulates (Wakeham et al., 1997; Hedges et al., 2000). Wakeham et al. (1997) were only able to characterize ~30% of Equatorial Pacific POC collected in sediment traps at 1000 m as either amino acids, lipids or carbohydrates.

This amount was significantly smaller than the characterizable quantity in shallower POC, from fresh phytoplankton biomass in the euphotic zone to sediment traps at 150 m (~80%). Large proportions (>50%) of MUC have been observed in other sections of the Equatorial Pacific with different rates of primary productivity, and elsewhere, including deep sections of the Cariaco and Santa Barbara Basins, the Eastern Subtropical Atlantic, Northeast Pacific, and Southern Ocean (Hwang and Druffel, 2003; Roland et al., 2008; Wang and Druffel, 2001).

It is likely that over half the POC in the deeper mesopelagic zone samples of each Great Calcite Belt profile is also molecularly uncharacterizable POC. This would imply that the processes that render POC uncharacterizable also manifest in a narrower, more homogenous POC distribution across activation energies of decomposition (Fig. 11). Hypotheses for MUC formation include structural linkages among biomolecules that make them difficult to access by exoenzymes or too large to pass through bacterial cell membranes, and the creation of metabolic intermediates during heterotrophic remineralization (Del Rio et al., 1996; Wakeham et al., 1997; Hedges et al., 2000; Freeman, 2001). Both processes would lead to the loss of characterizable compounds and lower the biomolecular diversity of mesopelagic zone POC, which may also manifest in a more homogenous distribution of POC about a smaller activation energy range. It is unlikely that abiotic cross-linking alone would decompose the highest activation energy pools of euphotic zone POC, which is thermodynamically unfavorable (Hedges et al., 2000). More likely, a combination of cross-linking and heterotrophic reworking of organic matter by enzymes accounts for the observed removal of both the lowest and highest activation energy POC pools with depth.

While the POC activation energy distribution narrows with depth, POC becomes more isotopically diverse (Section 4.4, Fig. 12d-e). If a significant proportion of MUC in the thermograms of deeper samples were formed by linkages of discrete biomolecules into larger, recalcitrant structures during remineralization in the water column, we would expect $\Delta \delta^{13}C_{POC}$ values to decrease with the combination of isotopically distinct biomolecules. The fact that $\Delta \delta^{13}C_{POC}$ increases instead points to conversion of euphotic zone POC to metabolic intermediates by heterotrophy and/or exchange of biomolecular fragments during cross-linking followed by break up into isotopically diverse smaller molecules, both of which could drive greater variations in isotopic composition of the POC pool.

4.6 Compositional differences in accumulated POC

Selective preservation implies that MUC is compositionally and structurally distinct from the identifiable biomolecule classes, and accumulates with depth because it is intrinsically recalcitrant. The individual $\delta^{13}C_{POC,T1}$ and $\delta^{13}C_{POC,T2}$ profiles (Fig. 12a-c) illustrate three different scenarios of selective POC preservation from three ecologically distinct sites of the Southern Ocean (Table 4), indicating that compositionally different biomolecules persist in deeper mesopelagic zone POC. With the information available, there are no clear relationships among phytoplankton community composition, the <51 µm [POC] attenuation (Table 1), POC stability and the identity of these biomolecules that accumulate with depth.

Differences in deep $\delta^{13}C_{POC}$ composition among stations could result from different sources of euphotic zone POC. Trull and Armand (2001) observed that different size-fractions of organic particles in the Southern Ocean exhibited different $\delta^{13}C$ values, ranging by as much as ~8‰ between 1-5 µm to >200 µm size classes, and that selective feeding of distinct size classes could shift bulk $\delta^{13}C$ with depth. In this data set, we saw that >51 µm POC is isotopically distinct (i.e., lighter) from <51 µm POC in the euphotic zone (Section 4.3; Fig. 8). But, because sample size limitations precluded the analysis of >51 µm POC by RPO in the mesopelagic zone, size-specific degradation processes are beyond the scope of this study. In the future, applications of ramped oxidation could be designed to explore the contribution of factors such as advection and size-partitioning to compositional changes in POC through the water column.

Differences in the community composition of heterotrophs in the mesopelagic zone could also cause differences among $\delta^{13}C_{POC,T1}$ and $\delta^{13}C_{POC,T2}$ profiles across stations. Indeed, the conversion of POC to heterotrophic biomass can comprise a significant fraction of POC at these depths (McCarthy et al., 1998; Hedges et al., 2000; Griffith et al., 2012). Accumulation of different heterotrophic communities at different rates in mesopelagic zone particles could mask any relationships between phytoplankton

community composition, thermal stability and the identity of biomolecules preserved from the euphotic zone. More work remains to determine whether the thermal stability and activation energy range for heterotrophic biomass would manifest in the distributions observed in the mesopelagic zone (Figs. 6, 11). Future ramped oxidation studies should focus on controlled "end-member" experiments, such as comparing axenic E. huxleyi biomass thermograms (Fig. 5) to cultures inoculated with bacterial biomass.

One lingering caveat in this discussion is the closed system perspective that <51 μ m POC in the mesopelagic zone derives from the POC produced in the euphotic zone. Deeper advection of even more highly degraded <51 μ m particles could further influence the organic matter matrix we observe in the mesopelagic zone and drive differences in $\delta^{13}C_{POC,T1}$ and $\delta^{13}C_{POC,T2}$ profiles in Fig. 12 (Griffith et al., 2012; Cavagna et al., 2013). If significant enough, advection would impact the conclusion that MUC derives from selective preservation (Roland et al., 2008). However, it would not change the position that the same processes that form and concentrate MUC with depth also concentrate organic matter within a narrow activation energy distribution (Section 4.2).

5 Conclusion

Ramped oxidation of marine POC in the water column reveals significant differences in the composition of euphotic and mesopelagic zone $<51 \mu m$ POC, supporting the perspective that organic matter degradation/preservation in the water column selects for specific biomolecules or fragments of them. Taking into account matrix effects of sample minerals, particularly calcite, is important for accurately interpreting POC composition using this method. These data set the stage for future applications of ramped oxidation to samples collected from more controlled environments and towards testing specific hypotheses of organic matter transformation and fate during particle transport through the water column.

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Table 1. Bulk properties of particulate organic carbon (POC) and deployment blank filters sampled from the euphotic zone to 1000 m at stations GB1-117, GB2-43 and GB2-100 from the Southern Ocean Great Calcite Belt region. Bolded numbers correspond to values at (or above/below at GB2-43) z_{PAR} , the export depth, defined by the depth of 0.3% PAR. $f_{C,blank}$ and f_{PIC} are the proportions of total carbon that are deployment blank carbon and particulate inorganic carbon, respectively. POC error is equivalent to one standard deviation across three [POC] concentrations measured by different methods described in Section 2.2 (Table 3).

tation	1 pling cation	Depth	ature	POC] ficient	ency *	ivity*	from alysis	[PIC]	[BSi])C,blank	fpic	:[PIC]	¹³ CPIC	³ CPOC	C:N
õ	San Date/Lo	Ι	Temper	<51 µm [attenuation coeff	Export efficie	Primary Product	<51 µm [POC] RPO an	<51 µm	<51 µm	fc		<51 µm [POC]:	Q	õ	
-	-	т	°C	μΜ	%	mg/m^2d	μM	μM	μΜ	-	-	-	‰	‰	-
GB1-117	13-Feb-11	25	18.7				5.09 ± 0.63	0.177		0.03	0.03	29	1.0	-21.0	7.1
GB1-117	38.97°S	62	14.2	1.14	9	218	$\textbf{2.46} \pm \textbf{0.28}$	0.084	0.010	0.03	0.03	29	0.1	-21.9	6.3
GB1-117	9.49°E	162	11.6				0.39 ± 0.04	0.045		0.12	0.09	9	-0.04	-21.91	6.7
GB1-117	zPAR=	300	9.7				0.41 ± 0.02	0.019		0.19	0.04	22	-1.96	-20.14	5.1
GB1-117	62m	1000	5.4				0.12 ± 0.03	0.017		0.28	0.09	7	-0.91	-23.07	N=BDL
GB1-117 blank		N/A					35 ± 9.9^{b}	1.9 ^b		0.95	0.05	18	2.87 ^a	-31.6	N=BDL
GB2-43	1-Mar-12	20	6.5				1.75 ± 0.05	0.011		0.09	0.01	166	-1.7	-26.9	5.7
GB2-43	47.53 °S	85	6.2	0.81	43	140	1.76 ± 0.10	0.022	0.15	0.09	0.01	82	-0.4	-27.3	5.7
GB2-43	64.01°E	120	4.6	0.81	43	140	$\textbf{0.89} \pm \textbf{0.03}$	0.037	0.17	0.13	0.03	24	0.7	-27.8	5.1
GB2-43	zPAR=	300	3.0				0.28 ± 0.07	0.013		0.29	0.03	21	-0.6 ^a	-24.1	7.2
GB2-43	108m	800	2.6				0.13 ± 0.02	0.012		0.47	0.04	11	-0.9	-24.1	N=BDL
GB2-43 blank		N/A					57 ± 7.8^{b}	2.6 ^{a,b}		0.96	0.04	22	2.87 ^a	-31.6	N=BDL
GB2-100	14-Mar-12	20	13.0				$\textbf{2.91} \pm \textbf{0.38}$	0.133		0.04	0.04	22	1.5	-22.5	6.1

GB2-100	44.62°S	90	12.7				2.76 ± 0.02	0.151		0.05	0.05	18	0.6	-22.5	6.3
GB2-100	100.50°E	110	11.4	0.89	11	171	0.50 ± 0.02	0.010	0.0074	0.12	0.02	50	0.4 ^a	-24.2	7.6
GB2-100	zPAR=	300	10.4				0.22 ± 0.07	0.030		0.20	0.09	7	0.4	-21.4	7.8
GB2-100	113m	800	7.9				0.11 ± 0.05	0.028		0.33	0.14	4	-0.7	-21.6	9.4
GB2-100 blank		N/A					36 ± 15^{b}	3.2 ^b		0.92	0.08	11	2.87 ^a	-33.3	N=BDL
*Values published in Rosengard et al. (2015).															
^a averaged value of other station measurements because not enough PIC for resolving measurement/gas lost after acid-hydrolysis.															
^b normalized to the	^b normalized to the total active area of the Whatman TM QMA filters, 125 cm ² .														
BDL=below detection limit.															

Table 2. Quantities of POC and PIC in each end-member and matrix (I-V) sample RPO analysis. The roman numerals following the Icelandic Spar and JGOFS sample ID's refer to the analysis number. The numerals following the *E. huxleyi* strain numbers refere to the culture replicate number. The total carbon loaded into the RPO reactor, C_{Total} , was calculated as the total CO₂ evolved from the sample during ramped oxidation. $f_{E,hux,Total}$ and $f_{Calcite,Total}$ of each matrix mixture was calculated using the weighted sum δ^{13} C values of all CO₂ gas fractions across the matrix thermograms (Equation 6, Table 3).

	0	()	_		_			7 \
oldr	Irce	ırce	Tota	Tota	Tota	NC C	PIC	DIG
San	105	10S	hux	cite,	0	al P r R	tal] r R	ÿ
•1	C	IC	f _{E.}	fcal		fo	fol	PO
	PC	4				L		
			_	_	μg	μg	ug	
Icelandic Spar - I	N/A	N/A	0.00	1.00	450	0	450	0
Icelandic Spar -II	N/A	N/A	0.00	1.00	250	0	250	0
JGOFS MC-1-8 -I	N/A	N/A	0.06^{a}	0.94	950	57	893	0.06
JGOFS MC-1-8 -II	N/A	N/A	0.06 ^a	0.94	250	0	235	0.06
E.hux373-I	N/A	N/A	1.00	0.00	593	593	0	no PIC
E.hux373-II	N/A	N/A	1.00	0.00	570	570	0	no PIC
E.hux373-III	N/A	N/A	1.00	0.00	828	828	0	no PIC
E.hux624-I	N/A	N/A	1.00	0.00	678	678	0	no PIC
Matrix I	E.hux373-III	Spar	0.50	0.50	506	253	253	1
Matrix II	E.hux373-III	JGOFS	0.59	0.41	464	274	190	1
Matrix III	E.hux373-II	Spar	0.85*	0.15*	863	732*	132*	6*
Matrix IV	E.hux373-II	JGOFS	0.88	0.12	339	299	41	7
Matrix V	E.hux373-I	JGOFS	0.88	0.12	652	573	78	7

* Approximation because of incomplete isotope quantification.

^a Fraction of organic carbon in sample, assumed to be negligible in the matrix mixtures II, IV, and V.

Table 3. Carbon quantities and bulk δ^{13} C values of Great Calcite Belt samples and PIC end-members measured by different approaches. The three POC quantities were normalized to QMA filter area (125 cm²) and either calculated as the total CO₂ evolved during ramped oxidation, measured by closed tube combustion (CTC), or measured by two different elemental analyzers. The weighted sums δ^{13} C values of thermogram gas fractions were compared to separately measured δ^{13} C_{bulk} values.

ution	epth	Csum	Cbulk	ence	irom RPO	from v EA	from 2 EA	hree tities	S.D.
St	Q	Weighted δ^{13}	Measured δ^{13}	$\delta^{13} \mathrm{C}_{\mathrm{bulk}}$ differ	<51 µm POC 1]	<51 µm POC 1 CTC/Costech	<51 µm POC 1 <i>Flash 1112</i>	S.D. across t POC quant	Relative
-	meters	‰	‰	‰	µmol	µmol	µmol	µmol	%
Spar	N/A	2.7	3.0	-0.25	N/A	N/A	N/A	N/A	N/A
JGOFS	N/A	-1.1	-0.9	-0.18	N/A	N/A	N/A	N/A	N/A
GB1-117	25	-21.2	-20.5	-0.74	1272	1497	1573	157	11%
GB1-117	62	no data*	-21.5	no data	1262	1488	1530	144	10%
GB1-117	162	-22.3	-20.9	-1.36	237	281	269	23	9%
GB1-117	300	-24.3	-21.3	-3.04	153	154	165	7	4%
GB1-117	1000	-23.7	-23.0	-0.75	83	42	67	21	32%
GB1-117 blank	N/A	no data*	-29.81	no data	35	54	42	10	23%
GB2-43	20	-28.4	-27.1	-1.32	625	604	642	19	3%
GB2-43	85	no data*	-27.3	no data	616	637	686	36	6%
GB2-43	120	-28.3	-27.1	-1.20	390	373	368	12	3%
GB2-43	300	-26.7	-25.1	-1.60	143	116	187	36	24%
GB2-43	800	no data*	-25.9	no data	62	69	79	9	12%
GB2-43 blank	N/A	-33.1	-30.12	-3.00	57	66	50	8	13%
GB2-100	20	-21.5	-21.8	0.28	1007	no data	1193	131	12%
GB2-100	90	no data*	-21.8	no data	749	no data	755	4	1%
GB2-100	110	-26.1	-24.5	-1.62	284	279	297	10	3%
GB2-100	300	-22.1	-21.2	-0.90	136	116	192	40	27%
GB2-100	800	-24.1	-21.6	-2.53	63	no data	106	30	36%
GB2-100 blank	N/A	-30.8	-30.41	-0.36	36	40	64	15	32%

*not calculated because one or more gas fractions were lost during ramped oxidation.

Table 4. δ^{13} C values of CO₂ evolved within intervals T₁ (100°C - 420°C) and T₂ (420°C – 700°C) of each sample thermogram. These values are corrected for the contributions of blank carbon sorption and PIC to the total carbon in each temperature interval (Equations 10-12). Starred recombined bulk values deviate the most from the bulk δ^{13} C_{POC} reported in Table 1. Underlined numbers refer to the numbers used to calculate the average euphotic zone δ^{13} C_{POC,T1} and δ^{13} C_{POC,T2} values reported in Fig. 12.

Station	Depth	δ ¹³ C _{POC,T1}	$\delta^{13}C_{POC,T2}$	$\Delta \delta^{13} C_{POC}$	Recombined bulk δ ¹³ C
-	meters	%0	%0	$(T_1 - T_2) \%$	‰
117	25	-22.7±0.1	-21.0± 0.3	-1.7	-21.9
117	62	<u>-21.8±0.1</u>	-23.8±0.3	2.0	-23.0
117	162	-22.4±0.4	-27.7±1.4	5.3	-24.5
117	300	-23.7±0.8	-24.0±0.9	0.3	-23.9
117	1000	-24.0±0.9	-28.8±1.6	4.8	-26.0
43	20	-29.52±0.14	-26.9±0.1	-2.6	-28.2
43	85	<u>-30.0±0.1</u>	-27.9±0.2	-2.1	-29.0
43	120	-27.1±0.2	-31.0±0.3	3.9	-28.9
43	300	-23.0±1.5	-26.0±0.7	3.0	-24.7
43	800	-8.0±-7.0	-23.1±1.0	No data	-17.3
100	20	<u>-20.8±0.3</u>	-22.9±0.4	2.1	-22.1
100	90	No data	No data	No data	No data
100	110	<u>-26.2±0.4</u>	-25.7±0.2	-0.5	-26.0
100	300	-18.0±2.8	-26.5±3.2	8.5	-22.2
100	800	-16.0 ± 5.0	-42.0±17.0	No data	-26.3

*>4 ‰ different from $\delta^{13}C_{bulk}$



Figure 1. (a) Cruise tracks GB1 and GB2 across the Southern Ocean Great Calcite Belt region (image from Rosengard et al., 2015). The "x" notation indicates each profile of <51 and >51 μ m diameter particles. The color bar represents surface seawater temperature (°C). (b) >51 μ m [PIC]/[BSi] ratios at the export depth z_{PAR} , or the depth of 0.3% photosynthetically available radiation, of every particle sampling station (values reported in Rosengard et al., 2015). The bubble sizes scale with the ratio. The filled bubbles indicate the ratios at stations selected for RPO analysis.



Figure 2. (a-c) Profiles of $\delta^{13}C_{bulk}$ (‰) measured in <51 µm POC samples and deployment blanks at stations GB2-43, GB2-100 and GB1-117. (d-f) $\delta^{13}C_{PIC}$ (‰) profiles in samples and blanks, measured after acid hydrolysis. (g-i) $\delta^{13}C_{POC}$ (‰) profiles, calculated by Equation 3. In all panels, the horizontal lines are the depth of z_{PAR} (Table 1), and the gray stars represent the respective $\delta^{13}C_{bulk}$, $\delta^{13}C_{PIC}$ or $\delta^{13}C_{POC}$ values of the deployment blank collected at each station.



Figure 3. Thermograms of three Great Calcite Belt samples analyzed once after acid fumigation and rinsing (solid gray line), and again without any treatment (solid colored line). All y-axis values have been normalized to the 125 cm² active QMA filter area. The normalized y-axis values for the untreated thermogram were blank-corrected by subtracting the contribution from the filter-normalized blank carbon thermogram (dashed colored line). Carbon yield, reported on each panel, is the CO_2 quantity released in the fumigated sample run relative to the CO_2 quantity released in the non-fumigated sample run.



Figure 4. (a-b) Thermograms of two quantities of 100% PIC Icelandic spar analyzed by ramped oxidation. δ^{13} C (‰) of gas fractions, only measured in the 450 µg analysis, are represented by the height of the bars. (c-d) Thermograms of two quantities of JGOFS sediment from a Southern Ocean core top at 60°S, 170°W. δ^{13} C of gas fractions were only measured in the 1000 µg analysis. All y-axis quantities in panels a and c are normalized to the total CO₂ released during ramped oxidation.



Figure 5. Thermograms and gas fraction δ^{13} C values (‰; height of bars) of three replicates (rep. I, rep. II, rep. III) of *E. huxleyi* strain CCMP373 and one replicate of *E. huxleyi* strain DHP624. All y-axis quantities are normalized to the total CO₂ released during ramped oxidation.



Figure 6. Thermograms of total carbon in fifteen $<51 \mu m$ particle samples (solid gray lines) at five depths at GB2-43 (column 1), GB2-100 (column 2), and GB1-117 (column 3). All y-axis values represent CO₂ concentrations normalized to 125 cm² active QMA filter area. The dashed gray lines are thermograms of deployment blank carbon at each station. Blank thermograms were subtracted from the particle thermograms to yield sample carbon thermograms, shown as colored lines. The PIC contribution in each sample is represented in the area of the gray bar shown above 420°C (Table 1). Column 4 also shows thermograms of $>51 \mu m$ particles from two depths at GB2-43, which were not blank or PIC-corrected.



Figure 7. Thermograms of deployment blank filters and δ^{13} C of CO₂ gas fractions (‰; height of bars) collected within three specific temperature intervals. The y-axis is the range of CO₂ concentration normalized to the active area of the QMA filters.



Figure 8. Thermograms of total <51 μ m sample carbon and δ^{13} C of gas fractions (‰; height of bars) collected in specific temperature intervals. The δ^{13} C of gas fractions collected in one >51 μ m sample thermogram (GB2-43, 85 m) is shown in the fourth column. The y-axis is the range of CO₂ concentration normalized to the active area of the QMA filter (Fig. 6).



Figure 9. Thermograms of five artificial mixtures with varying proportions of *E. huxleyi* culture-derived organic carbon and PIC (colored solid lines) (Table 2). The source in matrices I and III is an Icelandic Spar (a, c), while the PIC source in matrices II, IV and V (b, d, e) is a Southern Ocean core top sediment collected during JGOFS (Sayles et al., 2001). Roughly equal molar proportions of POC and PIC comprise mixtures I and II, while mixtures III, IV and V are more POC-rich. The bar heights represent the δ^{13} C (‰) of CO₂ gas fractions within specific temperatures. The dashed lines represent "null hypothesis" thermograms generated by the linear addition of each organic and PIC-rich end-member thermogram scaled to the POC:PIC ratios in each matrix (Table 2).



Figure 10. The absolute difference (µmol) between the masses of culture- or PIC-derived CO₂ observed within temperature intervals of each matrix thermogram and expected within similar intervals of each null hypothesis thermogram. These differences are equivalent to $mass_{Calcite,obs,i} - mass_{Calcite,null,i}$ for the blue bars and $mass_{E.hux.,obs,i} - mass_{E.hux.,null,i}$ for the red bars, as defined by Equations 7-9. No actual POC and PIC differences were calculated in panel C because of a lost δ^{13} C measurement. The cumulative POC or PIC lost across some fractions does not always equal the cumulative POC or PIC mass gain in other fractions because of imprecision in the calculation of $f_{Calcite,obs,i}$ in Equation 8, which affects the calculations in Equation 9.



Figure 11. Three metrics that describe the narrowing thermal stability distribution of POC with depth across GB2-43, GB2-100 and GB1-117 (Table 1, Fig. 1). (a-c) The temperature range over which thermogram CO_2 concentration exceeds the half maximum. (d-f) The temperature range over which thermogram CO_2 concentration below 420°C exceeds the half maximum below 420°C. (g-i) The fraction of total POC that oxidizes between 100-355°C, 355-420°C, and 420-700°C. 355°C is the average T_{max} of the lowest temperature peak in the euphotic zone sample thermograms, while 420°C is operationally defined as the lower limit of PIC decomposition. The highest temperature fraction is PIC-corrected. In all panels, the dashed line is the depth of z_{PAR} (Table 1). Illustrations of how each metric was calculated are provided in Appendix B, Fig. S4.



