

Global Biogeochemical Cycles

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Key Points:

- One fifth of organic matter on Arabian Sea seafloor could stem from bacterial carbon fixation in the oxygen minimum zone
- Evaluation of past anoxic events needs to take chemoautotrophic contribution into account in isotope balances
- Biogeochemical models ignoring dark carbon fixation could highly underestimate oxygen demand and thus expansion of oxygen minimum

Supporting Information:

· Supporting Information S1

Correspondence to:

S. K. Lengger, sabine.lengger@plymouth.ac.uk

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Dark Carbon Fixation in the Arabian Sea Oxygen Minimum Zone Contributes to Sedimentary Organic Carbon (SOM)

Sabine K. Lengger^{1,2,3}, Darci Rush³, Jan Peter Mayser², Jerome Blewett², Rachel Schwartz-Narbonne⁴, Helen M. Talbot^{4,5}, Jack J. Middelburg⁶, Mike S.M. Jetten⁷, Stefan Schouten^{3,6}, Jaap S. Sinninghe Damsté^{3,6}, and Richard D. Pancost²

¹Biogeochemistry Research Centre, School of Geography, Earth and Environmental Science, University of Plymouth, Plymouth, United Kingdom, ²Organic Geochemistry Unit, School of Chemistry, University of Bristol, Bristol, United Kingdom, ³NIOZ Royal Netherlands Institute for Sea Research, Dept. of Marine Microbiology and Biogeochemistry, and Utrecht University, Texel, The Netherlands, ⁴School of Natural and Environmental Sciences, Newcastle University, Newcastle-upon-Tyne, United Kingdom, ⁵Now at BioArCh, Environment Building, University of York, Heslington, United Kingdom, ⁶Department of Earth Sciences, Faculty of Geosciences, Utrecht University, Utrecht, The Netherlands, ⁷Department of Microbiology, IWWR, Radboud University Nijmegen, Nijmegen, The Netherlands

Abstract In response to rising CO₂ concentrations and increasing global sea surface temperatures, oxygen minimum zones (OMZ), or "dead zones", are expected to expand. OMZs are fueled by high primary productivity, resulting in enhanced biological oxygen demand at depth, subsequent oxygen depletion, and attenuation of remineralization. This results in the deposition of organic carbon-rich sediments. Carbon drawdown is estimated by biogeochemical models; however, a major process is ignored: carbon fixation in the mid- and lower water column. Here, we show that chemoautotrophic carbon fixation is important in the Arabian Sea OMZ; and manifests in a ¹³C-depleted signature of sedimentary organic carbon. We determined the δ^{13} C values of C_{org} deposited in close spatial proximity but over a steep bottom-water oxygen gradient, and the δ^{13} C composition of biomarkers of chemoautotrophic bacteria capable of anaerobic ammonia oxidation (anammox). Isotope mixing models show that detritus from anammox bacteria or other chemoautotrophs likely forms a substantial part of the organic matter deposited within the Arabian Sea OMZ (~17%), implying that the contribution of chemoautotrophs to settling organic matter is exported to the sediment. This has implications for the evaluation of past, and future, OMZs: biogeochemical models that operate on the assumption that all sinking organic matter is photosynthetically derived, without new addition of carbon, could significantly underestimate the extent of remineralization. Oxygen demand in oxygen minimum zones could thus be higher than projections suggest, leading to a more intense expansion of OMZs than expected.

Plain Language Summary Oxygen minimum zones are areas in the ocean in which algae produce large amounts of organic material. When this sinks towards the seafloor, all oxygen at depth is used up. This results in vast "dead zones" where almost no oxygen is available to sustain life. With global warming, and increased nutrients from rivers, dead zones are forecast to expand. Computer models can calculate this, by considering algal production, and the amount of material delivered to the seafloor. However, these models often ignore a major process: anaerobic bacteria in the deeper water column, that can live at the edge or in the middle of these dead zones, which can also produce organic material from the dissolved CO₂. In this study, we used the fact that these bacteria add a distinct signature to the organic material, to show that one fifth of the organic matter on the seafloor could stem from bacteria living in these dead zones. Thus, models that have missed out on considering this contribution could have underestimated the extent of oxygen depletion we are to expect in a future, warming world. A more intense expansion of dead zones than expected could have severe ecological, economical (fisheries), and climatic consequences.

1. Introduction

Marine primary production fixes 50 Pg carbon per year, of which only about 1% is buried in sediments (Dunne et al., 2007; Middelburg, 2011). The majority of organic carbon derived from the photic zone is remineralised during sedimentation, fueling heterotrophic bacterial activity in the water column (Keil et al., 2016). In marginal settings and OMZs, marine primary production in the photic zone can be significantly higher than in other settings. Organic carbon (OC) sedimentary accumulation rates within an OMZ can

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be in the range of tens to hundreds of mg C cm⁻² y⁻¹ (Hartnett et al., 1998; Hedges & Keil, 1995) higher than observed in other parts of the ocean. These high accumulation rates are most commonly attributed to attenuation in remineralization rates within the OMZ, and low bottom-water oxygenation, which results in decreased biodegradability of polymeric and matrix-protected substances (Burdige, 2007).