Figure 12. (a-c) $\delta^{13}C_{POC}$ values (‰) of CO₂ oxidized within intervals T₁ and T₂ ($\delta^{13}C_{POC,T1}$ and $\delta^{13}C_{POC,T2}$, respectively) of the <51 µm sample thermograms. The starred values have high error bars due to relatively large blank and PIC corrections, and are either omitted from or cautiously interpreted in the discussion. The dashed lines represent the export depth, z_{PAR} (Table 1). (d) Average $\delta^{13}C_{POC,T1}$ (circles) and $\delta^{13}C_{POC,T2}$ (triangles) across euphotic zone (labeled "E" above data points) and mesopelagic zone (labeled "M" above data points) sample thermograms. The number of sample values averaged in each depth zone is indicated below the data points.

Chapter 5:

Diagnosing and quantifying organic carbon sources to suspended sediments in the Amazon River main stem

Abstract

The Amazon River mobilizes organic carbon across one of the world's largest ³ terrestrial carbon reservoirs, supplying ~6-8% of the global riverine particulate organic carbon (POC) flux to the oceans. Constraining the contribution of different terrestrial sources to this flux, and their fate in the coastal Atlantic Ocean, is challenging in such a vast and dynamic system, but essential to understanding the role of Amazonia in the global carbon cycle. Here, we identify and quantify different sources to the POC flux at Óbidos, the most downstream gauging station of the Amazon main stem, by integrating bulk, compound-specific, and ramped pyrolysis/oxidation (RPO) analyses of riverine POC. The latter uses a controlled temperature ramp to decompose samples into constituents with different thermal stabilities that can be further analyzed for isotopic composition (stable carbon and radiocarbon). Ten to eleven suspended sediment samples were analyzed from within the cross-section at Óbidos (Brazil) during two different stages of the river's hydrologic cycle. These data were compared to analyses of surface suspended sediment samples from the Madeira, Solimões and Tapajós Rivers. Bulk isotope and C:N compositions suggest that riverine POC in the Amazon River is predominantly soil-derived, while the RPO data point to input from three thermally distinct soil pools from a range of landscapes and/or degradation histories. The RPO data also show that up to 4% of total main stem POC is petrogenic, confirming earlier assessments based on bulk OC radiocarbon analysis. Using Acoustic Doppler Current Profiler transects data from 2014, we calculate the flux of these soil and petrogenic components through the cross-section at Obidos. Finally, using RPO analyses of POC collected from three other major tributaries upstream and downstream of Obidos, we discuss the diagenetic stability of the four thermally distinct POC pools as they are exported to the ocean.

1 Introduction

The Amazon River Basin, a global biodiversity hotspot, is one of the largest reservoirs for reactive organic carbon on the planet. The river network includes eight

major tributaries and supplies $\sim 6.3 \times 10^{12} \text{ m}^3$ /year of water (Molinier et al., 1996) to the coastal Atlantic Ocean, comprising 20% of the global freshwater flux. Draining through six million square kilometers of flooded rainforest, drier savannah, high-elevation Andean landscape, as well as cropland and pasture, the river delivers $\sim 12-14 \text{ Tg/yr}$ particulate organic carbon (POC) from the basin to the Atlantic Ocean, $\sim 6-8\%$ of the global annual riverine POC export to the ocean (Richey et al., 1990; Galy et al., 2015). Accurately tracking the fate of this massive flux has bearings on the residence time of terrestrial organic carbon from the Amazon reservoir, and our understanding of the Amazon basin's role in the global carbon cycle (Cole et al., 2007; Aufdenkampe et al., 2007; Butman et al., 2016;).

There are several options for the fate of POC mobilized by the Amazon River (Cole et al., 2007): it can be respired by bacteria or photo-oxidized in transit (Richey et al., 1990; Amon and Benner, 1996; Mayorga et al., 2005) or in the coastal Atlantic Ocean (Aller et al., 1996; Blair and Aller, 2012), buried for the shorter-term in the river bed or flood plains (Guyot et al., 1996), or buried for longer time scales on the continental shelf (Kastner and Goñi, 2003). Quantifying how riverine POC partitions among these pathways relies on accurately measuring carbon fluxes and describing the POC composition in suspended river sediments. POC composition is particularly important because several studies have shown that certain classes of biomolecules in the organic matrix preferentially degrade during transit in the river (Mayorga et al., 2005; Ward et al., 2013), suggesting that POC quality is important for the fate of Amazon River POC.

Several challenges constrain our ability to accurately measure POC flux and composition in the Amazon River Basin. One challenge is where measurements and river sampling occur. Most studies of riverine geochemistry and POC transport through the Amazon River main stem occur near the municipality Óbidos in the state of Pará (Fig. 1a), the most downstream gaging station that is still outside the zone of salt intrusion from the Atlantic Ocean, ~900 km upstream of the river's mouth. The river's particle composition at Óbidos integrates sediment input from most of the major Amazonian tributaries, including the Solimões and Madeira rivers. Still, measurements at Óbidos exclude the influences of influx from the Tapajós, Trombetas, Xingu and Tocantins rivers. The omission of these downstream tributaries has bearings on interpretations of POC flux to the ocean from the Amazon River, and has encouraged more recent studies to sample along transects of the river system between Óbidos, or sites upstream, and the river plume (e.g., Ward et al., 2015). Further, interpreting point measurements at one location of the river network is limiting because large rivers systems like the Amazon are heterogeneous in space and time. Within the Amazon cross-section, hydrodynamic sorting causes large grain-size sediments to settle faster and thus concentrate at deeper depths (Rouse 1950; Gibbs, 1967; Curtis et al., 1979; Bouchez et al., 2011b), leading to compositional differences between the deeper, coarser sediments and the shallower, finer sediments (Bouchez et al., 2011a; Bouchez et al., 2014). Finally, seasonality presents another challenge to measuring fluxes. Water levels in the Amazon River alone fluctuate by at least a factor of two between the rainy and dry seasons (Moreira-Turcq et al., 2013), which manifests in variability in POC flux and composition as well (Richey et al., 1990; Kim et al., 2012; Ward et al., 2015).

This study addresses some of these challenges by combining measurements of POC flux and composition along three dimensions: among tributaries near Óbidos, within the cross-section of the main stem, and across time. Integrating a suite of geochemical analyses, from bulk POC composition to compound-specific lipid assessments, the results identify key sources of POC mobilized by the Amazon River. We show how coupling these measurements to measurements of POC thermal stability using the ramped pyrolysis/oxidation (RPO) technique (Rosenheim et al., 2008; Rosenheim and Galy, 2012; Rosenheim et al., 2013) enables us to partition POC flux at Óbidos among these sources. The application of RPO by geochemists has been particularly useful for environments that integrate diverse sources of organic carbon that vary in age and degradation history, such as large, complex river systems. This study utilizes RPO to construct an age distribution of POC exported at Óbidos and comments on the reactivity of these different sources in the river system.

2 Methods

2.1 Sampling location and timing

Suspended sediment sampling in 2014 focused on a region of the Amazon main stem near the city of Óbidos (Table 1, Fig. 1a). In 2014, main stem discharge at Óbidos varied by a factor of two between the low discharge dry season (Southern Hemisphere spring/summer) and the high discharge rainy season (Southern Hemisphere winter/fall) (Fig. 1b-c). When sampling occurred in March/April and July 2014, the river levels were high, near their peak in May/June. In April, the river levels were rising, and in July, river levels were falling.

The 2014 sampling effort at Óbidos was supplemented by several additional sampling locations (Table 1, Fig. 1a), including one additional sample that was collected from Óbidos during another sampling campaign in June 2005, one near the Madeira River mouth to the Amazon River (~Foz Madeira) and one near the Solimões River mouth to the main stem (~Manacapuru) from that same expedition. The Tapajós River was further sampled in July 2014 to allow for comparison between Óbidos and its most immediate downstream tributary. Additional details of the June 2005 sampling campaign are described in (Bouchez et al., 2011b).

2.2 Suspended sediment collection

Suspended sediments were filtered from two types of river water samples: a large volume (100-200 L) sample of surface river water and a small volume (10 L) sample at select depths below surface. The exact volumes were calculated by weighing the water samples after collection and converting mass to volume using the density of fresh water, 1 kg/L. With the exception of the 2005 samples, all water samples were filtered in pressurized Teflon units through 0.22 μ m diameter pore size Millipore PES membrane filters within two days after collection. Between collection and filtration, samples were either covered in a dark tarp or stored in the shade to minimize exposure to sunlight. After filtration, the particles on the filters were immediately frozen in a freezer on board the ship. Although previous studies have highlighted the compositional differences across particle size fractions in the Amazon River (Hedges et al., 1986; Hedges et al., 1994; Aufdenkampe et al., 2001; Aufdenkampe et al., 2007), we have chosen to pool all particles above 0.22 μ m in diameter for analysis (Bouchez et al., 2014).

The large volume samples were collected at Óbidos (April 2014, July 2014 and June 2005), and each upstream and downstream tributary site using a bucket submerged a few centimeters below the surface. Each sampling location was roughly in the center of each river channel. The 10 L samples were collected within three depth profiles at Óbidos in April and July 2014, using a depth-specific isokinetic sampler. Each profile, consisting of three to five depths from 2-3 m to >50 m, was located at a different position in the channel of the main stem, from near the right bank to near the left bank (Óbidos side of the river crossing) (Table 1). The depth-specific water sampler was equipped with a depth sensor to accurately record the collection depths.

In addition to suspended sediment samples, bedload samples from the Tapajós River and main stem at Óbidos were taken using a bedload sampler in July 2014. A flood deposit sample from the right bank of the main stem, across from Óbidos, was also collected and frozen on the ship.

2.3 Enzyme activity assays

Enzyme assays were conducted on suspended particles from the July 2014 Óbidos depth profiles to probe variations in heterotrophic activity with variations in particle composition in the Óbidos cross-section. Samples were analyzed using–AMC (7-amino-4-methylcoumarin) and –MUB (4-methylumbelliferone)-based fluoregenic substrate proxies (Steen and Arnosti, 2013; Table 2) and buffered with 100 mM carbonate buffer (pH=6.85). Assays were performed in triplicate for all samples and for one control, boiled to denature all enzymes, which served as a blank correction. Fluorescence was measured three to four times during 4 hour sample and control incubations in 1 mL cuvettes at ambient temperature (Table 1). Enzyme activities in each sample were inferred by the blank-subtracted substrate hydrolysis rates calculated from these four time points.

2.4 Flux calculations

During each sampling month in 2014, water velocity profiles were measured across the main stem channel using a *Sontek* RiverSurveyor® M9 Acoustic Doppler Current Profiler (ADCP). The ADCP operated on a 1 MHz frequency, and was equipped with a 0.5 MHz vertical beam sensor for river depth measurements and an external GPS for compass heading, latitude and longitude tracking. The external GPS did not function over most of the transect in April, so an average offset from the external GPS was applied to correct the internal *Sontek* compass, which did provide continuous heading measurements. The raw velocity cross-sections and depth gauges helped guide the choice of sample depths and locations for each depth profile (Table 1, Section 2.2).

The *Sontek* RiverSurveyor-M9 ADCP only measured three velocity components (E, N, U) down to 40 m depth. Horizontal velocities, V_{mag} , were calculated as the vector combination of the E (east) and N (north) components. Because 40 m was often shallower than the river bed and a further 10% of each velocity profile data was discarded prior to data export, the deepest measured velocities had to be extrapolated to the river bed, using the following relationship between horizontal velocity, V_{mag} , and vertical distance above river bed, *z* (Chen, 1989; Mueller et al., 2009):

$$V_{mag} = a_v z^{1/6} \tag{1}$$

The a_v values for each ADCP transect position were first extracted by fitting the measured V_{mag} profiles to Equation 1. Then, the extrapolated V_{mag} values for all depths between the deepest measured V_{mag} and the river bed were calculating using the fitted a_v values.

For each sampling month, the RiverSurveyor® software integrated velocities measured through the instrument's transect and modeled within the "edge" regions between each end of the instrument transect and the nearest river bank to calculate a water discharge value through the river's cross-section. Using the transect velocity data exported from the software, we further calculated suspended sediment fluxes at Óbidos during each sampling month by integrating the product of suspended sediment concentration and water velocity across channel position and over river depth (Bouchez et al., 2011b):

$$TSS \, flux = \iint V_{mag} C_z dx \, dz \tag{2}$$

In Equation 2, dx values were calculated by projecting the distance of the boat track using ADCP transect coordinates (Fig. 1d-e) against the azimuth of the main stem at Óbidos. Profiles of V_{mag} across the channel were a combination of measured and extrapolated values (Equation 1).

Suspended sediment concentration, C_z , values in Equation 2 were modeled according to the hypothesis that the Amazon River main stem was deep enough in April and July 2014 to allow for separation of sediment grain size and density by depth (Rouse 1950). Following Bouchez et al. (2011b), who also reported hydrodynamic sorting in the Amazon main stem, all depth-specific (from 10 L samples) suspended sediment concentrations from each sampling month were fitted to a Rouse equation, which relates sediment concentration to river depth:

$$\frac{C_z}{C_a} = \left(\frac{H-z}{z}\frac{a}{H-a}\right)^{Z_R}$$
(3)

In this relationship, C_z is suspended sediment concentration at z, the vertical distance above the river bed, which has a depth of H. a is the depth of the shallowest point measurement in the depth profile, C_a is the suspended sediment concentration at a, and z_R is the Rouse number. We used a nonlinear least-squares fit to calculate the Rouse number for April and July. The Rouse fits allowed us to model C_z across the entire main stem cross-section by varying H from the ADCP data, but using constant C_a values, averaged across the surface-most measurement of the three channel positions in each sampling month.

2.5 Bulk particle composition

In the laboratory, each suspended sediment sample was re-suspended from the filters in milli-Q water, and freeze-dried. Bed samples were directly freeze-dried. The dried sediment and bed samples were sieved through a 2 mm mesh to remove any coarse impurities such as small rocks fragments and large leaf debris, and homogenized before subsequent analyses. The homogenized quantities of suspended sediments were weighed to calculate suspended sediment concentrations as mg homogenized sediment/L water filtered, which were later applied to Equation 3.

The distribution of grain sizes in the homogenized particle samples was analyzed using a Beckman Coulter Laser Diffraction Particle Size Analyzer (LS 13 320). Samples were sonicated for 10 seconds in tap water before loading into the detector. The LS 13 320 detects particles in the 0.4 μ m to 2 mm size range, and reports mean and median grain size for each sample. The software additionally calculates the volumetric contribution of particle sizes throughout the distribution, as well as the d10, d50, and d90 diameters, which delineate the 10th, 50th and 90th percentiles of the size distribution.

Bulk weight % organic carbon (% OC), weight % nitrogen (% ON), δ^{13} C, and δ^{15} N in the homogenized suspended and bed sediments were analyzed using a *Fisons Instruments Carlo Erba 1108* elemental analyzer interfaced via a *Finnigan MAT Conflo II* to a *Delta-Plus Stable Light* isotope ratio mass spectrometer (IRMS). Prior to measuring % OC and δ^{13} C, sub-samples were weighed, loaded in Ag boats and fumigated in concentrated hydrochloric acid (12 N HCl) vapors for 72 hours at 60°C, and then dried in a desiccator at 60°C for 72 hours to remove the inorganic carbon contribution to the total carbon in the sediment (Whiteside et al., 2011). One bedload sediment from Óbidos (July 2014) was analyzed just after 12-16 hours of acid fumigation. Weighed sub-samples were not fumigated prior to measuring %N and δ^{15} N. All analyses were conducted in triplicate. The results only report the averages of the triplicate measurements, and their standard deviation as an estimate of analytical precision.

For bulk radiocarbon composition, a sub-sample of each sediment was similarly decarbonated via acid fumigation for 72 hours. After drying, the fumigated sample was sealed in an evacuated quartz tube with 2 g copper oxide, and baked at 850°C for 6 hours, which converted all the sample organic carbon to CO_2 gas (McNichol et al., 1995). The evolved CO_2 was then cryogenically purified under vacuum and subsequently graphitized by iron catalysis in pure H₂ gas at 450°C and analyzed for its radiocarbon composition at the National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) facility at Woods Hole Oceanographic Institution (McNichol et al., 1992).

2.6 Ramped Oxidation

Ramped pyrolysis/oxidation (RPO) offers a higher resolution approach to analyzing the composition of organic carbon in complex environmental settings. The technique oxidizes OC in a sediment sample at a steady, controlled temperature ramp from ~100°C to up to 1000°C, and monitors the release of CO₂ throughout the ramp. Plots of evolved CO₂ concentration against temperature, hereafter referred to as thermograms, separate organic carbon by its thermal stability. The reaction can be coupled to a vacuum line to trap CO₂ gas produced over specific user-defined temperature intervals for stable and radiocarbon isotope analysis.

All samples were analyzed by ramped oxidation at the National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) facility following methods described in Rosenheim et al. (2008) and Hemingway et al. (accepted). Sub-samples were placed into a quartz reactor receiving a constant flow of ultra high-purity helium carrier gas and oxygen (~92%:8%). As the temperature increases in the reactor, the carbon in the sample oxidizes and volatilizes into the carrier gas, passing through an 800°C oven equipped with a catalyst wire (nickel, cadmium and platinum) to ensure complete conversion of the combustion products to CO₂. Downstream of the ovens, a *Sable Systems*© *CA-10* infrared gas analyzer continuously measures the concentration of evolved CO₂ in the carrier gas matrix.

Beyond the gas analyzer, the CO₂ is cryogenically frozen between user-selected temperature intervals and re-expanded into a vacuum line, where the gas is purified (cryogenic water stripping) and flame-sealed in pyrex tubes with 50 mg copper oxide and 10 mg silver for analysis. These tubes were baked at 525°C for 1 hour as an additional step to convert any incomplete oxidation products to CO₂, as well as to remove any impurities from the gas fraction, prior to graphitization and isotope analysis. These gas fractions were then re-quantified in the vacuum line using a pressure transducer (baratron). One split of the gas fractions was converted to graphite for radiocarbon measurements (reported as fraction modern, or F_m), while the rest was analyzed by the *VG Isotech Prism II* IRMS to measure stable isotope composition (reported as δ^{13} C). The sum of all gas quantities recovered from each sample after the bake step was compared to the amount of POC loaded into the reactor (calculated as sediment mass times %OC of that sample) to monitor the yield of the ramped oxidation.

We have analyzed six depth-specific suspended samples from Óbidos 2014 depthprofiles center channel, and each large volume sample from 2005 and 2014 (Table 1). Four to five fractions of CO₂ gas were trapped and analyzed during each ramped oxidation analysis. One depth-specific sample was fumigated for 72 hours, dried for 72 hours and rinsed in Q-H₂O prior to ramped oxidation. Rinsing was necessary to remove residual Cl from fumigation, which reacts with the catalyst wire, forming precipitates at higher temperatures (>400°C) that clog the reactor. The thermogram generated by this treated sample was compared to the thermogram of another sub-sample that was analyzed without any acid treatment. Based on the results of this comparison, discussed in Section 3.4, the rest of the samples were not acid-treated prior to ramped oxidation.

2.7 Compound-specific lipid analysis

Abundances of specific biomolecules provide even higher-resolution details of organic matter composition. Straight-chain *n*-alkanes, *n*-alcohols, and fatty acids were quantified in each large-volume sample collected in 2014 and 2005 and four depth-specific samples collected in 2014 (Table 1). Total lipids were extracted from sediment into 15-20 mL of 9:1 dichloromethane (DCM)/methanol at 100°C for 20 minutes using a Microwave Accelerated Reaction System (MARS, CEMS Corp.). The total lipid extracts were then saponified in 15 mL of 0.5 M potassium hydroxide (KOH) in wet methanol at 70°C for two hours. After adding 20 mL milli-Q water and 0.5 g of sodium chloride to the KOH solution, the basic lipids were extracted from the aqueous phase via five hexane rinses. The remaining KOH solution was acidified to pH~2 using 12 N HCl to isolate the acidic lipids in five rinses with 4:1 hexane/DCM.

Each basic and acidic lipid fraction was separated into five biomolecular classes on the basis of polarity. The concentrated lipid fractions were loaded onto aminopropyl silica gel columns and sequentially flushed with hexane (*n*-alkanes), 4:1 hexane/DCM (ketones), 9:1 DCM/acetone (sterols, alcohols and other polar compounds), 2.6% oxalic acid in methanol (fatty acids), and 1:1 DCM/methanol (residual). Both acidic and basic fraction-derived fatty acids were re-combined and methylated in 95:5 methanol:HCl with a known δ^{13} C value and ¹⁴C composition overnight, for 12-16 hours, at 70°C. The fatty acid methyl esters (FAMES) were isolated on an additional aminopropyl silica gel column after methylation. Prior to compound-specific isotope analysis of the straightchain saturated FAMES and *n*-alkanes, two additional purification steps were required. First, for δ^{13} C analysis, all FAMES and *n*-alkanes were urea-adducted in 40 mg/mL methanol urea solution to separate the branched compounds from straight-chain compounds.

All compound abundances were measured using a flame ionization detector coupled to a *Hewlett Packard 5890 Series II* Gas Chromatograph (GC-FID). Ureaadducted, combined acid and base fractions of *n*-alkanes were injected in high purity hexane. The urea-adducted FAMES were directly injected in hexane. In addition, the non-adducted fractions of FAMES and *n*-alkanes, as well as the other fractions of the post-methylation FAMES columns were analyzed in the FID-GC. Any "residual" FAMES and *n*-alkane quantities in these other fractions were added to the FAMES and *n*alkane quantities from the purified fractions, and are reported as such in the next section.

All GC-FID analyses were accompanied by a suite of standard *n*-alkanes and FAMES to cross-reference sample peak retention times and quantify compound abundances by peak areas. Some analyses were accompanied by just one standard injection at one known concentration, while others were accompanied by injection of three different standard concentrations. When three standard concentration chromatograms were available, standard curves were applied to sample peak areas to estimate analyte mass (nanograms). When analyses included just one standard concentration, a response factor, equivalent to the average peak area/ng compound across standard compounds, was used to convert sample peak area to mass.

The δ^{13} C values of specific FAMES and *n*-alkanes were measured using a *HP6890* gas chromatograph fitted with a *Gerstel* PTV and interfaced via a *Finnigan MAT Conflo II* to a *Delta-Plus* IRMS. When compounds were abundant enough, they were analyzed in duplicate or triplicate.

2.8 Statistics and error analysis

We equate the error of the average %OC, %N, bulk δ^{13} C and δ^{15} N, and compound-specific δ^{13} C to the standard deviation of the replicate analyses of each sample. The relative error in bulk F_m values ranged from 0.2%-0.6%, while the relative error in the F_m values of RPO fractions ranged from 0.2%-1.2%. The errors of all RPO fraction δ^{13} C values of CO₂ fractions trapped during ramped oxidation were 0.1‰. Because these RPO δ^{13} C and F_m errors are relatively small, they are not reported in the figures and tables of the following sections. The average relative error of the lipid masses per extraction, quantified from the GC-FID, is ±10%. Errors were propagated for any metrics calculated from these values, such as methylation-corrected compound-specific δ^{13} C values and POC-normalized lipid abundances.

3 Results

3.1 Discharge at Óbidos

In 2014, maximum river depth of the Amazon main stem at Óbidos was 67-68 m (mode~61 m) in both April and July (Fig. 1f-g). The M9 RiverSurveyor ADCP logged >1400 velocity profiles from surface to riverbed in April and >1300 profiles in July. In April, the average water velocity was 1.9 ± 0.1 m/s across the transect, only slightly faster in July, 1.6 m/s \pm 0.1 m/s. Velocities were always highest (3-4 m/s) across the deeper part of the channel, where depths exceeded 40 m. Velocities near the river bed were slower, less than 1.5 m/s. Because of similar velocity distributions and river depths, total discharge at Óbidos did not change significantly between sampling months, ranging from 246,000 m³/s in April and 247,000 m³/s in July (Table 1). This agrees with continuous gauge measurements near Óbidos conducted by Brazil's Agencia Nacional de Aguas, which show that in 2014, monthly-averaged discharge peaked at ~290,000 m³/s in May and June. Water levels were also highest during these months. Thus, the two sampling months in 2014 reflect different river stages, rising in April and falling in July, on both sides of the peak discharge. The following sections describe variations in the quantity and organic composition of suspended particulate material across the channel at

Obidos, from surface to river bed, and between the two stages of the hydrological cycle, April/rising water and July/falling water.

3.2 TSS concentrations and fluxes at Óbidos

The total suspended sediment concentrations ([TSS]) throughout the main-stem cross-section at Óbidos varied from 55.5 to 318.5 mg/L in April, and from 16.4 to 741.4 mg/L in July (Fig. 2, Table 3). Concentrations are highest in the deepest samples of the near-right bank profiles in both April and July 2014. Because the right bank (across from the town of Óbidos) is located on the inside of a bend in the Amazon main stem, the combination of more turbulent water and lower horizontal velocities may entrain a greater amount of suspended sediments from the river bed, leading to higher TSS concentrations in this section of the river cross-section.

[TSS] concentrations increased with depth in all profiles at Obidos (Fig. 2, Table 3). Generally, all grain size metrics, including mean, median/d50, d10, and d90 values increased with depth and [TSS], providing clear evidence for hydrodynamic sorting of sediment in the cross-section (Rouse 1950; Bouchez et al., 2011b; Bouchez et al., 2011a). All the [TSS] values measured in 10 L profile samples within each sampling month were fitted to the Rouse equation (Equation 3). Channel depths varied more by channel position than by season (Fig. 1). z_a , the reference height above the river bed, varied from 36.9 m to 56.2 m in April and from 41.5 to 58.9 m in July. C(a), the suspended sediment concentration at z(a), varied from 71.6 to 106.8 mg/L in April and from 43.9 to 102.7 mg/L in July. We chose to combine profiles within each sampling rather than calculate one Rouse number per channel position profile per month because Bouchez et al. (2011b) found that the errors from fitting individual, more scarcely resolved depth profiles from a specific channel position exceeded errors from fitting a combined profile across the entire river transect.

Fits of April TSS concentrations yielded a Rouse number of 0.26, while fits of July TSS concentrations yielded a Rouse number of 0.34 (Table 1). The modeled concentration profiles fit the data well, suggesting that the Rousean description of hydrodynamic sorting adequately explains our observations (Fig. 3). The integrated

product of [TSS] and water velocity within each cross-section yielded suspended sediment fluxes of 53,000 kg/s in April and 48,000 kg/s in July (Equation 2). These values do not take into account the water discharge and sediment concentrations in the cross-sectional distance between each river bank and the ends of the ADCP transects (i.e., the "edge" discharge values), but discharge in these "edge" sections of the transect represents less than 1% of total discharge at the main stem.

3.3 Bulk POC composition

The weight % organic carbon (%OC) within suspended sediments at Óbidos ranged from 0.5 - 1.52% and decreased with depth in all profiles across channel positions and seasons, except within the right bank profile in July, which exhibited no significant change in %OC (Table 3, Fig. 2). At the same time, particulate organic carbon concentrations ([POC]), which ranged from 0.6 - 5.9 mg/L, increased with depth as TSS concentrations increased. Weight % nitrogen (% N) values in suspended sediments ranged from 0.05 - 0.24% and also decreased with depth. Molar C:N ratios exhibited a large range from 7.4 to 14.1, and tended to increase with depth in all profiles. Even though [TSS], %OC, and [POC] varied within profiles at Óbidos, the isotopic composition of organic carbon in the sediments did not vary significantly with depth. δ^{13} C values ranged from 3.1% to 4.3%. The range in fraction modern (F_m) values was also small (0.74 - 0.80), corresponding to a radiocarbon age span of 680 years (1790 to 2470 years).

The composition of suspended POC was compared to bed and bank sediments collected near Óbidos and the Tapajós River in July, as well as several large volume surface sediments collected from the Amazon River at Óbidos (April 2014, July 2014 and June 2005), the Tapajós River (July 2014), the Solimões River (June 2005), and the Madeira River (June 2005) (Table 1). The suspended sediment concentration at the main stem surface in 2005, reported in Bouchez et al. (2011b), was within the range observed in the surface and depth-specific sediment samples collected in 2014. %OC, %N and stable isotope composition of POC in all main stem surface samples from 2005 and 2014

were similar to the POC composition of depth-specific samples from 2014. The only difference was that the POC in these surface samples was consistently older, with F_m values as low as 0.71, than the POC analyzed from the depth-specific samples.

Surface suspended sediment concentrations in the Solimões and Madeira rivers, collected in 2005, were also similar to the range observed at Óbidos (Bouchez et al., 2011b). The % OC in these sediments ranged from 0.49% in the Madeira River to 1.1% in the Solimoes River (Table 3). %N ranged from 0.08% in the Madeira River to 0.17% in the Solimoes River. Stable carbon and radiocarbon measurements of POC in the Solimões and Madeira river sediments were slightly more ¹³C-enriched and older than the Óbidos depth-specific samples. The Tapajós River had the most compositionally distinct suspended sediments of the data set. Surface suspended sediment concentrations in July 2014 were much lower than observed at any other river, only 0.52 mg/L. But, % OC in these sediments was the highest measured, 15.7%. %N was also high, 2.7%. The F_m value was 0.89, corresponding to a younger ¹⁴C age than any of the values measured at Óbidos.

The weight % OC and %N in the three bed sediment samples from near the left bank of Óbidos, the right bank of Óbidos, and the Tapajós River were lower than corresponding values in all depth-specific samples, ranging from 0.06% to 0.14% organic carbon and from 0.01% to 0.02% nitrogen, consistent with other bedload observations in the basin (Bouchez et al., 2014). F_m values of the bed samples varied from 0.60 to 0.87 at Óbidos, both older and younger than the POC in the depth-specific samples, respectively, while the Tapajós bed sample was modern ($F_m > 1.0$), younger than POC in the Tapajos suspended sediment. %OC and %N of right bank flood deposit near the Óbidos crosssection were more similar to the range in depth-specific sediments, 0.5% and 0.05%, respectively. The stable isotope composition of the flood deposit was closer to the range observed across depth-specific samples, as well.

3.4 Ramped oxidation results

Suspended sediment sub-samples from three depths of the centermost profile of each sampling month at Óbidos, as well as four surface sediment samples were selected
for thermal stability analysis via ramped oxidation (Table 4). In addition, the deepest (54 m) sample collected in July 2014, was analyzed twice: once following fumigation in hydrochloric acid to remove particulate inorganic carbon, and once without any treatment (Table 4). The fumigated sample was further rinsed in Milli-Q water to remove any residual Cl (Section 2.5). The rest of the samples were analyzed untreated.

We note a small difference between the CO_2 evolved in the acid-treated vs. untreated thermograms of the deep July sample between 200°C and 400°C and above 550°C (Fig. 4). At the same time, the difference between the mass-weighted sums of δ^{13} C values from both thermograms, -29.3‰ to -29.1‰, is not statistically significant (Tables 3. 4). These sums are also comparable to the δ^{13} C value of bulk POC, -28.4‰, which was measured after sample fumigation but without rinsing (Tables 3, 4). This 0.7% - 0.9% difference is similar to the observed average difference between the sum of RPO fractions and the bulk δ^{13} C value for a compilation of data (Hemingway et al., accepted). The fact that the weighted sum δ^{13} C values from the untreated sample analysis are similar to the bulk δ^{13} C value of sample POC implies that the quantity of PIC is likely negligible relative to the POC quantity and that acid fumigation did not greatly shift the thermal stability distribution of POC in the sample. This agrees well with observations that the PIC content in Amazon riverine POC is negligible (Galy et al., 2007; Mayorga et al., 2005; Bouchez et al., 2014). Because all the untreated thermograms are very similar to each other (Fig. 4a- bottom right), we assume that the PIC quantity is small in all of them. For these reasons, we did not fumigate any other samples prior to ramped oxidation, and solely interpret and compare the untreated sample thermograms in the remaining discussion. In fact, all untreated sample thermogram carbon yields, calculated as the fraction of total mass of CO₂ gas quantified after trapping and baking relative to the amount expected from the bulk % OC and sediment mass, averaged around $99 \pm 5\%$, much greater than the yield observed for the fumigated sample, 86%.

The weighted sum F_m value across fractions of the untreated sample thermogram is 0.82, 0.02 units (2.5%) higher than the bulk POC F_m value of 0.80 (Tables 3, 4). This difference likely results from loss of a small fraction of older, ¹⁴C-poor carbon during fumigation. By comparison, the weighted F_m sum from fractions of the fumigated and rinsed sample thermogram is 0.77, 0.03 units (3.5%) lower than the separate F_m

measurements of the bulk POC and 0.05 units lower than the weighted sum F_m from the untreated sample thermogram. Further, CO₂ evolved in the two lower temperature intervals of the treated sample thermograms, which represents 52% of total sample carbon, has a lower F_m value than CO₂ evolved in corresponding intervals of the treated thermogram (Fig. 4a), suggesting that acid fumigation and rinsing remove younger, ¹⁴C-enriched and thermally unstable POC pools from the suspended sediment, which would also lower the carbon yield during ramped oxidation.

All six untreated sample thermograms overlap closely across depths and seasons, oxidizing from 100°C to 800°C (Fig. 4). Most of the carbon evolves at temperatures below 700°C, roughly the temperature at which several ramped oxidation analyses were terminated. But some analyses that continued recording CO₂ concentrations above 750°C show that a small fraction of total sample carbon (0.5%, averaged across thermograms) oxidizes to CO₂ between 700°C and 800°C. Stable carbon isotope values measured within specific CO₂ fractions of the thermograms ranged from -32.2‰ to -25.4‰ (Fig. 4b), while F_m values ranged from 0.39 to 0.94 (Fig. 4b).

All the surface sediments from Óbidos in 2005, and the Tapajós, Madeira, and Solimões Rivers generated broadly similar thermograms compared to thermograms from the depth-specific Óbidos samples (Fig. 4). The only exception was the Tapajós thermogram, which shows that little carbon oxidizes above 600°C. The δ^{13} C and F_m values in fractions of CO₂ collected across the 2005 Óbidos, Solimões and Madeira sample thermograms were similar to values observed in CO₂ fractions from all depthspecific thermograms. Again, the Tapajós sample thermogram generated the most isotopically distinct fractions of all runs, lighter (δ^{13} C= -35.1‰ to -31.49‰) and younger (F_m = 0.82- 0.91).

For some analyses, one to two fractions were lost during CO_2 recovery in the vacuum line, graphitization or preparation for radiocarbon analysis (Fig. 4). If just one fraction is lost, the F_m value of that fraction can be calculated using the separately measured F_m value of the bulk POC (Equation 4a). But, $F_{m,Bulk,POC}$ values do not equate the weighted sum F_m values from the RPO analyses of samples in which no fractions were lost, owing to kinetic fractionation during the temperature ramp (Hemingway et al., accepted) and different sample preparation prior to bulk and RPO analysis. Thus, we used

a linear regression of the weighted sum $F_{m,sum}$ values from all other RPO analyses where no CO₂ fractions were lost and the separately measured $F_{m,bulk,POC}$ values of these samples (Equation 4b), and applied the best fit line to Equation 4a.

$$F_{m,lost} = \frac{([F_{m,Bulk POC} \times 1.24] - 0.17) - \sum_{1}^{i} f_{i} F_{m,i}}{f_{lost}}$$
(4a)

$$F_{m,sum} = 1.24 \cdot F_{m,Bulk POC} - 0.17$$
 (4b)

Here, *i* is the number of measured isotope values from the CO₂ fractions, and f_i and f_{lost} are the fractional contribution of each measured and lost fraction to total sample carbon, respectively. The F_{m,Bulk POC} term in Equation 4a refers to the F_{m,bulk,POC} value of the sample with the lost CO₂ fraction. The regression (r²= 0.98) was applied to calculate the F_m value of CO₂ fractions lost from two April 2014 (49 m and 3 m) sample analyses, but not for the fraction lost in the Madeira River sample thermogram because two RPO gas fractions were lost and thus the mass proportions each fraction lost are not known (Fig. 4a). The linear regression between bulk and weighted sum δ^{13} C values was weaker, and thus was not used to calculate the δ^{13} C values of the missing CO₂ gas fractions. The regression was even weaker when assuming an intercept of 0, implying that δ^{13} C of lost gas fractions could not be calculated by a constant offset, either.

In general, CO₂ fractions tended to get more ¹⁴C-depleted (older) and more ¹³Cenriched with increasing temperature during ramped oxidation (Fig. 4a). The magnitude of variation in isotopic composition of CO₂ collected from thermograms of tributary samples was less than the variation from Óbidos sample thermograms. The range of F_m values across Óbidos sample thermograms varied from 0.42 to 0.53. The range was much lower in the Tapajós suspended sediment thermogram (0.14) and the Solimões 2005 thermogram (0.22). The range of all δ^{13} C values across the Óbidos thermograms varied from 3.63-6.39. Again, this range was slightly lower in the Tapajós and Solimões 2005 samples. No range was calculated for the Madeira River sample thermogram because the first CO₂ fraction, likely the youngest and most depleted, was lost prior to any isotopic analysis.

3.5 Lipid abundances and isotope composition

Total lipids were extracted from all large-volume suspended sediment samples from the surface of the Amazon River main stem (June 2005, April 2014 and July 2014), the Tapajós River, the Solimões River and the Madeira River, as well as from four depthspecific samples from the two 2014 Óbidos center profiles (Table 1). Compound-specific δ^{13} C values were not resolved for the April 2.7 m sample, where abundances extracted were too small.

Abundances of straight-chain *n*-alkanes in Óbidos samples (2014, 2005) were quantified for sixteen carbon chain lengths from 19 (C₁₉) to 35 (C₃₅), and ranged from 0.001 to 0.4 ng/µg of total POC (Appendix C, Table S1, Fig. 5c-e). The distribution of abundances above chain lengths C₂₃ displayed an odd-over-even carbon chain length predominance in all samples, characteristic of terrestrial vegetation (Eglinton and Hamilton, 1963) (Fig. 5c-e). Within each sampling month, the POC-normalized compound abundances measured at the surface were greater than the abundances measured deeper in the water column. Average chain lengths (ACL) for *n*-alkanes (Equation 5a; Jeng, 2006) ranged from 29.6 to 30.1 and decreased slightly with depth, but did not vary significantly between the two months (Fig. 6b). Stable isotope values of odd chain-length leaf wax-derived *n*-alkanes exhibited a larger range than all δ^{13} C values measured in bulk and across single thermograms from the Óbidos depth profiles, ranging from -37.3‰ to -28.3‰ (Appendix C, Table S2; Fig. 5 a-b).

$$ACL_{n-alkane} = \frac{25[C_{25}] + 27[C_{27}] + 29[C_{29}] + 31[C_{31}] + 33[C_{33}] + 35[C_{35}]}{[C_{25}] + [C_{27}] + [C_{29}] + [C_{31}] + [C_{33}] + [C_{35}]}$$
(5a)

$$ACL_{fatty acid} = \frac{24[C_{24}] + 26[C_{26}] + 28[C_{28}] + 30[C_{30}] + 32[C_{32}] + 34[C_{34}]}{[C_{24}] + [C_{26}] + [C_{28}] + [C_{30}] + [C_{32}] + [C_{34}]}$$
(5b)

Straight chain *n*-alkane abundances in the Solimões and Madeira river surface suspended sediments collected in 2005 spanned a similar range (Fig. 5c). *n*-alkanes from the Solimões River exhibited odd-over-even predominance, but the *n*-alkanes from the Madeira River did not. Average chain lengths in these samples were similar to the values calculated from Óbidos n-alkanes in 2014 (Fig. 7b). Compound-specific δ^{13} C values were not measured in these samples. Abundances of saturated straight-chain fatty acids (C_{14} - C_{34}) measured at Óbidos ranged from 0.008 to 4.0 ng/µg POC (Appendix C, Table S2; Fig. 7d-i). The most abundant compounds were C_{16} and C_{18} . At chain lengths above 23, the abundances exhibited even-over-odd predominance in all samples, reflecting an important contribution from terrestrial vegetation. River stage and depth-related differences in the abundance of even chain-length compounds were negligible at chain-lengths above C_{22} . The ACL of fatty acids (Fig. 6a; Equation 5b; Jeng, 2006) increased with depth in the April 2014 profile, from 27.6 at the surface to 28.6 at 49 m. The increase in ACL with depth in July 2014 was smaller. The stable isotope distribution across even chain-length fatty acids in the depth-specific samples spanned a greater range than bulk $\delta^{13}C_{POC}$ values in the same samples, -34.2‰ to -27.7‰ (Fig. 6b-c).

Straight-chain fatty acids were also quantified in surface suspended sediment samples from Óbidos (June 2005), and the Tapajós, Solimões and Madeira rivers. All Óbidos, Solimões River and Madiera river samples exhibited similar fatty acid abundances (Appendix C, Table S2; Fig. 7d,g). Only the Tapajós River sample showed very elevated concentrations of C_{14} , C_{16} and C_{18} that were ~4-40 times higher than the abundances of these fatty acids across all other 2014 and 2005 samples. All 2005 and tributary samples displayed strong even-over-odd predominance among the higher chainlength fatty acids. ACL values in all samples except Tapajós POC were similar to the range observed in the Óbidos profiles (Fig. 6a). The ACL calculated from Tapajós River fatty acids was 25.8, the lowest of the data set. All fatty acids in these samples were more depleted than the values observed in the 2014 Óbidos profiles (Fig. 7a). The $\delta^{13}C_{C22}$ value of Tapajós River POC was -39.4‰. The differences in $\delta^{13}C$ values across samples were smallest in the highest chain-length fatty acids (> C_{28}).

3.6 Enzyme activities

Activities of eight enzymes measured at various depths within the Obidos crosssection in July 2014 ranged from 0 to 0.09 μ mol/L-hr (Fig 8; Appendix C, Table S3). Values generally did not vary with depth or channel position. Activities of the N- and Pyielding enzymes (i.e., leucyl aminopeptidase, phenylalanyl aminopeptidase, and phosphomonoesterase) (Table 2), were exceeded activities of the C-yielding enzymes (i.e., cellulase, beta-glucosidase and xylanase) throughout profiles.

4 Discussion

The following sections first examine the dynamics underlying variation in TSS load and POC loading with cross-sectional depth. Then, coupling bulk and compoundspecific data to thermograms generated by ramped oxidation of these sediment samples, we quantify the sources of organic carbon exported at Óbidos and discuss the reactivity of these different sources in the river network.

4.1 From depth profiles to POC flux at Óbidos

Increasing [TSS] with depth in all profiles compiled at Óbidos in April and July 2014 are indicative of hydrodynamic sorting (Rouse 1950), and consistent with prior depth profiles of the same size-fraction of sediments compiled by Bouchez et al. (2011b) across the central Amazon River Basin (Fig. 2). This is supported by even lower TSS concentrations in the surface large-volume samples (<1 m depth) (Table 3) and a concurrent increase in grain size (i.e., mean and median diameter, as well as d10, d50 and d90) with depth. The Rouse numbers that we report are similar in magnitude and temporal variation to the numbers estimated from profiles at Óbidos in March 2006 ($z_R = 0.24$) and June 2005 ($z_R = 0.39$) (Bouchez et al., 2011b).

It is interesting that z_R values are so different between April and July 2014, despite the lack of change in river discharge between months. Significantly higher mean, d50 and d90 grain diameters in July relative to April (Student's t-test, p<0.05), especially in the deeper locations of the channel cross-section, could lead to greater hydrodynamic sorting of [TSS] in July and therefore higher z_R values. Considering that water levels in the Negro and Solimões rivers, the confluence of which forms the Amazon River main stem, upstream of Óbidos, are usually higher in July than in April, their increased input to Óbidos in July may contribute to larger grain sizes in the cross-sectional profiles (Moreira-Turcq et al., 2003). As the Negro River is very sediment-deplete (Kim et al., 2012), variations in Solimões discharge contribute relatively more to variations in sediment grain size through time in the main stem. By comparison, discharge and [TSS] in the Madeira River, the other major tributary to the main stem between the Solimões River and Óbidos, decreases between April and July (Moreira-Turcq et al., 2003; Bouchez et al., 2011b). Thus, a higher relative input of discharge and suspended sediments from the Solimões River to Óbidos between April and July could cause significant increases in the mean, median, d50 and d90 grain sizes in suspended sediments at Óbidos, creating a larger gradient in [TSS] between river surface and river bed and hence a higher z_R value in [TSS] profiles.

Suspended sediment fluxes, 53,000 kg/second in April and 48,000 kg/s in July (Table 1), are similar in magnitude but less variable than fluxes estimated by Bouchez et al. (2011b) in June 2005 and March 2006. Collectively, these quantities would greatly exceed flux estimates based on surface [TSS] measurements alone. For example, our fluxes are more than five times greater than surface-based fluxes calculated by Kim et al. (2012) during two expeditions to Óbidos in 2005 and 2009. The comparison is not perfect because the authors measured [TSS] in a slightly different size fraction of particles (>0.7 μ m). Nonetheless, the differences highlight the necessity and impact of accounting for cross-sectional variation in [TSS] when calculating TSS fluxes out of the river cross-section.

While TSS concentrations increase with depth, the %OC and %N per unit sediment weight decrease in all April and July 2014 profiles at Óbidos (Fig. 2, Table 3). Bouchez et al. (2014) argued that associations between POC and mineral surfaces and/or coincident concentration of fine carbon rich particles from buoyant organic debris drive declines in POC loading with coarser sediments with depth. But, the increase in C:N values across profiles suggest an accumulation of vegetative debris with depth. This is especially apparent in the deep section of Óbidos in July, which bears a higher F_m value than in April, indicating input of younger organic carbon at depth. Bulk POC F_m values are also older near the surface in July than in April, pointing instead to greater addition of finer, older and possibly more buoyant organic matter flushed in from floodplain soils during receding river levels.

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At the same time, [POC] (mg/L) values increase with depth because [TSS] concentrations are much higher in deeper sections of the river. For each sampling season, [POC] correlates positively with [TSS] in all profile samples (p<0.05). Linearly regressing the two concentrations yields a slope (mg [TSS]/mg [POC]) and intercept (mg/L [POC]), which can then be substituted into the Rouse-modeled C_z in Equation 3 to calculate POC fluxes during each sampling month:

$$[POC] = m[TSS] + b \tag{6}$$

In this equation, m is the slope of the linear relationship (0.0079 mg TSS/mg POC in April and 0.0072 mg TSS/mg POC in July) while b is the intercept (0.46 mg POC in April and 0.075 mg POC in July). The difference in the intercepts between months suggests that more POC in April is not associated with the mineral load in the suspended sediments.

The integrated POC flux in April 2014 was 540 kg/s, while the integrated POC flux in July 2014 was 370 kg/s (Table 1). This difference can be explained by a significant decrease in the average %OC in suspended sediments from $1.2 \pm 0.2\%$ (mean ± 1 S.D.) in April to $0.88 \pm 0.27\%$ in July (Student's t-test, p<0.05). Again, these fluxes are similar to those reported in Bouchez et al. (2014), and ~50-200% higher than those reported by Kim et al. (2012). Extrapolating these fluxes over one year results in an estimated annual POC flux from Óbidos of 12-17 Tg/year, which is likely an overestimation because April and July 2014 represent the high discharge period and therefore do not capture the full variation expected in TSS fluxes expected over an annual cycle in the Amazon River basin. Nonetheless, our annual fluxes are similar to earlier flux estimates of 14 Tg/year by Richey et al. (1990) who deployed a depth-integrated sampler to measure TSS concentrations (Richey et al., 1986).

4.2 Prominent soil-derived POC source to the main stem

Earlier studies of carbon cycling in the Amazon River partitioned riverine POC between two size fractions, a <63 μ m "fine" fraction and a >63 μ m "coarse" fraction, arguing that the smaller size fraction derived from older, degraded soils while the coarser fraction derived from fresher vegetation debris (Hedges et al., 1994; Aufdenkampe et al.,

2001; Aufdenkampe et al., 2007). Compound specific abundances of *n*-alkanes and fatty acids measured in one size-fraction (>0.22 μ m) of suspended sediments from various depths at Óbidos and at the surface of the Solimões and Madeira Rivers support the perspective that terrestrial vegetation is indeed an important source to riverine POC within the main stem cross-section as well as upstream. Odd-over-even predominance of the higher chain length *n*-alkanes (C₂₅-C₃₅), even-over-odd predominance of the higher chain length *a*cids (C₂₄-C₃₄), and relatively high average chain lengths (ACL) of each compound class reflect input from land plants (Figs. 5-7). Further, *n*-alkanes and fatty acids are more depleted than the bulk δ^{13} C_{POC} of each sample, especially at higher chain lengths (C₂₄-C₃₄ for fatty acids; C₂₅-C₃₅ for n-alkanes), which indicate that these compounds derive from terrestrial plants (Cranwell, 1982).

However, at the same time, the bulk composition of these samples vary little with depth and season (Table 3, Fig. 2) and are largely characteristic of soil organic matter (Quesada et al., 2010; Kim et al., 2012), suggesting that Amazon River POC derives predominantly from a mixture of soils. The radiocarbon age and stable isotope distribution of POC across sample thermograms generated by ramped oxidation of main stem samples supports this perspective. No CO₂ gas fraction is modern, exhibiting a bomb ¹⁴C component, and all gas fraction δ^{13} C values are within the range observed for Amazonian soils (Fig. 4, Table 5, Fig. 9). This suggests that fresh vegetation comprises a minor fraction of riverine POC, and biomarkers for vegetation observed in Figs. 5-7 instead trace inputs of degraded vegetation flushed into the river through soil organic carbon reservoirs.

The composition of Tapajós POC, very N-enriched (high N:C) and ¹³C-depleted, clearly falls off the mixing line in Fig. 9a, reflecting dominant input of *in situ* production, consistent with prior observations of Tapajós River POC composition (Mortillaro et al., 2011; Kim et al., 2012; Ward et al., 2015). The combination of a relatively deep euphotic zone depth and slower water velocities in the Tapajós River, especially during high water in July, encourages the growth of phytoplankton and cyanobacteria (Mortillaro et al., 2011). The highly depleted and anomalously low ACL fatty acids at the tributary surface (Figs. 6a; 7a,d,g) indicate that lower chain length lipids produced by aquatic phytoplankton (Cranwell, 1982) contribute much more to POC in this tributary and are

effectively absent in POC from the main stem and upstream (the Solimões and Madeira rivers).

4.3 Radiocarbon distribution of POC flux at Óbidos

The bulk and compound-specific analyses of riverine POC do not quantify the soil sources of POC to the Amazon main stem. Data generated by ramped oxidation of suspended sediment samples from various depths of the main stem at Óbidos (Table 1) and the surface of the Tapajós, Solimões and Madeira Rivers offer a unique opportunity to partition POC quantitatively among distinct sources from the drainage basin (Fig. 4).

To interpret thermograms, we assume that organic matrices span a range of activation energies (E_a) of decomposition, which causes different biomolecules to oxidize at different times during a controlled temperature ramp (Burnham and Braun, 1999; Cramer, 2004; Rosenheim et al., 2008; Hemingway, 2017 thesis). Thus, more thermally stable components of POC, which oxidize and release CO₂ at higher temperatures, have higher activation energies of decomposition. With this model, we transform thermograms to a distribution of activation energies, according an inverse algorithm coded in Python (Hemingway, 2016; Hemingway, 2017 thesis). The software deconvolves the calculated distribution of activation energies into a series of Gaussian distributions and reconstructs a thermogram as CO₂ concentration against temperature by summing these individual component distributions.

When transforming and deconvolving thermograms, users can customize a "smoothing factor" that limits the sensitivity of the Gaussian fit to features in the E_a distribution and therefore the number of components that the algorithm fits. The same "smoothing factor" was applied to all sample thermogram in this sample set to resolve a similar number and spread of components. The exception was the thermogram of the 2 m suspended sediment sample from the July 2014 Óbidos profile, to which we applied a lower "smoothing factor", derived from an optimization algorithm in the package, in order to deconvolve a similar number components to the other thermograms. Peaks that overlapped significantly in E_a space were combined before thermogram reconstruction. Although all of these parametrizations introduce user bias to the analysis, we justify these

choices by the fact that all sample thermograms exhibit a strikingly similar CO_2 distribution across temperatures and thus should be analyzed under similar statistical constraints (i.e., the number of peaks fitted and recombined to construct thermograms in temperature space).

Furthermore, the isotopic composition (δ^{13} C, F_m) of each deconvolved component can be calculated by inverting the following matrix, also described in Rosenheim et al. (2008):

In Equation 7, *i* is the number of CO₂ fractions trapped and analyzed, *j* is the number of components deconvolved, $\delta^{13}C_i$ is the stable isotope composition of each fraction (measured by IRMS), $\delta^{13}C_{Fj}$ is the isotopic composition of the component F and $f_{i,j}$ is the fraction of CO₂ measurement *i* in component *j*. The sum of all fractions in the same row of the matrix equals 1. Altogether, this approach allows us to interpret thermal stability distributions using several metrics: (1) the number of components per thermogram, (2) the mean activation energy and maximum temperature (T_{max}) of each component, (3) the contribution of each component to the total carbon (assumed to be ~100% organic carbon according to Section 3.4), and (4) the isotopic composition ($\delta^{13}C$ and F_m) of each component.

Nine thermograms of suspended sediment samples, all analyzed untreated, were deconvolved using this approach. The thermogram of suspended sediments collected from the Madeira River was not deconvolved because, as illustrated in Fig. 4, we cannot trust CO₂ concentration measurements beyond 500°C. The deconvolutions yielded three major components throughout all other thermograms, which from hereon forth we call components F_1 , F_2 and F_3 (Table 5). The root mean square error of the peak fits ranged from 7.5 E-04 to 5.0 E-03. The T_{max} of the two lower temperature components varies little across all nine samples: $299 \pm 10^{\circ}$ C (average ± 1 standard deviation) for F_1 and 404 $\pm 12^{\circ}$ C for F_2 . The fractions of total carbon that these peaks represent are relatively constant across samples too: $66 \pm 4\%$ for F_1 and $25 \pm 6\%$ for F_2 . Only the Óbidos 2005 sample thermogram yielded a low F_1 fraction of 57% (Fig. 4a). The difference in the Óbidos 2005 thermogram deconvolution could have resulted from longer river water

sample storage times prior to filtration (Section 2.2), prolonging *in situ* degradation and/or primary production, which would have shifted the POC composition in the sample relative to the POC composition at the time of sampling. The relative consistency of these components across all 2014 Óbidos samples suggest that ~2/3 of the POC flux from the main stem in April and July 2014, corresponding to F₁, consists of younger, more ¹³Cdepleted soil-derived POC (F_m= 0.90 ± 0.07). ~1/4 of the total POC, corresponding to F₂, consists of an older, more degraded and ¹³C-enriched source (F_m= 0.75 ± 0.12).

The highest activation energy and temperature component, F₃, is the most variable across thermogram deconvolutions, with an average T_{max} of $511 \pm 20^{\circ}C$ and fractional contribution of $9 \pm 7\%$ across samples. The relatively high standard deviations can be attributed to inherently higher errors associated with resolving components with a lower fractional contribution to total sample carbon. In fact, for four thermograms, a fourth, even higher activation energy peak was resolved by deconvolution, which comprised 0.5 - 4% of the total sample carbon (Table 5; Fig. 10c-d). But, because this four-peak fit yields stable isotope values that vastly deviate from the expected δ^{13} C range of riverine POC, we have combined the third and fourth peak, F₃' and F₄, into the component we report as F₃. The fractional contribution of this fourth peak to total sample carbon is also very variable because fitting this peak depends on the highest temperature to which CO_2 data was collected. For instance, during ramped oxidation of the Óbidos 2005 sample, the gas analyzer measured concentrations of CO_2 released until 795°C, which extends beyond the temperature range of several other thermograms for which the fourth peak could not be resolved (Figs. 4). Taken together, these factors would account for some of the observed variability in component F₃ fractions and T_{max} values among thermograms that resolve F₄ and thermograms that do not.

Because the properties of F_1 and F_2 are very similar across Óbidos sample thermograms, and because methodological artifacts impacted the deconvolution of F_3 vs. $F_{3'}$, we suggest that this fourth component is likely present in all samples from the main stem. Further, we posit that F_4 represents radiocarbon-dead petrogenic organic carbon, which is expected to be most thermally recalcitrant (Galy et al., 2008; Rosenheim and Galy, 2012) (Fig. 4a). Much of the petrogenic organic carbon source to the Amazon River likely derives from the Andes, where rapid flushing of steep bedrock occurs (Clark et al., 2013; Bouchez et al., 2014). It is possible that some of the CO_2 evolved at higher temperatures in the thermograms derives from charring products of intrinsically lower thermal stability POC in the sample (Williams et al., 2014). Nonetheless, the small fraction we propose to be F_4 is consistent with the sharp decrease in F_m values with increasing temperature observed across thermogram gas fractions, and independent estimates of petrogenic organic carbon in suspended sediments at Óbidos (Bouchez et al., 2014).

Assigning an $F_{m,F4}$ value of 0 in the four thermograms that resolve F_4 (Table 5; Fig. 10c-d), the proportion of F_4 prior to combination with $F_{3'}$ can be used to calculate the F_m value of $F_{3'}$ (Equation 8). These calculated $F_{m,3'}$ values range from 0.37-0.50 in Obidos samples, similar to the original F_3 values, and the fractional contribution of F_4 to total sample POC ($f_{F_{\tau}}$ in Equation 8) ranges from 0.5% to 4%. Again, this relatively wide range results from the Óbidos 2005 thermogram deconvolution, which yielded anomalously large F_{3} , and F_{4} components (22% and 4% of total carbon, respectively), but is still consistent with estimates of petrogenic organic carbon content reported by Bouchez et al. (2014). Fraction $F_{3'}$ has a higher F_m value of 0.60 in the Solimões sample, where the fractional contribution of F₄ is 2%. Because δ^{13} C values for petrogenic carbon are variable and only weakly constrained in the Amazon basin (Bouchez et al., 2014), we refrain from using the same mass balance approach to calculate the δ^{13} C value of F₃'. But, Bouchez et al. (2014) estimated that petrogenic organic carbon in the Amazon River Basin would have a δ^{13} C of ~-25‰. Considering the consistent enrichment of CO₂ with temperature across the thermograms (Fig. 4b), it is likely that F_4 would be more enriched than three F3 values resolved from the Obidos sample thermograms (-34.6‰ to -25.9‰; Table 5), which agrees with previous estimates.

$$F_{m,F3} = F_{m,F3'} \frac{f_{F3'}}{f_{F3'} + f_{F4}} + F_{m,F4} \frac{f_{F4}}{f_{F3'} + f_{F4}}$$
(8)

Altogether, the deconvolutions of Óbidos sample thermograms can be used to apportion POC flux at Óbidos into three soil organic matter pools and a petrogenic organic carbon pool. The component proportions F_1 , F_2 , and F_3/F_3 , across thermograms amount to fluxes of ~240-360 kg/second of relatively young and recently degraded soil POC, 90-130 kg/s and 30-50 kg/s of two older soil pools that are more ¹³C-enriched with age, respectively. The highest thermal stability and more fractionally variable component F_4 amounts to a flux between 2 and 22 kg/s of petrogenic organic carbon (calculated from the range of F_4 fractions resolved from three main stem thermograms) at Óbidos between April and July 2014.

4.4 Differentiating soil organic matter sources to riverine POC

In general, bulk and RPO fraction F_m values trace a negative relationship with bulk and RPO fraction δ^{13} C values from Óbidos and tributary samples (Fig. 9b), indicating that bulk POC in most of these samples consists of a mixture of young, low activation energy ¹³C-depleted soil organic matter and old, high activation energy ¹³Cenriched soil pools. Component F_m values also tend to decrease as component δ^{13} C values increase (Table 5). Fig. 9a illustrates a weak but significant correlation between bulk N:C and $\delta^{13}C_{POC}$ values measured in suspended sediments at Óbidos, also highlighting a mixing relationship between a fresher, low N:C, and ¹³C-depleted endmember and a more degraded, high N:C and ¹³C-enriched end-member (p<0.05, r²=0.28), consistent with the thermogram data. This is consistent with perspective that soil organic matter is a mixture of distinct pools at different stages of degradation, and thus exhibits a range of radiocarbon ages (Plante et al., 2013).

Two non-mutually exclusive hypotheses could explain the compositional variations in the thermally distinct soil POC sources to riverine POC (Table 5; Fig. 9). First, main stem POC could reflect mixing of soil sources from different landscapes that bear distinct $\delta^{13}C_{POC}$ signals and degradation histories. The drainage basin includes both ^{13}C -depleted, C3 landscapes in the low-lying floodplain (most of Amazonia) and more ^{13}C -enriched C4 and higher altitude Andean landscapes traversed by the western tributaries, including the Solimões and Madiera rivers (Fig. 1a, O'Leary, 1981; Körner et al., 1991). Isotopic differences among the Solimões River, Madeira River and main stem RPO gas fractions provide evidence for differences among soil age distributions across geographical space (Fig. 9b).

Mixing of geographically and isotopically diverse soil sources in the main stem could explain the slight differences in the bulk POC composition observed at Óbidos samples between sampling months. Lower average molar C:N ratios of 8.7 ± 0.88 in

April, compared to 10.7 ± 1.2 in July (p<0.05, Student's t-test), as well as significantly less depleted bulk $\delta^{13}C_{POC}$ values in April (-28.2 ± 0.1‰) than in July (-28.8 ± 0.2‰), suggest a small shift in soil source between river stages. The lower bulk $\delta^{13}C_{POC}$ and N:C values throughout the July profile (Table 3; Fig. 9a) are consistent with increased drainage of the adjacent floodplain during receding water levels, as the floodplain and varzea lakes are dominated by soils replete with the residual organic carbon from C3 plants (Moreira-Turcg et al., 2013). By contrast, the data imply that main stem POC in April, during rising waters, is influenced by a greater proportion of POC from the upstream tributaries, including POC transported by the Solimões and Madiera Rivers to the main stem, which are more ¹³C-enriched and N-enriched (Table 3; Fig. 9a). Increased incidence of landslides in the Andes during the months leading up to April, i.e., the Amazon rainy season, would increase the input of high altitude ¹³C-enriched POC from these tributaries to Óbidos (Clark et al., 2013; Clark et al., 2016). The higher N:C of POC in this sampling month would also imply that these sources are more degraded (Ometto et al., 2006; Cleveland and Liptzin, 2007), consistent with lower bulk and RPO gas fraction F_m values observed for the Solimões and Madeira river samples. Thus, we posit that preferential transport of the more degraded pools of this Andean soil source to Óbidos would explain the both more degraded and ¹³C-enriched signatures of POC observed in April (Feng et al., 2016; Mayorga et al., 2005). Note that these bulk differences do not manifest in observable differences in thermogram shape and the isotope composition of RPO fractions or components among the Óbidos samples (Fig. 4, Table 5).

Second, main stem POC could also comprise a mixture of soils from different depth horizons rather than from different landscapes. This hypothesis provides a strong mechanistic explanation for concurrent decreases in F_m and increases in $\delta^{13}C$ with increasing thermal stability (Figs. 4, 9a-b), but does not exclude the influence of the first hypothesis described above. This would be expected because $\delta^{13}C$ of soil organic matter gets more enriched with depth across the Amazon River Basin (De Camargo et al., 1999; Ehleringer et al., 2000; Ometto et al., 2006; Appendix C, Fig. S1). Thus, more thermally stable pools of POC in the Óbidos thermograms would derive from deeper, more degraded and ¹³C-enriched soil pools. This hypothesis could similarly explain the temporal shifts in bulk $\delta^{13}C$ and C:N values of POC at Óbidos, particularly the greater input of ¹³C-depleted, higher C:N and less degraded soil POC in July, as the river draws more carbon from the superficial layers of the floodplain. Input of this fresher floodplain-derived organic matter to the river in July is consistent with accumulation of higher F_m POC, possibly from coarser and less degraded vegetation debris, in the deeper sections of the river (Table 3, Fig. 2).

4.5 Bioreactivity of POC in the main stem

Sections 4.2-4.4 show that the Amazon River predominantly mobilizes organic carbon from longer residence time reservoirs in soils and bedrock, rather than from the more actively recycled reservoirs of living/aboveground terrestrial biomass. The mass of POC ultimately transported to the ocean, ~12-14 Tg POC/yr (Richey et al. 1990; Galy et al., 2015), is the mass difference between the quantity of soil and petrogenic POC flushed into the river and heterotrophic respiration of this POC in the river, which equates to a CO_2 outgassing flux of ~500 Tg/yr (Richey et al., 1990; Richey et al. 2002; Mayorga et al. 2005). Observations that certain pools of riverine POC contribute disproportionately to *in situ* oxidation (Richey et al., 1990; Mayorga et al., 2005; Ward et al. 2013) have led to the hypothesis that preferential degradation of organic matter plays an important role in this balance, and thus the river basin's role in the global organic carbon cycle (Cole et al., 2007).

The lack of any modern or bomb ¹⁴C-influenced component in riverine suspended POC at Óbidos implies that either younger sources of POC (e.g., fresh vegetation) are rapidly decomposed in the river over other POC sources (Mayorga et al., 2005), or that these fresher inputs to the river system are insignificant, perhaps because they are rapidly degraded in soils prior to being flushed into the river (e.g., Feng et al., 2016). The similarity in thermograms of POC between Óbidos and the two major tributaries upstream (Solimões and Madeira rivers) supports this argument, as the two upstream thermograms also show a lack of a significant modern POC component across RPO fractions. However, that the isotopic differences between the upstream and Óbidos sample thermograms could reflect turnover of older soil-derived components from

different soil horizons or geographical sources (Section 4.4) between the Solimões River mouth, Madeira River mouth and Óbidos.

If we consider thermal stability a proxy for diagenetic stability or biological reactivity (Rosenheim et al., 2008; Rosenheim and Galy, 2012), the invariant thermograms of samples from Óbidos suggest that the diagenetic reactivity of POC in the main stem is relatively homogenous throughout the Amazon main stem (Fig. 4). Hydrodynamic sorting does not impose an observable control on the recycling of carbon within the river cross-section. This perspective is supported by lack of depth-dependent variation in bulk and compound-specific metrics for POC composition (Figs. 2, 5-7), and invariant enzyme activities throughout the Óbidos cross-section in July 2014 (Fig. 8; Appendix C, Fig. S3) that suggest that heterotrophic activity on POC remains constant with depth (e.g., Steen and Arnosti, 2013; Thao et al., 2015). At the same time, it is worth considering whether the three thermally distinct components of riverine POC at Óbidos have different fates downstream of Óbidos. The sample thermograms suggest that ~2/3 of POC flux from the Amazon River main stem, derived from a less degraded soil pool, is more likely to degrade in transit towards or within the coastal ocean water column than the other third.

Finally, the lack of change in thermogram shape among samples upstream and downstream of Óbidos would suggest that mixing of isotopically distinct POC from different tributaries does not alter the thermal stability of POC in the Amazon main stem. Alternatively, the similarity in sample thermogram shape could be a methodological limitation of the extent of degradation that thermal stability can resolve. Perhaps organic matter pools beyond a specific degradation state generate invariant thermograms despite isotopic differences in the CO₂ evolved during ramped oxidation. It is particularly perplexing that the Tapajós POC thermogram is similar to the upstream sample thermograms, except that it lacks a high temperature (petrogenic OC) component, while the isotopic composition of gas fractions indicates that *in situ* production must dominate the CO₂ evolved (Fig. 4; Fig. 9). At the same time, bulk POC in the Tapajós River is not modern (F_m = 0.88) and even-over-odd predominance observed across higher chain-length fatty acids (Fig. 7g) points to a significant input of vegetation-derived POC to the river. As POC concentrations are so low in the Tapajós River (0.08 mg/L), DOC leached from

degraded soils in the drainage basin and sorbed onto particles produced by *in situ* production could noticeably influence the apparent bulk F_m and fatty acid distribution in POC. Enough sorbed DOC could even shift the thermal stability of the sample via matrix interactions between the older DOC and fresher POC in the sample (see Chapter 4 Section 4.2; Williams et al., 2014), causing the thermogram to resemble the others in Fig. 4.

5 Conclusions

Combining ramped oxidation with bulk and compound-specific analyses of POC composition has enabled us to quantify three thermally and isotopically distinct soilderived components and one relatively small petrogenic organic carbon-derived component of POC exported by the Amazon River main stem. The data show that 2/3 of POC flux at Óbidos is less thermally stable and younger, presumably representing a less degraded soil pool, than the other two soil pools, suggesting that biomolecules from this fraction are more likely to remineralize between Óbidos and the coastal Atlantic Ocean. The small highest thermal stability petrogenic POC component in the river flux would be least likely to degrade. As petrogenic POC most likely originates from the Andean-influenced Solimões and Madeira river basins (Clark et al., 2013; Bouchez et al., 2014), its presence at Óbidos suggests that it persists between the confluence of these upstream tributaries with the main stem and Óbidos, and may continue to persist in route to the Atlantic Ocean.

However, thermal stability does not necessarily equate to diagenetic stability. Williams et al. (2015) analyzed the ¹⁴C distribution of POC stored mudbank sediments from the French Guiana region of the Amazon River Basin shelf by ramped pyrolysis and found that petrogenic organic carbon was absent in these sediments, suggesting that this pool of old terrestrial organic matter turns over between the floodplain and the coastal ocean. Ward et al. (2013) also suggested that even lignins, considered the most recalcitrant terrestrial biomolecules (Gough et al., 1993), are actively remineralized during river transport across the basin. Thus, perspectives of environmental reactivity and selective degradation vs. preservation in the Amazon need review, perhaps with the unique perspectives and analytical opportunities offered by ramped oxidation.

Looking to the future, RPO analyses of riverine POC collected over a higherresolution transect upstream and downstream of Óbidos will be essential to constrain the different sources of soil POC we observe across the thermograms from Óbidos, assess their reactivity through the river system, and resolve the paradox of identical thermograms across isotopically distinct POC sources in the current data set. Finally, as the Amazon landscape shifts in response to climate change and land use change policies ((Davidson et al., 2012; Gloor et al., 2013; Brienen et al., 2013), export of sediments and organic carbon to the river system are expected to change (Butman et al., 2015; Dias et al., 2015), as well. RPO offers a tool to assess whether these transitions would manifest in changes in the quantity, quality and recycling of soil organic carbon flushed into the river system.

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Table 1. Sample locations and times. Depth-specific samples at Óbidos were collected in three positions across the main stem channel, referenced to the right bank across from Óbidos (Fig. 1). The 2005 data are taken from Bouchez et al. (2011b; 2014). Discharge, total suspended solid (TSS) and particulate organic carbon (POC) fluxes, and water temperatures are provided when possible, as well. z_R , or Rouse numbers, were calculated by fitting TSS concentration profiles in each sampling month to Equation 3, following Bouchez et al. (2011b).

River	Location	Latitude	Longitude	Date	Channel position	ample depth	emperature	Vol.	Discharge	zR	TSS flux	POC flux
						Ś	-					4.2
		°S	°W		km	m	$^{\circ}C$		m^3/s	unitless	kg/s	kg/s
main stem	Óbidos	1.9367*	55.503*	6/8/05	-	0	no data	>100 L	128,000	0.39	61,300	604
Madeira	Foz Madeira	3.450*	58.808*	6/6/05	-	0	no data	>100 L	21,800	0.09	17,400	168
Solimões	Manacapuru	3.314*	60.554*	6/4/05	-	0	no data	>100 L	124,700	0.10	1,300	14
main stem	Óbidos	1.946	55.510	4/2/14 10:30	0.32	35.49	25.9	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.946	55.510	4/2/14 10:30	0.32	15.3	27.3	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.946	55.510	4/2/14 10:30	0.32	2.1	no data	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.942	55.503	4/2/14 10:30	1.1	<u>49.1</u>	no data	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.942	55.503	4/2/14 10:30	1.1	<u>30.15</u>	27.5	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.942	55.503	4/2/14 10:30	1.1	13.33	26.8	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.942	55.503	4/2/14 10:30	1.1	<u>2.7</u>	26.4	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.938	55.496	4/2/14 10:30	2.00	27.8	27.5	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.938	55.496	4/2/14 10:30	2.00	10.4	26.1	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.938	55.496	4/2/14 10:30	2.00	2.8	26.1	~10 L	246,000	0.26	53,000	540
main stem	Óbidos	1.94	55.501	4/1/14 10:30	-	0	28.0	>100 L	246,000	N/A	53,000	540

Tapajós	Tapajós	no data	no data	7/27/14 13:00	-	0	no data	>100 L	no data	N/A	no data	no data
Tapajós	Tapajós	no data	no data	7/27/14	-	bed	no data	N/A	no data	N/A	no data	no data
main stem	Óbidos	1.946	55.509	7/28/14 14:00	0.34	42.8	28.9	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.945	55.510	7/28/14 14:00	0.34	20.6	no data	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.945	55.509	7/28/14 14:00	0.34	3.5	no data	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.944	55.494	7/28/14 14:00	1.6	<u>54</u>	28.9	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.942	55.497	7/28/14 14:00	1.6	39.83	29	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.941	55.499	7/28/14 14:00	1.6	<u>30.4</u>	28.9	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.941	55.499	7/28/14 14:00	1.6	14.95	no data	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.939	55.501	7/28/14 14:00	1.6	<u>3.14</u>	no data	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.933	55.497	7/28/14 14:00	2.2	53.06	29	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.931	55.499	7/28/14 14:00	2.2	29.83	no data	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.933	55.498	7/28/14 14:00	2.2	3.33	no data	~10 L	247,000	0.34	48,000	370
main stem	Óbidos	1.942	55.496	7/28/14 13:00	-	0	29.2	>100 L	247,000	0.34	48,000	370
main stem	Óbidos	no data	no data	7/29/14	-	bed	no data	N/A	247,000	N/A	48,000	370
main stem	Óbidos	no data	no data	7/29/14	-	bed	no data	N/A	247,000	N/A	48,000	370
main stem	Óbidos	no data	no data	7/29/14	1	flood	no data	N/A	247,000	N/A	48,000	370
*=approximate	; 											
bolded = samp	mples analyz les that were	ed by ramp lipid extrac	ed oxidatic	on.								
"bed"=bedload	l.; "flood"=fl	oodplain de	posit.									

Table 2. Substrates used to measure enzyme activities associated with suspended sediments collected within the Óbidos cross-section in July 2014. AMC is -amino-4-methylcoumarin and MUB is 4-methylumbelliferone. Table credit: Andrew Steen.

Substrate	Abbreviation	Enzyme	Element
leucine-AMC	Leu-AMC	leucyl aminopeptidase	Ν
Phenylalanine-AMC	Phe-AMC	phenyalanyl aminopeptidase	Ν
Ala-Ala-Phe-AMC MUB-beta-N-acetyl	AAF-AMC	chymotrypsin	Ν
glucosamine	MUB-NAG	N-acetylglucosaminidase	Ν
MUB-cellobiose	MUB-cello	cellulase	С
MUB-beta glucose	MUB-beta-glu	beta-glucosidase	С
MUB-beta xylose	MUB-xyl	xylanase	С
MUB-PO4	MUB-PO4	phosphomonoesterase	Р

Table 3. Total suspended sediment concentration, organic matter composition and grain size of all depth-specific samples collected at Óbidos in 2014, large-volume samples, and bed /floodplain samples. Errors for C:N and $\delta^{13}C_{POC}$ are reported. Relative errors of depth-specific F_m values are < 3%, and are not reported. Refer to Table 1 for sampling times of individual samples.

Location	Date	Depth	Channel Position	[TSS]	%POC	Nd%	C:N	δ ¹³ C _{POC}	Fraction Modern	Grain- mean	Grain-d50	Grain-d10	Grain-d90
•							µmol/						
		m	km	mg/L	mg/mg	mg/mg	µmol	%0	unitless	μm	μm	μm	μm
Óbidos	6/8/05	0	-	no data	0.85 ± 0.01	0.14 ± 0.005	7.3 ± 0.5	-28.5 ± 0.05	0.71	no data	no data	no data	no data
Solimões	6/6/05	0	-	no data	1.1 ± 0.06	0.17 ± 0.003	7.9 ± 1.8	$\textbf{-27.0} \pm 0.05$	0.72	no data	no data	no data	no data
Madeira	6/4/05	0	-	no data	0.50 ± 0.01	$\textbf{0.08} \pm \textbf{0.0004}$	7.1 ± 1.9	$\textbf{-27.7} \pm 0.3$	0.67	no data	no data	no data	no data
main stem	4/1/14	0	-	55	1.4 ± 0.01	0.19 ± 0.004	$\textbf{8.5}\pm\textbf{0.2}$	-28.1 ± 0.06	0.72	no data	no data	no data	no data
main stem	4/2/14	35.49	0.32	319	0.98 ± 0.02	0.12 ± 0.001	$\textbf{9.6} \pm \textbf{0.2}$	$\textbf{-28.3}\pm0.07$	no data	44	29	4	109
main stem	4/2/14	15.3	0.32	150	1.0 ± 0.02	0.12 ± 0.001	9.6 ± 0.2	$\textbf{-28.3}\pm0.2$	no data	48	31	5	121
main stem	4/2/14	2.1	0.32	79	1.3 ± 0.02	0.20 ± 0.001	7.8 ± 0.1	$\textbf{-28.4} \pm 0.09$	no data	50	29	4	131
main stem	4/2/14	49.1	1.1	208	0.96 ± 0.01	0.13 ± 0.001	$\textbf{8.8} \pm \textbf{0.1}$	$\textbf{-28.2}\pm0.05$	0.77	63	44	6	153
main stem	4/2/14	30.15	1.1	233	0.93 ± 0.01	0.11 ± 0.001	10 ± 0.1	$\textbf{-28.2}\pm0.03$	0.78	48	30	5	119
main stem	4/2/14	13.33	1.1	111	1.4 ± 0.01	0.19 ± 0.002	$\textbf{8.4} \pm \textbf{0.1}$	$\textbf{-28.0} \pm 0.08$	0.78	43	27	4	106
main stem	4/2/14	2.7	1.1	72	1.5 ± 0.01	0.24 ± 0.004	7.4 ± 0.1	-28.1 ± 0.03	0.77	70	49	4	171
main stem	4/2/14	27.8	2.00	149	1.1 ± 0.01	0.15 ± 0.001	8.5 ± 0.1	-28.2 ± 0.1	no data	61	39	5	154
main stem	4/2/14	10.4	2.00	126	1.2 ± 0.05	0.16 ± 0.003	8.7 ± 0.4	$\textbf{-28.1}\pm0.02$	no data	60	43	6	142
main stem	4/2/14	2.8	2.00	107	1.2 ± 0.03	$\textbf{0.17} \pm \textbf{0.002}$	8.3 ± 0.3	-28.1 ± 0.1	no data	44	28	4	110
Tapajós	7/27/14	bed	-	N/A	0.14 ± 0.002	0.017 ± 0.0003	$\textbf{9.8}\pm\textbf{0.3}$	$\textbf{-29.4} \pm 0.2$	1.02	no data	no data	no data	no data
Tapajós	7/27/14	0	-	1	16 ± 0.13	2.7 ± 0.005	6.8 ± 0.1	-31.0 ± 0.1	0.89	no data	no data	no data	no data
main stem	7/28/14	0	-	16	1.8 ± 0.02	0.23 ± 0.003	$\textbf{9.0}\pm\textbf{0.1}$	-29.1 ± 0.06	0.77	no data	no data	no data	no data
main stem	7/28/14	42.8	0.34	741	0.79 ± 0.04	0.07 ± 0.002	14 ± 0.9	-28.5 ± 0.1	no data	80	69	11	165

main stem	7/28/14	20.6	0.34	135	1.0 ± 0.003	0.11 ± 0.002	11 ± 0.2	$\textbf{-28.8} \pm 0.03$	no data	71	52	9	163
main stem	7/28/14	3.5	0.34	103	$\textbf{0.85} \pm \textbf{0.02}$	0.10 ± 0.001	10 ± 0.3	-28.8 ± 0.03	no data	66	51	8	146
main stem	7/28/14	54	1.6	305	0.51 ± 0.02	0.05 ± 0.001	11 ± 0.5	$\textbf{-28.4} \pm 0.2$	0.80	106	96	15	212
main stem	7/28/14	39.83	1.6	182	0.54 ± 0.01	0.06 ± 0.002	9.8 ± 0.3	-28.6 ± 0.1	0.78	80	66	9	176
main stem	7/28/14	30.4	1.6	169	0.77 ± 0.02	0.09 ± 0.001	11 ± 0.3	$\textbf{-28.9} \pm 0.05$	0.78	77	65	9	168
main stem	7/28/14	14.95	1.6	53	1.1 ± 0.02	0.12 ± 0.004	10 ± 0.4	$\textbf{-29.1} \pm 0.05$	0.77	60	45	7	139
main stem	7/28/14	3.14	1.6	44	1.3 ± 0.02	0.16 ± 0.002	9.4 ± 0.2	$\textbf{-28.9} \pm 0.06$	0.74	55	38	6	133
main stem	7/28/14	53.06	2.2	286	0.61 ± 0.01	0.07 ± 0.001	11 ± 0.3	-28.6 ± 0.1	no data	93	80	13	194
main stem	7/28/14	29.83	2.2	108	1.0 ± 0.01	0.11 ± 0.001	11 ± 0.2	$\textbf{-28.8} \pm 0.07$	no data	71	50	8	171
main stem	7/28/14	3.33	2.2	56	1.3 ± 0.02	0.14 ± 0.0003	11 ± 0.2	$\textbf{-28.9} \pm 0.07$	no data	62	42	7	153
main stem	7/29/14	flood	0	N/A	0.50 ± 0.03	0.05 ± 0.001	12 ± 0.9	$\textbf{-28.1}\pm0.08$	no data				
main stem	7/29/14	bed	0	N/A	0.06 ± 0.002	0.01 ± 0.0004	7.5 ± 0.5	-26.6 ± 0.09	0.60	no data	no data	no data	no data
main stem	7/29/14	bed	0	N/A	0.02 ± 0.001	BDL	no data	-26.9 ± 0.02	0.87	no data	no data	no data	no data

BDL= below detection limit. "bed"=bedload.; "flood"=floodplain deposit.

Table 4. Stable δ^{13} C and fraction modern (F_m) of CO₂ released within specific temperature intervals during two ramped oxidation analyses of the deepest (54 m) suspended sediment sample collected at Óbidos in July 2014. In the first run, the sample was untreated; in the second run, the sample was fumigated in concentrated hydrochloric acid, rinsed in Milli-Q water and freeze-dried prior to analysis. The mass-weighted sums of the δ^{13} C and F_m values across fractions were compared to the bulk δ^{13} C_{POC} and F_{m,POC} values, which were measured after acid fumigation only (no rinsing). Carbon yield = total µmol CO₂ gas quantified in RPO fractions/µmol expected from the bulk % OC and sediment mass inserted into RPO system.

Treatment	Temperature Interval	% total C	Fraction Modern	δ ¹³ C	
	$^{\circ}C$	%	unitless	%0	
Fumigation	116-277°C	24%	$\textbf{0.89} \pm \textbf{0.002}$	$\textbf{-30.7} \pm 0.1$	
+	277-332°C	28%	0.86 ± 0.002	$\textbf{-29.5}\pm0.1$	
Rinsing	332-394°C	24%	0.79 ± 0.002	$\textbf{-28.3}\pm0.1$	
	394-807°C	24%	0.53 ± 0.002	$\textbf{-28.0} \pm 0.1$	
	Weighted	Weighted	Bulk	Bulk	C Yield
	sum F _m	sum δ ¹³ C	F _{m,POC}	$\delta^{13}C_{POC}$	(%)
	0.77	-29.1	0.80	-28.4	86
Treatment	Temperature Interval	% total C	Fraction Modern	δ ¹³ C	
Untreated	142-276°C	22%	0.94 ± 0.002	-31.9 ± 0.1	
	276-325°C	30%	0.92 ± 0.002	-30.0 ± 0.1	
	325-449°C	36%	0.80 ± 0.002	-28.1 ± 0.1	
	449-704°C	12%	0.44 ± 0.002	-25.9 ± 0.1	
	Weighted	Weighted	Bulk	Bulk	C Yield
	sum F _m	sum δ ¹³ C	F _{m,POC}	$\delta^{13}C_{POC}$	(%)
	0.82	-29.3	0.80	-28.4	106 (100)

Table 5. Stable and radiocarbon isotope composition of deconvolved components from thermograms of all samples underlined in Table 1, except Madeira River POC. No deconvolved stable isotope data are available for samples where CO_2 gas fractions were lost prior to analysis. T_{max} is the temperature of each component peak maximum. In general, the deconvolution algorithm yielded at least three major components (F1, F2, F3) across all thermograms. In four samples, the algorithm divided F3 between two components, F3' and F4 (bolded), yielding a four-component fit. Refer to Table 1 for sampling times of individual samples.

					Fraction of		
Location	Month	Depth	Peak	Tmax	Total C	δ ¹³ C	Fm
	mm-yyyy	т		°C	%	%0	unitless
main stem	06 2005	0	Fl	308	57%	-31.8 ± 0.1	0.80 ± 0.002
main stem	06 2005	0	F 2	385	21%	-27.2 ± 0.2	0.79 ± 0.01
main stem	06 2005	0	F3	483	22%	-28.9 ± 0.1	0.36 ± 0.003
main stem	06 2005	0	F3'	N/A	18%	N/A	$\boldsymbol{0.37 \pm 0.004}$
main stem	06 2005	0	F4	N/A	4%	N/A	0
Solimões	06 2005	0	Fl	311	65%	$\textbf{-28.9} \pm 0.1$	0.80 ± 0.002
Solimões	06 2005	0	F2	397	21%	-26.8 ± 0.1	0.69 ± 0.004
Solimões	06 2005	0	F3	497	14%	-27.3 ± 0.1	0.51 ± 0.004
Solimões	06 2005	0	F3'	N/A	12%	N/A	0.60 ± 0.005
Solimões	06 2005	0	F4	N/A	2%	N/A	0
main stem	04 2014	49.1	F1	288	65%	no data	0.95 ± 0.01
main stem	04 2014	49.1	F2	404	30%	no data	0.86 ± 0.002
main stem	04 2014	49.1	F3	523	5%	no data	0.39 ± 0.003
main stem	04 2014	30.15	Fl	297	65%	no data	0.93 ± 0.002
main stem	04 2014	30.15	F2	404	29%	no data	0.68 ± 0.003
main stem	04 2014	30.15	F3	524	6%	no data	0.14 ± 0.006
main stem	04 2014	2.7	F1	296	70%	no data	0.93 ± 0.001
main stem	04 2014	2.7	F2	416	25%	no data	$\textbf{0.86} \pm \textbf{0.002}$
main stem	04 2014	2.7	F3	507	6%	no data	$\textbf{0.46} \pm \textbf{0.003}$
main stem	04 2014	2.7	F3'	N/A	5.4%	N/A	0.50 ± 0.003
main stem	04 2014	2.7	F4	N/A	0.5%	N/A	0
Tapajós	07 2014	0	F1	315	62%	-34.1 ± 0.1	$\textbf{0.94} \pm \textbf{0.003}$
Tapajós	07 2014	0	F2	421	35%	-31.1 ± 0.1	0.93 ± 0.004
Tapajós	07 2014	0	F3	548	2%	-36.4 ± 1.7	0.34 ± 0.03
main stem	07 2014	54	F1	293	66%	-31.1 ± 0.1	0.94 ± 0.002
main stem	07 2014	54	F2	391	25%	-26.6 ± 0.2	0.72 ± 0.004
main stem	07 2014	54	F3	500	10%	-25.9 ± 0.1	0.33 ± 0.004

main stem	07 2014	54	F3'	N/A	8%	N/A	0.38 ± 0.005
main stem	07 2014	54	F4	N/A	1%	N/A	0
main stem	07 2014	30.14	F1	288	70%	$\textbf{-30.8}\pm0.1$	0.94 ± 0.002
main stem	07 2014	30.14	F2	416	27%	-26.8 ± 0.3	0.58 ± 0.01
main stem	07 2014	30.14	F3	527	3%	-34.6 ± 3.5	0.17 ± 0.04
main stem	07 2014	3.14	F1	295	71%	no data	0.83 ± 0.001
main stem	07 2014	3.14	F2	399	15%	no data	0.62 ± 0.004
main stem	07 2014	3.14	F3	493	14%	no data	0.38 ± 0.004



Figure 1. (a) The Amazon River Basin and all field sites from June 2005, April 2014 and July 2014. Image credit: Paul Lefebvre and Greg Fiske (Woods Hole Research Center). (b-c) Monthly-averaged main stem discharge and water level time-series in 2014, compiled by the Brazilian Agencia Nacional de Aguas. The symbols represent monthly averages and the dashed lines represent linear interpolations between the monthly averages. The vertical solid lines represent the two sampling months, April and July. (d-e) Acoustic Doppler Current Profiler (ADCP) transects in latitude/longitude across the Amazon River at Óbidos in April and July 2014. Transects started at the right bank across from Óbidos, and ended at the left bank near Óbidos. The colorbar represents river depth (m). (f-g) Measured and extrapolated (Equation 1) water velocities within the cross-section at Óbidos in April 2014 and July 2014. The colorbar represents water velocity, 0-4 m/s.



Figure 2. Profiles of TSS concentration, organic matter composition, and median grain size (d50) through depth profiles at Óbidos in April and July 2014. The profiles for each position in the transect are plotted separately as triangles (near right bank), circles (near left bank) and crosses (near center). Refer to Table 1 and Fig. 1 for channel position and GPS coordinates of the profiles. All y-axes are normalized to the river depth at the channel position of each profile. The errors in %POC, %PN and fraction modern are relatively small and therefore not plotted. δ^{13} C values are expressed in units of ‰.


Figure 3. Fits of all depth-specific TSS concentration (C_z) profiles measured at Óbidos in 2014 to the Rouse Equation (Equation 3; Rouse, 1950; Bouchez et al., 2011b). Observed and modeled C_z/C_a values are plotted as a function of z/H. z is the height above the river bed H of each profile, and C_a is the surface-most measured C_z value of each profile.





Figure 4. Ten thermograms (solid lines) generated by ramped oxidation of untreated suspended sediments from the Amazon main stem, and the Tapajós, Solimões and Madeira Rivers (Table 1). One sample from a deeper section of the main stem was additionally analyzed following acid-treatment (dashed line). The left-hand y-axes for the thermograms are all normalized to the total CO₂ released during ramped oxidation analysis. The height of the bars indicates the (A) F_m and (B) δ^{13} C values (‰) of CO₂ trapped within distinct temperature intervals during ramped oxidation. The bottom-right panel in (A) shows all thermograms, excluding the Madeira River thermogram, plotted on the same axes.



Figure 5. (a-b) δ^{13} C values (‰) and (c-e) abundances of straight chain *n*-alkanes measured at Óbidos (surface and depth-specific), and the Madeira and Solimões Rivers (surface). The horizontal lines in a-b represent the δ^{13} C value of the bulk POC, reported in Table 3.



Figure 6. Profiles of average chain length (ACL) calculated in higher chain length fatty acids (C_{24} - C_{34}) and *n*-alkanes (C_{25} - C_{33}) in all 2005 and 2014 surface samples and four depth-specific samples collected at Óbidos in 2014. Refer to Equation 5 for ACL calculations.



Figure 7. (a-c) δ^{13} C values (‰) and (d-i) abundances of straight chain fatty acids measured at Óbidos (surface and depth-specific), and the Madeira, Solimões and Tapajós Rivers. The horizontal lines in a-c represent the δ^{13} C values of the bulk POC, reported in Table 3.



Figure 8. Activities of eight enzymes measured on particles within the Óbidos crosssection in July 2014. Data analysis credit: Andrew Steen.



Figure 9. (a) Bulk N/C ratios as a function of $\delta^{13}C_{POC}$ (‰) in all samples collected in June 2005, April 2014 and July 2014. The correfer to location and/or time (Table 1; Fig. 1), while the symbols refer to sample type (circles for depth-specific, triangles for surface and diamonds for bed/floodplain samples). (b) F_m and $\delta^{13}C$ (‰) values of CO₂ collected and analyzed within specific temperature intervals during ramped oxidation of all samples shown in Fig. 4. The color bar represents starting temperature (°C) of each CO₂ g interval, with warmer colors indicating higher starting temperatures.

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Figure 10. Deconvolution of three suspended sediment thermograms. (a, c, e) Probability distribution function (PDF) of activation energy of CO_2 evolved (lines) and components (shaded peaks). In all examples, the sum of components (red line) fits well with the PDF of the raw thermogram (black line). (b, d, f) Fraction of CO_2 evolved/temperature as a function of temperature for components resolved in panels a, c and e. The reconstructed thermograms (red lines) also compare well to the original thermograms (black lines). Only three components were resolved in a and e, while four were resolved in c. In c, the third and fourth components were combined prior to reconstructing the thermogram in d. This range of deconvolution scenarios applied to the rest of the thermograms presented in Fig. 4.

Chapter 6: Conclusions

Because of its sheer size, the deep ocean carbon reservoir regulates the carbon cycle and climate over centuries to millennia. Our view of this reservoir as a long-term buffer against the fossil fuel CO₂ emissions relies on the assumption that it will remain unchanged in the Anthropocene (Steffen et al., 2011). However, climate change is likely to shift export and transfer efficiency of POC fixed by marine and terrestrial primary producers, which would directly impact the supply of carbon sequestered in this deep reservoir. It is difficult to detect changes in marine POC export fluxes, as large-scale, continuous monitoring of primary productivity over the ocean surface (i.e., by satellite) has been absent over most of the industrial era (Henson et al., 2010). However, Laufkötter et al. (2016) projected that export is likely to decrease in future climate change scenarios, and found that the magnitude and mechanism of change varied geographically. On land, climate change and landscape alteration by human development are not only expected to affect the total export of POC in rivers (e.g., Dias et al., 2015; Appendix C, Fig. S2), but also the source and composition of organic matter that is mobilized from land into the river system (Butman et al., 2015). In a large system like the Amazon River Basin, these changes could shift the balance between POC decomposition in the river and POC burial on the continental shelf (Langerwisch et al., 2016).

Thus, understanding the physical and biogeochemical mechanisms that control POC export and transfer to the deep ocean has bearing on our predictions of biological pump strength and river POC transport in future climate change scenarios. In this thesis, I have commented on the relative importance of different mechanisms that control POC transfer efficiency. The direction of the earliest chapter (chapter 2) led me to focus the remaining chapters (chapters 3-5) on the hypothesis that the quality of POC produced by different marine and terrestrial primary producers determines its fate during transport to the deep ocean. As characterizing POC by its quality and reactivity in the environment is challenging, much of the effort in these chapters was the search for a proxy that described both POC composition and reactivity. Consequently, this thesis further explores various applications of ramped pyrolysis/oxidation (RPO), an expanding technique for

characterizing POC composition and reactivity by thermal stability (Rosenheim et al., 2008; Rosenheim and Galy, 2012; Rosenheim et al., 2013).

Perspectives of marine and terrestrial organic matter recycling in water remain largely separate in scientific literature. In their recent overview paper, Burd et al. (2016) highlighted the common challenges that both research communities face in modeling "vertical transport" in the ocean and "lateral transport" in rivers. They highlight several priorities for development, such as improving empirical measurements and models of microbe-mediated organic matter transformations (e.g., enzyme kinetics), and characterizing the biomolecular composition of POC over different gradients in degradation. The combined RPO analyses in chapters 4 and 5 describe POC composition across a spectrum of marine and terrestrial samples, ranging from the freshest POC produced in the ocean's euphotic zone and in axenic phytoplankton cultures to more highly degraded, century to millennium year old POC from soil organic carbon reservoirs in the Amazon River Basin (Fig. 1). In the middle of this spectrum is mesopelagic zone POC, which is more degraded than euphotic zone POC, but not as degraded as soilderived POC in the Amazon River.

The direct link between the euphotic and mesopelagic zone samples in Chapter 4 provided an opportunity to highlight different processes that may transform POC during degradation in the ocean's water column. The three thermogram profiles imply that a similar series of processes drive selective accumulation of a middle activation energy pool of POC with depth and selective loss of a lower activation energy POC pool. Two of three thermogram depth profiles also exhibit selective loss of a higher activation energy POC pool with depth. These transformations occur regardless of the composition of POC produced at the surface by phytoplankton. But, with the small number of samples in the data set, it is difficult to further assess the influence of different phytoplankton assemblages on the quality and recycling of POC transferred to the deep ocean.

If increasing degradation state manifests in a lower spread of activation energies of decomposition in organic matter, we would hypothesize that the processes that consistently transform POC in the three Southern Ocean depth profiles would lead to an even narrower activation energy distribution in soil-derived POC from the Amazon (Fig. 1). The Chapter 5 thermograms fit these expectations (Fig. 2d-f). Because thermogram gas fractions in Amazon River POC exhibit a large range in radiocarbon age as well as stable isotope composition, providing one more dimension for thermogram comparison than in the chapter 4 thermograms (Fig. 2a-c), the data allowed for apportionment of riverine POC among different terrestrial organic matter sources that vary in thermal stability. One implication of this partitioning in the context of chapter 4 and Fig. 1 is that the youngest soil-derived source, which makes up $\sim 2/3$ of total POC flux in the main stem, is more likely to degrade between Óbidos and the continental shelf than the other higher thermal stability POC components in the river. The spectrum illustrated in Fig. 1 also suggests that thermogram "narrowness" may potentially serve as a better proxy for degradation state of organic matter than thermal stability itself (Rosenheim et al. 2008; Rosenheim and Galy 2012).

The data sets across marine and terrestrial environments highlight important future steps in the application of the RPO method to studying the fate of organic matter. First, future RPO experiments should be more controlled to minimize environmental variables that confound data interpretations. This approach would help constrain the identity of biomolecules that oxidize at different temperatures, and the real-world processes that accompany transformations in thermal stability. For example, RPO analyses of POC collected from marine algae cultures inoculated with bacteria and left to degrade through time would help isolate the effects of degradation on POC thermal stability. This approach could be coupled to measurements of bacterial vs. algae cell populations to tease apart the effects of algae biomass degradation from conversion of algae biomass to heterotrophic biomass. This biomass could even be placed in a rolling tank (Engel et al., 2009) to simulate particle sinking and possibly molecular cross-linking processes (Hedges et al., 2000; Freeman, 2001). Meanwhile, in the marine environment, collecting larger quantities of the sinking POC size class from sediment traps or McLane pumps would allow for analysis of the particles that contribute directly to the biological pump.

Further, it is surprising that Amazon River POC oxidizes across overall lower temperatures than marine POC, even though it is much older and presumably more degraded than the samples in chapter 4 (Fig. 2d-f). This distinction indicates that the thermal stability of marine and terrestrial POC may not be comparable in absolute terms, and that the link between thermal stability and environmental reactivity needs finetuning. Explanations for these differences in marine and terrestrial POC thermal stability are beyond the scope of this thesis, but include the influence of different source materials, (phytoplankton vs. fresh vegetation) different heterotroph communities (marine vs. soil bacterial biomass), and different mineral matrices. Higher spatial-resolution sampling of the Amazon River along a transect from upstream of Óbidos to the plume, or coastal Atlantic sediments, would provide a better link between the source of soil-derived organic matter in riverine POC, its reactivity in the river system and its thermal stability determined by RPO analysis.

Finally, the applications of RPO analysis to three different sample sets in this thesis compel us to consider directions for improvement in future applications of the method. Perhaps the most important challenge to overcome is the presence of particulate inorganic carbon (PIC) in samples from different environments. Chapters 4 and 5 have shown that acid fumigation and rinsing prior to RPO analysis not only hydrolyze PIC in a sample, but also remove specific pools of POC and/or shift the thermal stability distribution of remaining POC in the sample. At the same time, it is difficult to isolate PIC from thermograms of samples that have complex organic matrices without fumigation. Continuous measurements of δ^{13} C of CO₂ evolved during ramped oxidation may help to better isolate and correct for PIC in these sample mixtures by its ¹³C-enriched signature.

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Figure 1. Conceptual schematic of differences in the activation energy or thermal stability distribution of POC at three stages of degradation. As POC degradation state increases, its activation energy distribution narrows in thermograms, as expressed below by arrow length at the CO₂ concentration half-maximum of each thermogram (Appendix B, Fig. S4). Chapter 4 explored processes that may cause this decrease in activation energy range between fresh and mesopelagic zone POC, which potentially would further lead to differences between the mesopelagic zone and Amazon River POC.



Figure 2. (a-c) Thermograms (solid lines) and fraction modern values of CO₂ gas fractions (bar heights) collected during ramped oxidation of two Southern Ocean particle samples (a-b) and an Amazon River main stem sample (c). The variation in both δ^{13} C and F_m in the Amazon samples provide an additional dimension for comparing thermograms and POC source contributions to thermogram shape. (d-f) The thermal stability distributions of a euphotic zone sample thermogram (d), deeper mesopelagic zone sample thermogram (e), and an Amazon main stem sample thermogram (f). The arrows span the temperature range of the thermogram at half maximum CO₂ concentrations (Appendix B, Fig. S4).

Appendix A

Table S1.²³⁴Th activity and flux profiles estimated at 27 stations along cruises GB1 and GB2. Fluxes were estimated by measuring total seawater ²³⁴Th activity deficits relative to total seawater ²³⁸U activity, as described in Section 2.4. ²³⁴Th flux errors are propagated from ²³⁴Th activity errors.

			²³⁴ Th	²³⁴ Th Activity	²³⁸ U		²³⁴ Th Flux
Cruise	Station	Depth	Activity	Error	Activity	²³⁴ Th Flux	Error
-	-	т	dpm L ⁻¹	$dpm L^{-1}$	dpm L ⁻¹	$dpm m^{-2} d'$	$dpm m^{-2} d'$
GB1	6	13.6	1.04	0.02	2.36	681	27
GB1	6	22.3	1.28	0.05	2.36	924	31
GB1	6	29.2	1.12	0.03	2.36	1,243	34
GB1	6	40.2	1.23	0.03	2.36	1,503	37
GB1	6	45.2	1.29	0.06	2.36	1,702	39
GB1	6	53.1	1.53	0.04	2.36	2,096	48
GB1	6	78	1.86	0.12	2.36	2,437	100
GB1	6	100.6	1.90	0.05	2.36	2,684	106
GB1	6	115	2.08	0.10	2.36	2,804	116
GB1	6	130	2.15	0.05	2.36	2,911	121
GB1	6	150.3	2.09	0.07	2.36	3,182	145
GB1	6	200	2.04	0.18	2.37	3,419	195
GB1	16	11.1	0.99	0.02	2.35	548	20
GB1	16	17.07	1.10	0.02	2.35	762	22
GB1	16	23.04	0.94	0.02	2.35	1,107	25
GB1	16	34.2	1.61	0.04	2.35	1,311	30
GB1	16	42.2	1.61	0.05	2.36	1,611	42
GB1	16	62.3	1.97	0.05	2.36	1,933	71
GB1	16	100.17	2.06	0.05	2.36	2,277	105
GB1	16	140.8	2.18	0.06	2.37	2,487	139
GB1	16	180.1	2.24	0.08	2.37	2,636	178
GB1	16	220.5	2.21	0.21	2.37	2,819	308
GB1	16	260.7	2.37	0.13	2.37	2,810	394
GB1	16	340.3	2.16	0.08	2.37	3,040	409
GB1	25	10.3	1.70	0.05	2.45	283	26
GB1	25	16.1	1.95	0.06	2.45	380	30
GB1	25	23.8	1.91	0.03	2.45	525	34
GB1	25	34.5	2.14	0.05	2.47	611	38
GB1	25	41.7	2.07	0.07	2.47	773	51
GB1	25	80	2.42	0.03	2.48	941	74

GB1	25	115	2.46	0.04	2.45	937	91
GB1	25	140.5	2.38	0.05	2.43	975	103
GB1	25	165.2	2.32	0.07	2.44	1,074	125
GB1	25	200.2	2.44	0.04	2.42	1,029	177
GB1	25	300	2.36	0.03	2.38	1,051	195
GB1	32	12.15	1.96	0.08	2.48	229	41
GB1	32	18	1.96	0.06	2.48	327	44
GB1	32	25.25	2.00	0.07	2.49	475	51
GB1	32	39.2	2.02	0.10	2.49	614	61
GB1	32	46	2.05	0.19	2.49	808	104
GB1	32	69.67	1.83	0.04	2.47	1,304	116
GB1	32	100.2	2.33	0.03	2.48	1,435	127
GB1	32	130.5	2.38	0.03	2.47	1,520	141
GB1	32	171	2.41	0.09	2.46	1,581	186
GB1	32	210	2.43	0.03	2.46	1,612	198
GB1	32	251	2.29	0.11	2.45	1,818	250
GB1	32	300.8	2.52	0.03	2.44	1,761	253
GB1	38	20.4	2.36	0.08	2.52	127	72
GB1	38	32.2	2.17	0.03	2.52	247	74
GB1	38	44.2	2.20	0.04	2.52	418	83
GB1	38	69	2.28	0.10	2.53	550	102
GB1	38	81	2.27	0.08	2.54	638	107
GB1	38	92	2.34	0.03	2.53	743	113
GB1	38	121	2.43	0.05	2.51	809	126
GB1	38	150	2.41	0.09	2.50	886	151
GB1	38	175	2.57	0.05	2.50	811	168
GB1	38	220	2.43	0.06	2.50	911	206
GB1	38	280.2	2.47	0.05	2.48	938	247
GB1	38	350.7	2.74	0.08	2.48	678	265
GB1	46	10	1.40	0.02	2.43	395	20
GB1	46	16.8	1.47	0.03	2.43	573	23
GB1	46	23	1.54	0.03	2.43	813	28
GB1	46	35.7	1.41	0.02	2.42	1,090	32
GB1	46	42.03	1.37	0.02	2.42	1,338	34
GB1	46	52.1	1.36	0.03	2.42	1,652	38
GB1	46	62.7	1.74	0.07	2.42	2,123	69
GB1	46	100.4	2.44	0.03	2.41	2,089	102
GB1	46	150.4	2.51	0.05	2.40	1,937	146
GB1	46	200.4	2.46	0.05	2.40	1,850	174
GB1	46	250.2	2.37	0.11	2.39	1,874	283
GB1	46	325.3	2.53	0.12	2.38	1,712	314
GB1	59	10.4	1.46	0.03	2.34	335	21
GB1	59	16.1	1.43	0.06	2.34	493	25

GB1	59	22.5	1.64	0.21	2.34	673	61
GB1	59	34	1.61	0.12	2.34	857	69
GB1	59	40	1.50	0.03	2.34	1,013	70
GB1	59	47	1.49	0.05	2.34	1,256	73
GB1	59	60	1.49	0.09	2.34	1,844	102
GB1	59	95	2.12	0.05	2.35	2,108	128
GB1	59	140	2.05	0.06	2.36	2,509	159
GB1	59	185	2.24	0.09	2.37	2,673	202
GB1	59	225	2.16	0.05	2.37	3,113	244
GB1	59	325	2.34	0.04	2.40	3,189	262
GB1	70	17	1.86	0.06	2.36	314	47
GB1	70	27	1.85	0.06	2.36	460	52
GB1	70	37	1.76	0.05	2.36	721	61
GB1	70	57.5	1.81	0.06	2.36	967	69
GB1	70	68	1.90	0.05	2.37	1,119	72
GB1	70	80	2.05	0.05	2.37	1,266	80
GB1	70	100	2.36	0.06	2.38	1,280	94
GB1	70	125	2.29	0.18	2.38	1,361	190
GB1	70	160	2.35	0.06	2.39	1,412	208
GB1	70	200	2.40	0.09	2.40	1,414	248
GB1	70	250	2.45	0.09	2.40	1,342	286
GB1	70	300	2.27	0.06	2.41	1,439	292
GB1	77	17	1.71	0.06	2.35	399	45
GB1	77	26	1.68	0.05	2.35	573	49
GB1	77	35	1.84	0.05	2.35	794	57
GB1	77	56	1.79	0.06	2.35	1,047	66
GB1	77	66	1.86	0.06	2.35	1,197	70
GB1	77	77	1.97	0.06	2.36	1,391	79
GB1	77	101	2.25	0.08	2.37	1,485	105
GB1	77	130	2.30	0.06	2.39	1,561	122
GB1	77	160	2.31	0.06	2.39	1,630	139
GB1	77	190	2.16	0.05	2.40	1,903	162
GB1	77	240	2.40	0.07	2.40	1,917	213
GB1	77	300	2.40	0.13	2.41	1,922	243
GB1	85	12.6	1.30	0.05	2.35	476	32
GB1	85	19	1.55	0.07	2.35	629	35
GB1	85	26	1.41	0.06	2.35	925	43
GB1	85	41	1.52	0.07	2.35	1,197	52
GB1	85	49	1.76	0.07	2.35	1,340	56
GB1	85	58	1.91	0.07	2.35	1,493	63
GB1	85	73	1.86	0.08	2.35	1,858	94
GB1	85	110	2.17	0.09	2.36	2,034	133
GB1	85	140	2.34	0.09	2.36	2,059	168

GB1	85	180	2.36	0.09	2.37	2,076	207
GB1	85	220	2.46	0.09	2.38	1,931	273
GB1	85	300	2.60	0.10	2.39	1,683	300
GB1	92	10	1.68	0.07	2.34	239	32
GB1	92	15	1.61	0.07	2.34	365	35
GB1	92	22	1.54	0.07	2.34	571	41
GB1	92	33	1.78	0.04	2.34	709	43
GB1	92	39	1.65	0.05	2.34	837	45
GB1	92	46	1.59	0.04	2.34	1,053	48
GB1	92	59	1.59	0.06	2.34	1,639	77
GB1	92	100	2.65	0.07	2.35	1,332	117
GB1	92	130	2.28	0.06	2.36	1,396	134
GB1	92	160	2.41	0.09	2.36	1,339	170
GB1	92	200	2.42	0.06	2.37	1,248	232
GB1	92	302	2.29	0.06	2.38	1,391	259
GB1	101	13.6	1.57	0.04	2.36	405	33
GB1	101	22	1.46	0.04	2.36	611	35
GB1	101	29.5	1.58	0.05	2.36	867	42
GB1	101	45	1.61	0.05	2.36	1,133	49
GB1	101	54	1.74	0.06	2.37	1,296	52
GB1	101	63	1.89	0.06	2.37	1,484	61
GB1	101	81	1.96	0.07	2.37	1,763	82
GB1	101	110	2.24	0.07	2.38	1,878	107
GB1	101	140	2.43	0.05	2.37	1,826	122
GB1	101	171	2.43	0.05	2.37	1,774	135
GB1	101	200	2.44	0.05	2.37	1,641	184
GB1	101	300	2.20	0.04	2.37	1,883	205
GB1	109	11.1	1.58	0.03	2.39	318	23
GB1	109	16.2	1.69	0.04	2.39	445	25
GB1	109	23.7	1.62	0.04	2.39	672	31
GB1	109	36.5	1.77	0.04	2.40	854	35
GB1	109	44	1.76	0.04	2.48	1,158	44
GB1	109	66	2.03	0.04	2.40	1,326	53
GB1	109	76	1.99	0.04	2.40	1,524	61
GB1	109	100	2.17	0.04	2.39	1,648	71
GB1	109	115	2.19	0.04	2.40	1,737	76
GB1	109	130	2.43	0.05	2.40	1,722	83
GB1	109	150	2.40	0.05	2.40	1,719	94
GB1	109	175	2.40	0.05	2.40	1,719	97
GB1	117	10.5	1.80	0.04	2.43	243	24
GB1	117	16.3	1.69	0.03	2.43	377	26
GB1	117	23	1.99	0.04	2.43	484	30
GB1	117	33	1.77	0.04	2.41	659	34

GB1	117	42	1.95	0.04	2.41	758	37
GB1	117	48	1.86	0.04	2.42	918	41
GB1	117	62	1.87	0.04	2.43	1,177	50
GB1	117	80	2.08	0.04	2.44	1,346	58
GB1	117	95	2.31	0.05	2.44	1,404	65
GB1	117	110	2.46	0.05	2.44	1,394	71
GB1	117	125	2.61	0.05	2.43	1,290	82
GB1	117	150	2.52	0.05	2.43	1,258	86
GB2	5	20	1.37	6.03	2.47	948	5,206
GB2	5	40	1.70	0.08	2.47	1,403	5,206
GB2	5	61	1.84	0.07	2.47	1,768	5,206
GB2	5	71	1.89	0.15	2.48	2,015	5,206
GB2	5	80	2.18	0.07	2.47	1,889	5,207
GB2	5	90	2.56	0.07	2.47	1,862	5,207
GB2	5	100	2.52	0.08	2.48	1,850	5,207
GB2	5	111	2.54	0.08	2.47	1,832	5,207
GB2	5	120	2.44	0.09	2.47	1,849	5,207
GB2	5	150	2.50	0.07	2.48	1,821	5,208
GB2	5	201	2.82	0.14	2.47	1,058	5,218
GB2	5	300	2.71	0.09	2.46	695	5,220
GB2	27	20	1.70	0.04	2.34	554	53
GB2	27	40	1.56	0.04	2.34	1,114	69
GB2	27	70	1.68	0.06	2.34	1,637	92
GB2	27	85	1.53	0.24	2.34	2,047	155
GB2	27	95	1.97	0.05	2.34	1,828	159
GB2	27	105	2.20	0.06	2.35	1,869	160
GB2	27	115	2.21	0.08	2.35	1,910	162
GB2	27	125	2.28	0.06	2.35	1,931	164
GB2	27	135	2.27	0.05	2.35	1,972	168
GB2	27	160	2.18	0.06	2.35	2,130	181
GB2	27	200	2.41	0.06	2.36	2,063	205
GB2	27	250	2.69	0.08	2.36	1,826	216
GB2	36	20	1.83	0.04	2.34	434	55
GB2	36	40	1.86	0.06	2.34	639	65
GB2	36	50	1.82	0.08	2.34	824	73
GB2	36	65	1.91	0.05	2.34	981	77
GB2	36	75	2.19	0.08	2.35	1,035	84
GB2	36	90	2.48	0.06	2.35	988	89
GB2	36	100	2.29	0.08	2.35	1,034	117
GB2	36	125	2.04	0.07	2.35	1,372	148
GB2	36	150	2.23	0.06	2.35	1,161	169
GB2	36	175	2.42	0.06	2.35	1,110	178
GB2	36	200	2.39	0.05	2.36	1,077	194

GB2	36	250	2.58	0.07	2.36	923	202
GB2	43	20	1.89	0.04	2.33	383	55
GB2	43	40	1.90	0.08	2.33	632	76
GB2	43	60	1.93	0.08	2.33	896	99
GB2	43	85	2.06	0.07	2.34	1,034	108
GB2	43	95	2.03	0.05	2.34	1,147	111
GB2	43	110	2.17	0.24	2.34	1,221	153
GB2	43	125	2.25	0.05	2.34	1,258	156
GB2	43	140	2.25	0.10	2.34	1,308	169
GB2	43	165	2.45	0.06	2.34	1,233	177
GB2	43	190	2.35	0.06	2.35	1,230	186
GB2	43	220	2.33	0.07	2.35	1,247	200
GB2	43	250	2.68	0.09	2.36	1,108	205
GB2	53	20	1.87	0.08	2.34	407	80
GB2	53	40	1.88	0.14	2.34	668	117
GB2	53	60	1.99	0.05	2.34	915	126
GB2	53	90	1.98	0.05	2.34	1,120	132
GB2	53	100	2.27	0.08	2.34	1,139	134
GB2	53	110	2.45	0.06	2.34	1,109	136
GB2	53	120	2.40	0.06	2.34	1,056	151
GB2	53	170	2.38	0.11	2.35	1,013	220
GB2	53	210	2.52	0.06	2.35	827	237
GB2	53	250	2.30	0.06	2.36	860	241
GB2	63	20	1.88	0.09	2.34	394	88
GB2	63	40	1.94	0.08	2.34	625	102
GB2	63	60	1.95	0.08	2.34	848	115
GB2	63	80	1.92	0.05	2.34	1,089	121
GB2	63	100	2.12	0.13	2.34	1,187	135
GB2	63	110	2.20	0.08	2.34	1,229	138
GB2	63	120	2.25	0.10	2.34	1,285	151
GB2	63	130	2.31	0.10	2.34	1,316	174
GB2	63	150	2.37	0.09	2.35	1,267	191
GB2	63	175	2.38	0.09	2.35	1,248	204
GB2	63	200	2.32	0.14	2.36	1,292	262
GB2	63	250	2.50	0.09	2.37	1,198	272
GB2	73	20	1.80	0.06	2.35	433	63
GB2	73	35	1.79	0.06	2.35	675	71
GB2	73	50	1.94	0.09	2.35	823	79
GB2	73	60	1.96	0.07	2.35	966	85
GB2	73	75	2.30	0.11	2.37	997	99
GB2	73	90	2.43	0.09	2.38	977	108
GB2	73	105	2.63	0.07	2.39	822	120
GB2	73	120	2.58	0.12	2.39	677	156

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GB2	73	135	2.43	0.06	2.40	794	168
GB2	73	160	2.34	0.06	2.40	853	184
GB2	73	200	2.48	0.06	2.40	807	189
GB2	87	20	1.93	0.07	2.35	354	71
GB2	87	40	1.92	0.05	2.35	599	81
GB2	87	60	1.75	0.06	2.35	939	91
GB2	87	80	1.89	0.07	2.35	1,134	98
GB2	87	90	1.88	0.05	2.35	1,268	100
GB2	87	100	2.28	0.08	2.35	1,290	103
GB2	87	110	2.33	0.13	2.35	1,299	115
GB2	87	125	2.35	0.08	2.36	1,302	121
GB2	87	140	2.36	0.08	2.36	1,297	136
GB2	87	170	2.32	0.08	2.36	1,336	159
GB2	87	200	2.48	0.09	2.37	1,213	196
GB2	87	250	2.55	0.08	2.38	1,088	208
GB2	93	20	1.95	0.07	2.36	352	71
GB2	93	40	2.04	0.09	2.36	554	96
GB2	93	65	2.14	0.08	2.36	710	116
GB2	93	90	1.95	0.08	2.36	944	127
GB2	93	105	2.01	0.08	2.36	1,071	132
GB2	93	115	2.17	0.10	2.37	1,142	137
GB2	93	130	2.57	0.09	2.39	1,061	144
GB2	93	145	2.44	0.08	2.39	1,033	152
GB2	93	165	2.42	0.08	2.39	1,003	170
GB2	93	180	2.40	0.09	2.39	992	213
GB2	93	200	2.82	0.09	2.39	469	249
GB2	93	250	2.61	0.09	2.38	309	259
GB2	100	20	1.89	0.05	2.42	464	58
GB2	100	40	1.90	0.07	2.42	764	75
GB2	100	60	2.06	0.07	2.42	922	83
GB2	100	70	1.80	0.05	2.42	1,101	85
GB2	100	80	2.27	0.08	2.43	1,145	89
GB2	100	90	2.33	0.08	2.43	1,174	94
GB2	100	100	2.66	0.28	2.43	1,110	124
GB2	100	110	2.43	0.08	2.43	1,112	130
GB2	100	130	2.30	0.09	2.43	1,205	151
GB2	100	160	2.48	0.09	2.43	1,157	182
GB2	100	200	2.47	0.08	2.43	1,132	190
GB2	106	20	1.87	0.05	2.47	603	68
GB2	106	50	1.77	0.05	2.47	1,005	79
GB2	106	60	1.83	0.06	2.47	1,189	82
GB2	106	70	2.09	0.08	2.48	1,301	86
GB2	106	80	2.17	0.08	2.48	1,413	92

GB2	106	95	2.45	0.09	2.48	1,426	107
GB2	106	115	2.62	0.09	2.48	1,335	126
GB2	106	140	2.50	0.06	2.48	1,320	138
GB2	106	165	2.36	0.06	2.48	1,423	154
GB2	106	200	2.49	0.07	2.47	1,405	186
GB2	106	250	2.64	0.17	2.46	1,274	226
GB2	112	10	2.11	0.08	2.42	136	40
GB2	112	20	2.39	0.09	2.42	149	54
GB2	112	35	2.29	0.08	2.42	199	65
GB2	112	45	1.91	0.07	2.43	347	69
GB2	112	55	1.80	0.07	2.42	526	73
GB2	112	65	2.25	0.09	2.42	589	82
GB2	112	80	2.20	0.08	2.43	717	97
GB2	112	105	2.47	0.09	2.43	682	126
GB2	112	135	2.66	0.09	2.42	466	160
GB2	112	170	2.63	0.09	2.42	270	186
GB2	112	200	2.69	0.10	2.42	-39	223
GB2	112	250	2.66	0.10	2.42	-211	237
GB2	119	10	1.73	0.19	2.42	347	96
GB2	119	25	1.73	0.05	2.42	644	101
GB2	119	40	1.77	0.06	2.42	923	106
GB2	119	55	2.00	0.07	2.42	1,072	110
GB2	119	65	2.16	0.07	2.43	1,147	113
GB2	119	74	2.21	0.07	2.43	1,226	117
GB2	119	90	2.43	0.08	2.43	1,223	124
GB2	119	105	2.32	0.08	2.42	1,280	135
GB2	119	130	2.60	0.08	2.43	1,127	158
GB2	119	165	2.69	0.09	2.42	853	189
GB2	119	200	2.50	0.08	2.42	756	218
GB2	119	250	2.75	0.09	2.42	516	231

Table S2. ²³⁴Th and POC fluxes at $z_{Th/U}$, estimated at 27 stations along cruises GB1 and GB2. $z_{Th/U}$ is the depth where ²³⁴Th and ²³⁸U activities re-establish secular equilibrium. Calculations of POC flux from ²³⁴Th fluxes are described in Section 2.4. POC flux errors are propagated from ²³⁴Th flux, and POC:²³⁴Th errors.

Cruise	Station	Z _{Th/U}	²³⁴ Th Flux	²³⁴ Th Flux Error	>51µm POC: ²³⁴ Th	POC Flux	POC Flux Error
-	-	т	$dpm m^{-2} d^{-1}$	$dpm m^{-2} d^{-1}$	µmol dpm⁻¹	$mmol m^{-2} d^{-1}$	$mmol m^{-2} d'^{1}$
GB1	6	130	2,911	121	2.0	5.7	0.25
GB1	16	140.8	2,487	139	2.6	6.6	0.37
GB1	25	115	937	91	1.7	1.6	0.18
GB1	32	131	1,581	186	1.3	2.0	0.20
GB1	38	121	809	126	2.7	2.2	0.35
GB1	46	100	2,089	102	2.8	5.9	0.32
GB1	59	95	2,108	128	2.9	6.1	0.42
GB1	70	100	1,280	94	4.3	5.5	0.44
GB1	77	100	1,485	105	6.0	9.0	1.3
GB1	85	140	2,059	168	5.9	12	1.0
GB1	92	100	1,332	117	3.8	5.1	0.47
GB1	101	140	1,826	122	1.8	3.3	0.23
GB1	109	130	1,722	83	1.0	1.6	0.09
GB1	117	110	1,394	71	1.0	1.4	0.09
GB2	5	90	1,862	5,207	1.3	2.5	6.9
GB2	27	105	1,869	160	1.9	3.5	0.32
GB2	36	90	988	89	2.0	2.0	0.18
GB2	43	125	1,258	156	3.7	4.6	0.58
GB2	53	100	1,139	134	3.9	4.4	0.53
GB2	63	130	1,316	174	4.6	6.1	0.81
GB2	73	75	997	99	4.9	4.9	0.49
GB2	87	100	1,290	103	2.9	3.8	0.33
GB2	93	130	1,061	144	1.2	1.2	0.18
GB2	100	90	1,174	94	2.8	3.3	0.27
GB2	106	95	1,426	107	0.9	1.2	0.10
GB2	112	105	682	126	0.8	0.6	0.11
GB2	119	90	1,223	124	2.5	3.0	0.31



А





С



Figure S1. Profiles of >51 μ m POC:²³⁴Th, PIC:²³⁴Th and BSi:²³⁴Th above 400 m. Black open circles represent measurements. Colored circles represent values at three possible depths (colored lines; see legend) in the POC:²³⁴Th panels, and at z_{PAR} (blue line) for the PIC:²³⁴Th and BSi:²³⁴Th panels. These values were interpolated when there were no measurements at these depths (refer to Tables 2 and 3 for specific stations). At station GB2-106, the BSi:²³⁴Th interpolation calculation excluded the anomalously low value at 80 m. Error bars shown are the propagated errors of >51 µm [POC], [PIC], [BSi] and particulate ²³⁴Th activity measurements. Note that "negative" values are below the instrument detection limit (see Section 2.3), and are equal to 0 within error.



Figure S2. (A) Diatom cell counts integrated from surface to z_{PAR} (cells m⁻²) as a function of >51 µm [BSi] at z_{PAR} . (B) Integrated coccolithophore cell counts as a function of >51 µm [PIC] at z_{PAR} . Outliers for >51 µm [BSi] and [PIC] at stations GB1-85 and GB1-16, respectively, are defined according to Chauvenet's Theorem (Glover et al., 2011). Total euphotic zone cell counts of diatoms and coccolithphores are significantly correlated to >51 µm [BSi] and [PIC] at z_{PAR} , respectively. The significant linear relationships are plotted here as blue lines, with corresponding p and r² values indicated. The regressions remain significant (p<0.05) even when excluding the outliers GB1-85 (Fig. S2a) and GB1-16 (Fig. S2b) from analysis (black lines). We note that cell counts are only available for the <36 µm size-fraction because of the methodology used for enumeration (see Supplemental Methods), while the biomineral measurements are of the >51 µm [BSi] and [PIC] at z_{PAR} do scale with the abundance of diatoms and coccolithophores in the euphotic zone of the water column.



Figure S3. The natural log of the ratio of >51 μ m [PIC]:[BSi] at z_{PAR} as a function of the natural log of the ratio of integrated coccolithophore: diatom cell counts in the euphotic zone. The significant linear relationship is plotted as a blue line, with a corresponding p and r² value indicated. This further supports the application of >51 μ m size-fraction biomineral concentrations at z_{PAR} as a proxy for describing euphotic zone ecosystem composition in Section 4.7 and Fig. 10. Despite the different size-fractions that are represented by the biomineral measurements and the cell counts (see Supplemental Methods), the significant correlation nonetheless supports the use of >51 μ m biomineral concentration ratios to describe the proportional abundance of certain phytoplankton types.



Figure S4. Percentage of total cells that are diatoms at z_{PAR} as a function of the %>51 μ m [POC] at z_{PAR} . The significant linear relationship is plotted as a blue line, with a corresponding p and r² value indicated. This relationship shows that FlowCAM® measurements of diatom abundance support our interpretation that the size fractionation of POC (%>51 μ m [POC] at z_{PAR}) determined from in-situ pump particle measurements (Section 2.2) reflects the relative abundance of diatoms. We note that there is no significant relationship between relative coccolithophore abundance and the size fractionation of POC.

Supplementary Methods for Figs. S2-S4

The FlowCAM® imaging cytometer enumerated nano- and microplankton cells from 10 mL Niskin cast samples at all McLane pump stations except GB2-36, GB2-27 and GB2-119 (refer to Table 1 for station locations) (Poulton and Martin, 2010). Moreover, at stations GB1-38 and GB1-70, cell counts were missing at z_{PAR} , and only measured at depths above and below z_{PAR} . The size range of counts was 5.6-35.5 µm. While particles >36 µm (up to 200 µm) could be seen in the FlowCAM®, they were rare in 10 mL samples, such that their normalized abundance could not be accurately calculated. Total diatom cell counts in the euphotic zone were approximated by summing FlowCam® -derived diatom concentrations (cells/mL) at all depths above z_{PAR} :

total cell counts =
$$\sum_{1}^{\text{end}} \text{mean}\left[\frac{\text{cells}}{\text{mL}_{n}}, \frac{\text{cells}}{\text{mL}_{n-1}}\right] \times [z_{n} - z_{n-1}]$$

where n is the measurement index number from the surface depth at n=1 downward towards z_{PAR} at n="end". The unit for this summation is equivalent to cells m⁻². Coccoliths and plated cells in the same Niskin samples were counted by birefringence microscopy (Balch et al., 2011). Total coccolithophore counts in the euphotic zone (cells m⁻²) were similarly estimated by summing the microscopy-based concentrations at all depths above z_{PAR} .

Supplementary References

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- Poulton, N. J., and Martin, J. L. Imaging flow cytometry for quantitative phytoplankton analysis - FlowCAM. In: Karlson B, Cusack C, Bresnan E (eds) Microscopic and molecular methods for quantitative phytoplankton analysis, Vol Chapter 8. Intergovernmental Oceanographic Commission of UNESCO, Paris, France, 49-54, 2010.
Appendix B



Figure S1. Quantity of PIC-derived CO₂ released by acid hydrolysis of one Great Calcite Belt sample ($<51 \mu m$) over ~ 25 hours.



Figure S2. Cell concentrations over four time points during a six day incubation of *E. huxleyi*, strains CCMP373 (panel A) and DHP624 (panel B), in f/50 media (Guillard, 1975; Harvey et al., 2015). All cell counts are significantly higher than the blank, non-inoculated f/50 media.



Figure S3. Two thermograms generated by ramped oxidation of the same Great Calcite Belt sample. RPO instrument conditions were kept as constant as possible between analyses. The y-axis is the CO₂ concentration normalized to total CO₂ evolved during the entire temperature ramp. The total CO₂ evolved was 54 μ mol in the first analysis and 30 μ mol in the second analysis.



Figure S4. Examples of three metrics illustrated in Fig. 11, calculated just for the surfacemost POC-rich sample from GB2-43 (top three panels) and the deepest sample from GB2-43 (bottom three panels). The thermograms are blank-corrected and the y-axis is CO₂ concentration normalized to the QMA filter active area, 125 cm². The first column exemplifies the calculation of the temperature range of the entire thermogram half maximum CO₂ concentration (Fig. 11a-c). The second column demonstrates calculation of the temperature range of the thermogram half maximum CO₂ concentration below 420°C (Fig. 11d-f). The third row illustrates calculation of the proportion of total CO₂ evolved within a middle temperature range, 355°C to 420°C (Fig. 11g-i).

Supplementary References

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- Harvey, E. L., Bidle, K. D., and Johnson, M. D.: Consequences of strain variability and calcification in Emiliania huxleyi on microzooplankton grazing, Journal of Plankton Research, 37, 1137-1148, 2015.

Appendix C

Table S1. Abundances and δ^{13} C values of straight-chain *n*-alkanes (C₁₉-C₄₁) in large-volume (>100 L) and depth-specific (10 L) samples collected in June 2005, April 2014 and July 2014. Refer to Table 1 in chapter 5 for sampling times of individual samples.

			Chain		Abundance		δ ¹³ C
Location	Date	Depth	Length	Abundance	Error	δ ¹³ C	Error
			#	ng/µg POC	ng/µg POC	‰	‰
Main stem	7/28/14	0	19	2.87E-03	2.88E-04	-32.5	0.1
Main stem	7/28/14	0	20	4.12E-03	4.14E-04	no data	no data
Main stem	7/28/14	0	21	5.16E-03	5.18E-04	-31.2	0.8
Main stem	7/28/14	0	22	4.86E-03	4.87E-04	no data	no data
Main stem	7/28/14	0	23	6.74E-03	6.76E-04	no data	0.6
Main stem	7/28/14	0	24	4.71E-03	4.72E-04	no data	no data
Main stem	7/28/14	0	25	9.36E-03	9.40E-04	-31.4	0.4
Main stem	7/28/14	0	26	7.33E-03	7.36E-04	no data	no data
Main stem	7/28/14	0	27	1.53E-02	1.53E-03	-31.1	0.1
Main stem	7/28/14	0	28	9.55E-03	9.59E-04	no data	no data
Main stem	7/28/14	0	29	3.70E-02	3.72E-03	-33.1	0.0
Main stem	7/28/14	0	30	1.15E-02	1.15E-03	no data	no data
Main stem	7/28/14	0	31	3.97E-02	3.99E-03	-33.4	0.1
Main stem	7/28/14	0	32	1.12E-02	1.13E-03	no data	no data
Main stem	7/28/14	0	33	2.00E-02	2.01E-03	-32.6	0.2
Main stem	7/28/14	0	34	7.02E-03	7.05E-04	no data	no data
Main stem	7/28/14	0	35	6.09E-03	6.11E-04	-31.0	0.3
Main stem	7/28/14	0	36	3.00E-03	3.01E-04	no data	no data
Main stem	7/28/14	0	37	3.69E-03	3.71E-04	-30.1	2.0
Main stem	7/28/14	0	38	2.39E-03	2.40E-04	no data	no data
Main stem	7/28/14	0	39	1.97E-03	1.98E-04	-31.6	1.8
Main stem	7/28/14	0	40	1.39E-03	1.39E-04	no data	no data
Main stem	7/28/14	0	41	1.01E-03	1.01E-04	no data	no data
Main stem	4/1/14	0	19	1.09E-02	1.09E-03	-28.3	0.2
Main stem	4/1/14	0	20	1.53E-02	1.53E-03	no data	no data
Main stem	4/1/14	0	21	1.16E-02	1.16E-03	-28.9	0.1
Main stem	4/1/14	0	22	1.14E-02	1.15E-03	no data	no data
Main stem	4/1/14	0	23	1.28E-02	1.28E-03	-29.6	0.2
Main stem	4/1/14	0	24	9.95E-03	9.97E-04	no data	no data
Main stem	4/1/14	0	25	1.21E-02	1.21E-03	-29.8	0.1
Main stem	4/1/14	0	26	9.98E-03	1.00E-03	no data	no data
Main stem	4/1/14	0	27	1.58E-02	1.59E-03	-30.3	0.4

Main stem	4/1/14	0	28	1.04E-02	1.04E-03	no data	no data
Main stem	4/1/14	0	29	3.26E-02	3.27E-03	-33.1	0.3
Main stem	4/1/14	0	30	1.07E-02	1.07E-03	no data	no data
Main stem	4/1/14	0	31	3.38E-02	3.39E-03	-33.7	0.2
Main stem	4/1/14	0	32	9.28E-03	9.29E-04	no data	no data
Main stem	4/1/14	0	33	1.67E-02	1.67E-03	-32.4	0.3
Main stem	4/1/14	0	34	4.97E-03	4.98E-04	no data	no data
Main stem	4/1/14	0	35	6.05E-03	6.06E-04	-31.2	0.4
Main stem	4/1/14	0	36	3.25E-03	3.26E-04	no data	no data
Main stem	4/1/14	0	37	3.37E-03	3.38E-04	-31.0	0.3
Main stem	4/1/14	0	38	2.50E-03	2.50E-04	no data	no data
Main stem	4/1/14	0	39	2.14E-03	2.14E-04	-31.3	0.7
Main stem	4/1/14	0	40	1.71E-03	1.72E-04	no data	no data
Main stem	4/1/14	0	41	1.33E-03	1.34E-04	-34.1	0.7
Main stem	6/8/05	0	19	2.09E-02	2.09E-03	no data	no data
Main stem	6/8/05	0	20	1.21E-02	1.21E-03	no data	no data
Main stem	6/8/05	0	21	1.10E-02	1.10E-03	no data	no data
Main stem	6/8/05	0	22	1.40E-02	1.40E-03	no data	no data
Main stem	6/8/05	0	23	1.71E-02	1.71E-03	no data	no data
Main stem	6/8/05	0	24	1.10E-02	1.11E-03	no data	no data
Main stem	6/8/05	0	25	1.59E-02	1.59E-03	no data	no data
Main stem	6/8/05	0	26	1.01E-02	1.01E-03	no data	no data
Main stem	6/8/05	0	27	1.95E-02	1.95E-03	no data	no data
Main stem	6/8/05	0	28	1.78E-02	1.79E-03	no data	no data
Main stem	6/8/05	0	29	4.07E-02	4.08E-03	no data	no data
Main stem	6/8/05	0	30	2.45E-02	2.46E-03	no data	no data
Main stem	6/8/05	0	31	4.15E-02	4.16E-03	no data	no data
Main stem	6/8/05	0	32	1.61E-02	1.62E-03	no data	no data
Main stem	6/8/05	0	33	2.14E-02	2.14E-03	no data	no data
Main stem	6/8/05	0	34	7.28E-03	7.30E-04	no data	no data
Main stem	6/8/05	0	35	9.55E-03	9.57E-04	no data	no data
Main stem	6/8/05	0	36	2.00E-03	2.00E-04	no data	no data
Main stem	6/8/05	0	37	3.28E-03	3.29E-04	no data	no data
Main stem	6/8/05	0	38	1.55E-03	1.56E-04	no data	no data
Main stem	6/8/05	0	39	2.74E-03	2.74E-04	no data	no data
Main stem	6/8/05	0	40	2.00E-03	2.00E-04	no data	no data
Main stem	6/8/05	0	41	1.65E-03	1.65E-04	no data	no data
Solimões	6/4/05	0	19	8.29E-03	9.40E-04	no data	no data
Solimões	6/4/05	0	20	5.56E-03	6.31E-04	no data	no data
Solimões	6/4/05	0	21	5.76E-03	6.53E-04	no data	no data
Solimões	6/4/05	0	22	7.86E-03	8.91E-04	no data	no data
Solimões	6/4/05	0	23	1.12E-02	1.27E-03	no data	no data
Solimões	6/4/05	0	24	8.57E-03	9.72E-04	no data	no data

Solimões	6/4/05	0	25	1.81E-02	2.05E-03	no data	no data
Solimões	6/4/05	0	26	1.86E-02	2.11E-03	no data	no data
Solimões	6/4/05	0	27	2.95E-02	3.35E-03	no data	no data
Solimões	6/4/05	0	28	2.59E-02	2.94E-03	no data	no data
Solimões	6/4/05	0	29	4.58E-02	5.19E-03	no data	no data
Solimões	6/4/05	0	30	2.09E-02	2.37E-03	no data	no data
Solimões	6/4/05	0	31	3.80E-02	4.31E-03	no data	no data
Solimões	6/4/05	0	32	1.11E-02	1.26E-03	no data	no data
Solimões	6/4/05	0	33	1.67E-02	1.89E-03	no data	no data
Solimões	6/4/05	0	34	3.20E-03	3.63E-04	no data	no data
Solimões	6/4/05	0	35	5.25E-03	5.95E-04	no data	no data
Solimões	6/4/05	0	36	1.35E-03	1.53E-04	no data	no data
Solimões	6/4/05	0	37	1.81E-03	2.05E-04	no data	no data
Solimões	6/4/05	0	38	7.15E-04	8.11E-05	no data	no data
Solimões	6/4/05	0	39	1.05E-03	1.19E-04	no data	no data
Solimões	6/4/05	0	40	6.93E-04	7.85E-05	no data	no data
Solimões	6/4/05	0	41	6.80E-04	7.71E-05	no data	no data
Madeira	6/6/05	0	19	2.13E-02	2.16E-03	no data	no data
Madeira	6/6/05	0	20	1.73E-02	1.76E-03	no data	no data
Madeira	6/6/05	0	21	1.78E-02	1.80E-03	no data	no data
Madeira	6/6/05	0	22	1.87E-02	1.90E-03	no data	no data
Madeira	6/6/05	0	23	2.25E-02	2.29E-03	no data	no data
Madeira	6/6/05	0	24	1.65E-02	1.67E-03	no data	no data
Madeira	6/6/05	0	25	2.25E-02	2.29E-03	no data	no data
Madeira	6/6/05	0	26	1.94E-02	1.97E-03	no data	no data
Madeira	6/6/05	0	27	3.19E-02	3.24E-03	no data	no data
Madeira	6/6/05	0	28	4.18E-02	4.25E-03	no data	no data
Madeira	6/6/05	0	29	7.80E-02	7.92E-03	no data	no data
Madeira	6/6/05	0	30	6.65E-02	6.75E-03	no data	no data
Madeira	6/6/05	0	31	7.74E-02	7.86E-03	no data	no data
Madeira	6/6/05	0	32	4.47E-02	4.54E-03	no data	no data
Madeira	6/6/05	0	33	3.43E-02	3.48E-03	no data	no data
Madeira	6/6/05	0	34	1.38E-02	1.41E-03	no data	no data
Madeira	6/6/05	0	35	9.87E-03	1.00E-03	no data	no data
Madeira	6/6/05	0	36	3.91E-03	3.97E-04	no data	no data
Madeira	6/6/05	0	37	3.89E-03	3.96E-04	no data	no data
Madeira	6/6/05	0	38	2.38E-03	2.42E-04	no data	no data
Madeira	6/6/05	0	39	2.75E-03	2.79E-04	no data	no data
Madeira	6/6/05	0	40	2.36E-03	2.40E-04	no data	no data
Madeira	6/6/05	0	41	1.16E-03	1.18E-04	no data	no data
Main stem	7/28/14	54	19	1.15E-03	1.27E-04	no data	NaN
Main stem	7/28/14	54	20	1.57E-03	1.73E-04	no data	NaN
Main stem	7/28/14	54	21	2.68E-03	2.95E-04	-30.9	0.2

Main stem	7/28/14	54	22	2.33E-03	2.56E-04	-29.6	0.4
Main stem	7/28/14	54	23	2.75E-03	3.02E-04	-30.9	0.5
Main stem	7/28/14	54	24	2.72E-03	2.99E-04	-29.9	0.5
Main stem	7/28/14	54	25	4.67E-03	5.14E-04	no data	NaN
Main stem	7/28/14	54	26	4.42E-03	4.86E-04	-29.6	0.6
Main stem	7/28/14	54	27	6.18E-03	6.80E-04	-34.1	0.2
Main stem	7/28/14	54	28	3.07E-03	3.38E-04	-28.5	3.7
Main stem	7/28/14	54	29	1.62E-02	1.78E-03	-34.0	0.5
Main stem	7/28/14	54	30	3.26E-03	3.59E-04	-37.3	0.5
Main stem	7/28/14	54	31	1.57E-02	1.73E-03	-34.6	0.3
Main stem	7/28/14	54	32	2.79E-03	3.07E-04	no data	NaN
Main stem	7/28/14	54	33	6.95E-03	7.64E-04	-34.6	1.2
Main stem	4/2/14	30.15	19	1.67E-03	1.69E-04	no data	NaN
Main stem	4/2/14	30.15	20	2.22E-03	2.24E-04	no data	NaN
Main stem	4/2/14	30.15	21	3.07E-03	3.09E-04	-29.6	0.4
Main stem	4/2/14	30.15	22	3.02E-03	3.05E-04	-29.4	0.1
Main stem	4/2/14	30.15	23	3.62E-03	3.65E-04	-31.1	0.1
Main stem	4/2/14	30.15	24	2.61E-03	2.63E-04	-30.0	0.5
Main stem	4/2/14	30.15	25	3.84E-03	3.87E-04	-30.3	0.3
Main stem	4/2/14	30.15	26	3.50E-03	3.53E-04	-30.1	0.4
Main stem	4/2/14	30.15	27	6.71E-03	6.77E-04	-32.3	0.2
Main stem	4/2/14	30.15	28	3.07E-03	3.09E-04	-31.8	0.4
Main stem	4/2/14	30.15	29	1.86E-02	1.88E-03	-34.5	0.1
Main stem	4/2/14	30.15	30	3.47E-03	3.49E-04	-33.6	0.4
Main stem	4/2/14	30.15	31	1.86E-02	1.88E-03	-34.9	0.0
Main stem	4/2/14	30.15	32	2.81E-03	2.83E-04	-34.2	0.2
Main stem	4/2/14	30.15	33	8.04E-03	8.11E-04	-34.7	0.3
Main stem	4/2/14	2.7	19	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	20	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	21	3.59E-03	3.61E-04	no data	NaN
Main stem	4/2/14	2.7	22	2.85E-03	2.86E-04	no data	NaN
Main stem	4/2/14	2.7	23	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	24	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	25	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	26	9.37E-03	9.41E-04	no data	NaN
Main stem	4/2/14	2.7	27	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	28	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	29	5.26E-03	5.28E-04	no data	NaN
Main stem	4/2/14	2.7	30	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	31	6.35E-03	6.37E-04	no data	NaN
Main stem	4/2/14	2.7	32	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	2.7	33	0.00E+00	0.00E+00	no data	NaN
Main stem	4/2/14	49.1	19	1.58E-03	1.58E-04	no data	NaN

Main stem	4/2/14	49.1	20	2.36E-03	2.37E-04	no data	NaN
Main stem	4/2/14	49.1	21	2.85E-03	2.86E-04	-30.4	0.7
Main stem	4/2/14	49.1	22	2.92E-03	2.93E-04	-30.0	0.0
Main stem	4/2/14	49.1	23	3.48E-03	3.49E-04	-31.0	0.2
Main stem	4/2/14	49.1	24	2.45E-03	2.46E-04	-29.8	0.1
Main stem	4/2/14	49.1	25	4.22E-03	4.23E-04	-31.2	0.1
Main stem	4/2/14	49.1	26	4.74E-03	4.75E-04	-30.5	0.3
Main stem	4/2/14	49.1	27	6.24E-03	6.25E-04	-34.4	0.5
Main stem	4/2/14	49.1	28	2.68E-03	2.68E-04	-31.8	0.4
Main stem	4/2/14	49.1	29	1.60E-02	1.60E-03	-34.2	0.1
Main stem	4/2/14	49.1	30	2.88E-03	2.88E-04	no data	NaN
Main stem	4/2/14	49.1	31	1.69E-02	1.69E-03	-35.9	0.1
Main stem	4/2/14	49.1	32	2.05E-03	2.06E-04	-34.3	0.7
Main stem	4/2/14	49.1	33	6.70E-03	6.71E-04	-34.2	0.1

			Chain		Abundance		δ ¹³ C
Location	Date	Depth	Length	Abundance	Error	δ ¹³ C	Error
			#	ng/µg POC	ng/µg POC	‰	%0
Main stem	4/1/14	0	14	0.31171	0.031	-27.7	0.1
Main stem	4/1/14	0	16	2.4949	0.250	-29.2	0.2
Main stem	4/1/14	0	18	3.3937	0.340	-28.6	0.3
Main stem	4/1/14	0	20	0.097067	0.010	-29.2	0.2
Main stem	4/1/14	0	22	0.20945	0.021	-31.6	0.5
Main stem	4/1/14	0	24	0.14455	0.014	-28.8	0.4
Main stem	4/1/14	0	25	0.037764	0.004	no data	no data
Main stem	4/1/14	0	26	0.10798	0.011	-30.2	0.5
Main stem	4/1/14	0	27	0.024168	0.002	no data	no data
Main stem	4/1/14	0	28	0.091665	0.009	-32.0	0.5
Main stem	4/1/14	0	29	0.026517	0.003	no data	no data
Main stem	4/1/14	0	30	0.081735	0.008	-33.2	0.2
Main stem	4/1/14	0	31	0.021069	0.002	no data	no data
Main stem	4/1/14	0	32	0.061277	0.006	-33.5	2.3
Main stem	4/1/14	0	33	0.012159	0.001	no data	no data
Main stem	4/1/14	0	34	0.031169	0.003	-33.1	0.1
Main stem	7/28/14	0	14	0.1952	0.020	-33.0	0.0
Main stem	7/28/14	0	16	3.4245	0.344	-30.8	0.4
Main stem	7/28/14	0	18	3.9505	0.397	-29.2	0.2
Main stem	7/28/14	0	20	0.096981	0.010	-30.9	0.0
Main stem	7/28/14	0	22	2.3583	0.237	-34.2	0.5
Main stem	7/28/14	0	24	0.15264	0.015	-32.5	0.6
Main stem	7/28/14	0	25	0.037073	0.004	no data	no data
Main stem	7/28/14	0	26	0.13925	0.014	-33.0	0.4
Main stem	7/28/14	0	27	0.029239	0.003	no data	no data
Main stem	7/28/14	0	28	0.13231	0.013	-32.9	0.6
Main stem	7/28/14	0	29	0.037333	0.004	no data	no data
Main stem	7/28/14	0	30	0.13233	0.013	-33.0	0.2
Main stem	7/28/14	0	31	0.034962	0.004	no data	no data
Main stem	7/28/14	0	32	0.10498	0.011	-33.3	0.3
Main stem	7/28/14	0	33	0.021529	0.002	no data	no data
Main stem	7/28/14	0	34	0.054067	0.005	-32.8	0.2
Tapajós	7/27/14	0	14	2.2663	0.227	-36.7	0.1
Tapajós	7/27/14	0	16	23.249	2.333	-33.6	0.3
Tapajós	7/27/14	0	18	11.378	1.142	-29.7	0.3

Table S2. Abundances and δ^{13} C values of fatty acids (C₁₄-C₃₄) from large-volume (>100 L) and depth-specific (10 L) samples collected in June 2005, April 2014 and July 2014. Refer to Table 1 in chapter 5 for sampling times of individual samples.

Tapajós	7/27/14	0	20	0.33744	0.034	-34.1	0.6
Tapajós	7/27/14	0	22	0	0.000	-39.4	0.2
Tapajós	7/27/14	0	24	0.93488	0.094	-36.2	0.6
Tapajós	7/27/14	0	25	0.18406	0.018	no data	no data
Tapajós	7/27/14	0	26	0.62779	0.063	-35.5	0.7
Tapajós	7/27/14	0	27	0.069807	0.007	no data	no data
Tapajós	7/27/14	0	28	0.4066	0.041	-36.6	1.2
Tapajós	7/27/14	0	29	0.031529	0.003	no data	no data
Tapajós	7/27/14	0	30	0.095123	0.010	-35.5	0.2
Tapajós	7/27/14	0	31	0.040663	0.004	no data	no data
Tapajós	7/27/14	0	32	0.031174	0.003	-34.9	0.1
Tapajós	7/27/14	0	33	0.0079396	0.001	no data	no data
Tapajós	7/27/14	0	34	0.012265	0.001	no data	no data
Main stem	6/8/05	0	14	0.19	0.019	-35.9	1.6
Main stem	6/8/05	0	15	0.063	0.006	no data	no data
Main stem	6/8/05	0	16	0.871	0.087	-34.3	7.4
Main stem	6/8/05	0	17	0.018	0.002	no data	no data
Main stem	6/8/05	0	18	0.18	0.018	-31.0	1.5
Main stem	6/8/05	0	19	0.05	0.005	no data	no data
Main stem	6/8/05	0	20	0.049	0.005	-33.0	0.4
Main stem	6/8/05	0	21	0.047	0.005	no data	no data
Main stem	6/8/05	0	22	0.097	0.010	-37.5	0.8
Main stem	6/8/05	0	23	0.028	0.003	no data	no data
Main stem	6/8/05	0	24	0.175	0.018	-37.0	1.5
Main stem	6/8/05	0	25	0.035	0.004	no data	no data
Main stem	6/8/05	0	26	0.194	0.019	-36.5	1.6
Main stem	6/8/05	0	27	0.04	0.004	no data	no data
Main stem	6/8/05	0	28	0.223	0.022	-37.2	1.9
Main stem	6/8/05	0	29	0.049	0.005	no data	no data
Main stem	6/8/05	0	30	0.166	0.017	-35.3	1.4
Main stem	6/8/05	0	31	0.038	0.004	no data	no data
Main stem	6/8/05	0	32	0.099	0.010	-34.3	0.8
Main stem	6/8/05	0	33	0.023	0.002	no data	no data
Main stem	6/8/05	0	34	0.048	0.005	-34.2	0.4
Main stem	6/8/05	0	35	0.066	0.007	no data	no data
Solimões	06 2005	0	14	0.187	0.021	-34.9	2.1
Solimões	06 2005	0	15	0.046	0.005	no data	no data
Solimões	06 2005	0	16	0.885	0.100	-34.7	9.9
Solimões	06 2005	0	17	0.017	0.002	no data	no data
Solimões	06 2005	0	18	0.169	0.019	-30.3	1.9
Solimões	06 2005	0	19	0.012	0.001	no data	no data
Solimões	06 2005	0	20	0.049	0.006	-30.9	0.5
Solimões	06 2005	0	21	0.018	0.002	no data	no data

Solimões	6/4/05	0	22	0.087	0.010	-34.5	1.0
Solimões	6/4/05	0	23	0.031	0.004	no data	no data
Solimões	6/4/05	0	24	0.19	0.022	-34.1	2.1
Solimões	6/4/05	0	25	0.044	0.005	no data	no data
Solimões	6/4/05	0	26	0.316	0.036	-34.2	3.5
Solimões	6/4/05	0	27	0.046	0.005	no data	no data
Solimões	6/4/05	0	28	0.27	0.031	-35.4	3.0
Solimões	6/4/05	0	29	0.059	0.007	no data	no data
Solimões	6/4/05	0	30	0.203	0.023	-34.6	2.3
Solimões	6/4/05	0	31	0.05	0.006	no data	no data
Solimões	6/4/05	0	32	0.141	0.016	-34.4	1.6
Solimões	6/4/05	0	33	0.032	0.004	no data	no data
Solimões	6/4/05	0	34	0.064	0.007	-34.5	0.7
Solimões	6/4/05	0	35	0.102	0.012	no data	no data
Madeira	6/6/05	0	14	0.089	0.009	-37.7	0.4
Madeira	6/6/05	0	15	0.024	0.002	no data	no data
Madeira	6/6/05	0	16	0.508	0.052	-37.6	2.5
Madeira	6/6/05	0	17	0.013	0.001	no data	no data
Madeira	6/6/05	0	18	0.099	0.010	-31.0	0.5
Madeira	6/6/05	0	19	0.006	0.001	no data	no data
Madeira	6/6/05	0	20	0.032	0.003	-31.2	0.2
Madeira	6/6/05	0	21	0.01	0.001	no data	no data
Madeira	6/6/05	0	22	0.05	0.005	-34.1	0.2
Madeira	6/6/05	0	23	0.018	0.002	no data	no data
Madeira	6/6/05	0	24	0.09	0.009	-35.5	0.4
Madeira	6/6/05	0	25	0.021	0.002	no data	no data
Madeira	6/6/05	0	26	0.085	0.009	-35.0	0.4
Madeira	6/6/05	0	27	0.018	0.002	no data	no data
Madeira	6/6/05	0	28	0.087	0.009	-35.0	0.4
Madeira	6/6/05	0	29	0.023	0.002	no data	no data
Madeira	6/6/05	0	30	0.075	0.008	-34.5	0.4
Madeira	6/6/05	0	31	0.018	0.002	no data	no data
Madeira	6/6/05	0	32	0.053	0.005	-33.7	0.3
Madeira	6/6/05	0	33	0.012	0.001	no data	no data
Madeira	6/6/05	0	34	0.027	0.003	-33.5	0.1
Madeira	6/6/05	0	35	0.034	0.003	no data	no data
Main stem	7/28/14	54	14	0.054832	0.006	-32.4	0.4
Main stem	7/28/14	54	16	0.63583	0.070	-30.4	0.2
Main stem	7/28/14	54	18	0.39584	0.044	-29.2	0.1
Main stem	7/28/14	54	20	0.036354	0.004	-30.6	1.1
Main stem	7/28/14	54	22	0.74563	0.082	-32.4	0.4
Main stem	7/28/14	54	24	0.086899	0.010	-31.5	0.3
Main stem	7/28/14	54	25	0.022954	0.003	no data	no data

Main stem	7/28/14	54	26	0.10196	0.011	-31.1	0.8
Main stem	7/28/14	54	27	0.023842	0.003	no data	no data
Main stem	7/28/14	54	28	0.12934	0.014	-31.5	0.1
Main stem	7/28/14	54	29	0.031932	0.004	no data	no data
Main stem	7/28/14	54	30	0.13955	0.015	-32.2	0.2
Main stem	7/28/14	54	31	0.028885	0.003	no data	no data
Main stem	7/28/14	54	32	0.094013	0.010	-32.8	0.3
Main stem	7/28/14	54	33	0.019822	0.002	no data	no data
Main stem	7/28/14	54	34	0.045886	0.005	-32.6	0.8
Main stem	4/2/14	30.15	14	0.070165	0.007	-29.9	1.3
Main stem	4/2/14	30.15	16	1.7788	0.179	-29.2	0.4
Main stem	4/2/14	30.15	18	1.7946	0.181	-28.8	0.3
Main stem	4/2/14	30.15	20	0.053737	0.005	-30.1	0.3
Main stem	4/2/14	30.15	22	0.2279	0.023	-31.5	0.6
Main stem	4/2/14	30.15	24	0.097517	0.010	-32.0	0.3
Main stem	4/2/14	30.15	25	0.023434	0.002	no data	no data
Main stem	4/2/14	30.15	26	0.097554	0.010	-32.0	0.3
Main stem	4/2/14	30.15	27	0.023332	0.002	no data	no data
Main stem	4/2/14	30.15	28	0.11045	0.011	-32.7	0.2
Main stem	4/2/14	30.15	29	0.029705	0.003	no data	no data
Main stem	4/2/14	30.15	30	0.11388	0.011	-33.3	0.2
Main stem	4/2/14	30.15	31	0.027346	0.003	no data	no data
Main stem	4/2/14	30.15	32	0.083013	0.008	-33.3	0.3
Main stem	4/2/14	30.15	33	0.015861	0.002	no data	no data
Main stem	4/2/14	30.15	34	0.038895	0.004	-33.4	0.9
Main stem	4/2/14	2.7	14	0.21517	0.022	no data	no data
Main stem	4/2/14	2.7	16	3.6379	0.365	no data	NaN
Main stem	4/2/14	2.7	18	3.4641	0.348	no data	NaN
Main stem	4/2/14	2.7	20	0.0699	0.007	no data	no data
Main stem	4/2/14	2.7	22	0.55703	0.056	no data	NaN
Main stem	4/2/14	2.7	24	0.078868	0.008	no data	NaN
Main stem	4/2/14	2.7	25	0.018353	0.002	no data	no data
Main stem	4/2/14	2.7	26	0.056486	0.006	no data	NaN
Main stem	4/2/14	2.7	27	0.022969	0.002	no data	no data
Main stem	4/2/14	2.7	28	0.050916	0.005	no data	NaN
Main stem	4/2/14	2.7	29	0.012918	0.001	no data	no data
Main stem	4/2/14	2.7	30	0.05081	0.005	no data	NaN
Main stem	4/2/14	2.7	31	0.011095	0.001	no data	no data
Main stem	4/2/14	2.7	32	0.034628	0.003	no data	NaN
Main stem	4/2/14	2.7	33	0.0078758	0.001	no data	no data
Main stem	4/2/14	2.7	34	0.016382	0.002	no data	NaN
Main stem	4/2/14	49.1	14	0.10537	0.011	-28.7	1.3
Main stem	4/2/14	49.1	16	3.1154	0.312	-29.4	0.3

Main stem	4/2/14	49.1	18	3.0324	0.304	-28.9	0.2
Main stem	4/2/14	49.1	20	0.064186	0.006	-30.1	0.3
Main stem	4/2/14	49.1	22	0.25316	0.025	-32.8	0.9
Main stem	4/2/14	49.1	24	0.090366	0.009	-32.1	0.3
Main stem	4/2/14	49.1	25	0.022487	0.002	no data	no data
Main stem	4/2/14	49.1	26	0.092227	0.009	-32.1	0.1
Main stem	4/2/14	49.1	27	0.024705	0.002	no data	no data
Main stem	4/2/14	49.1	28	0.10805	0.011	-32.7	0.2
Main stem	4/2/14	49.1	29	0.031813	0.003	no data	no data
Main stem	4/2/14	49.1	30	0.11688	0.012	-33.2	0.2
Main stem	4/2/14	49.1	31	0.031116	0.003	no data	no data
Main stem	4/2/14	49.1	32	0.08853	0.009	-33.7	0.2
Main stem	4/2/14	49.1	33	0.020959	0.002	no data	no data
Main stem	4/2/14	49.1	34	0.045168	0.005	-33.2	0.3

Table S3. Activities of eight enzymes measured on particles within the Óbidos crosssection in July 2014. Refer to chapter 5, Table 2 for enzyme substrate information. Data analysis credit: Andrew Steen.

Channel Position	Depth	Substrate	Avg. Activity	S.D. Activity
km	m		µmol/L/hr	µmol/L/hr
0.34	42.8	Leu	0.077	0.0091
0.34	42.8	Phe	0.049	0.0042
0.34	42.8	AAF	0.021	0.0033
0.34	42.8	NAG	0.008	0.0039
0.34	42.8	Cello	0.000	0.0002
0.34	42.8	b-glu	0.001	0.0005
0.34	42.8	xyl	0.000	0.0000
1.6	54	Leu	0.040	0.0065
1.6	54	Phe	0.031	0.0019
1.6	54	AAF	0.013	0.0006
1.6	54	NAG	0.003	0.0005
1.6	54	Cello	0.001	0.0000
1.6	54	b-glu	0.003	0.0001
1.6	54	xyl	0.001	0.0001
1.6	54	PO4	0.028	0.0008
1.6	39.83	Leu	0.056	0.0037
1.6	39.83	Phe	0.045	0.0020
1.6	39.83	AAF	0.020	0.0101
1.6	39.83	NAG	0.005	0.0006
1.6	39.83	Cello	-0.001	0.0003
1.6	39.83	b-glu	0.004	0.0002
1.6	39.83	xyl	0.000	0.0002
1.6	39.83	PO4	0.030	0.0035
1.6	30.4	Leu	0.053	0.0034
1.6	30.4	Phe	0.040	0.0045
1.6	30.4	AAF	0.012	0.0023
1.6	30.4	NAG	0.005	0.0002
1.6	30.4	Cello	0.001	0.0001
1.6	30.4	b-glu	0.003	0.0003
1.6	30.4	PO4	0.007	0.0032
1.6	14.95	Leu	0.040	0.0029
1.6	14.95	Phe	0.031	0.0014
1.6	14.95	Cello	0.006	0.0088
1.6	14.95	b-glu	0.002	0.0006
1.6	14.95	xyl	-0.001	0.0033
1.6	14.95	PO4	0.018	0.0021

1.6	0.15	Leu	0.090	0.0042
1.6	0.15	Phe	0.065	0.0028
1.6	0.15	AAF	0.014	0.0022
1.6	0.1,5	NAG	0.006	0.0002
1.6	0.15	Cello	0.001	0.0004
1.6	0.15	b-glu	0.005	0.0003
1.6	0.15	xyl	0.001	0.0002
1.6	0.15	PO4	0.036	0.0150
2.2	53.06	Leu	0.063	0.0047
2.2	53.06	Phe	0.028	0.0214
2.2	53.06	AAF	0.003	0.0025
2.2	53.06	NAG	0.005	0.0004
2.2	53.06	Cello	0.001	0.0001
2.2	53.06	b-glu	0.003	0.0001
2.2	53.06	xyl	0.001	0.0001
2.2	53.06	PO4	0.023	0.0019
2.2	29.83	Leu	0.064	0.0035
2.2	29.83	Phe	0.044	0.0115
2.2	29.83	AAF	0.010	0.0029
2.2	29.83	NAG	0.004	0.0009
2.2	29.83	Cello	0.002	0.0016
2.2	29.83	b-glu	0.002	0.0002
2.2	29.83	xyl	0.000	0.0001
2.2	29.83	PO4	0.015	0.0028
2.2	3.33	Leu	0.071	0.0288
2.2	3.33	Phe	0.040	0.0050
2.2	3.33	AAF	0.020	0.0186
2.2	3.33	NAG	0.003	0.0001
2.2	3.33	Cello	0.001	0.0004
2.2	3.33	b-glu	0.003	0.0012
2.2	3.33	xyl	0.001	0.0003
2.2	3.33	PO4	0.018	0.0010



Figure S1. Profiles of weight % particulate organic carbon (POC), bulk $\delta^{13}C_{POC}$ (‰) and C/N of soils sampled at various depths from the surface to 8 m in a soil pit from the Tanguro ranch in the Southern Amazon. Refer to Fig. 1 in chapter 5 for approximate ranch location.



Figure S2. (a) Total suspended sediment concentrations in three streams draining through three land cover types across the Tanguro Ranch: pristine, soy fields, and rotating soy and corn fields. (b) % organic carbon, (c) organic C:N and (d) $\delta^{13}C_{POC}$ (‰) measured in total suspended sediments, bedload and leaf litter collected from these three streams. Refer to Fig. 1 in chapter 5 for ranch location.

Supplementary Methods for Figs. S1-S2

During a field sampling trip to the Tanguro Ranch from November 12-20 2013, suspended sediment, bedload and suspended leaf litter samples were collected in three streams running through different catchments. Each catchment flushed through a different land cover type: pristine rain forest, a soy field, and an alternating soy/crop field. All samples were gathered, prepared and analyzed according to methods described in chapter 5. Leaf litter samples was separated from suspended sediments using a 2 mm diameter sieve after the suspended material was freeze-dried.

In addition, a soil pit on the ranch was sampled on November 19 and 20, 2013 at five depths from surface to 8 m. The soils were oven-dried overnight prior to organic composition analysis using methods described in chapter 5.

Supplementary acknowledgements for Figs. S1-S2

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