As a consequence of increasing atmospheric CO_2 concentrations and, consequently, temperature, oceanic OMZs are forecast to expand in a fashion similar to the past (Breitburg et al., 2018; Queste et al., 2018; Schmidtko et al., 2017; Shaffer et al., 2009; Stramma et al., 2010). Whilst the expansion of OMZs will result in widespread habitat loss of marine life and could cause an increase in emissions of greenhouse gases such as N_2O and CH_4 , it could also act as a long-term negative feedback on global warming via the enhanced drawdown and storage of organic carbon in sediments.

The biogeochemical system in subsurface waters, where light does not penetrate, has recently emerged to be substantially more complex – and possibly more important for the global carbon cycle – than previously assumed. In particular, dark water-column microbial activity is higher than what can be accounted for by heterotrophs (Herndl & Reinthaler, 2013), suggesting an important role for chemoautotrophy, i.e. fixation of dissolved inorganic carbon (DIC). It has been suggested to contribute substantially to the global carbon budget, with estimates ranging from 0.11 to 1.1 Pg C y⁻¹, equating to ca. 2% of total estimated yearly marine primary production (Middelburg, 2011; Reinthaler et al., 2010). The predominant chemoautotrophic process in the oxic, dark, pelagic ocean is thought to be nitrification (Middelburg, 2011; Pachiadaki et al., 2017). When oxygen is limited, nitrification still occurs, but other chemoautotrophic processes dominate, such as anaerobic oxidation of ammonia and sulfide oxidation (Ulloa et al., 2012; Wright et al., 2012).

Under hypoxic conditions, such as in the water column of OMZs, both archaeal (aerobic) and anaerobic oxidation of ammonia are thought to dominate dark inorganic carbon fixation processes (Lam & Kuypers, 2010; Pitcher et al., 2011). Here, nitrite accumulates, and other anaerobic autotrophic processes such as sulfide oxidation and methanogenesis are suppressed, most likely due to the abundance of nitrate and ammonia (Canfield, 2006; Ulloa et al., 2012).

Of the inorganic carbon converted to organic matter within the OMZ, only a negligible fraction is presumably transported to the sediments and preserved, as this newly produced material is regarded as more labile than the sinking OC derived from the photic zone (Cowie & Hedges, 1992; Keil et al., 1994; Middelburg, 1989). Dark carbon fixation rates are challenging to quantify: they have been determined experimentally (Reinthaler et al., 2010; Taylor et al., 2001), or have been estimated from the reaction stoichiometry of respiration based on Redfield organic matter and growth yields of nitrifiers (Middelburg, 2011; Wuchter et al., 2006). In OMZs, such as the Peruvian margin (Lam et al., 2009), the Arabian Sea (Jensen et al., 2011), or the sulfidic Black Sea (Lam et al., 2007), the activity of some chemoautotrophs was determined via ¹⁵N- labelling, and formation of the products of their biogeochemical reactions. However, incubation methods may suffer from bias, because *in situ* conditions such as pressure are difficult to maintain.

However, carbon from within anoxic waters has been observed in some settings to contribute to the particulate OC flux: for example, in eutrophic lakes (Hollander & Smith, 2001) and anoxic fjords (van Breugel et al., 2005). Furthermore, discrepancies between modelled and observed organic carbon fluxes suggest that dark carbon fixation in anoxic marine settings significantly contributes to sinking material (Keil et al., 2016; Taylor et al., 2001).

One way to constrain this input into sedimentary organic matter is to use isotope mixing models. Photosynthetically fixed carbon generally has stable carbon isotopic compositions of ca -19 to -21 % due to Rubisco fixation. However, chemoautotrophs dwelling in OMZs typically have a lower $\delta^{13}C$ values. This is the result of multiple factors: they either use ^{13}C -depleted CO_2 generated by remineralization, have larger fractionation factors due to the higher abundance of CO_2 at depth (Freeman et al., 1994), or use carbon fixation pathways such as the acetyl coenzyme A pathway, which results in ^{13}C -depleted biomass (Hayes, 2001). This characteristic chemoautotrophic isotopic signature in the organic carbon could allow us to quantify the contribution of dark carbon fixation to sedimentary organic matter.

Here, we investigated the δ^{13} C value of sedimentary organic matter of surface sediments deposited within the OMZ of the Arabian Sea and employed a simple isotope mixing model to investigate the extent of input from OMZ carbon fixation into sedimentary organic matter. As the major process in the Arabian Sea OMZ



known to produce isotopically light biomass is anaerobic oxidation of ammonia (anammox; Ulloa et al., 2012; Villanueva et al., 2014), in order to determine the isotopic signature of this pathway, we developed and applied a method to determine the δ^{13} C values of a novel biomarker, bacteriohopanetetrol stereoisomer (BHT'), which has been found to be unique to anammox bacteria in the marine environment in culture and environmental studies (Rush et al., 2014, 2019; Rush & Sinninghe Damsté, 2017). It has been found in the Arabian Sea and other marine anoxic settings (Matys et al., 2017; Sáenz et al., 2011). We also used stable isotope probing experiments to exclude sedimentary anammox as an important contributor to this process. This allowed us to investigate the contribution of these dark carbon fixers to sedimentary organic carbon.

2. Materials and Methods

2.1. Sediment sampling and stable isotope probing incubations

Sediments were collected with multicore devices on the R/V Pelagia in the Northern Arabian Sea in January 2009 during the PASOM cruise 64PE301 along the Murray Ridge (Figure 1f), which protrudes into the core of the OMZ. Two cores, one each from P900 (885 m water depth) and P1800 (1786 m water depth), hereinafter referred to as anoxic and oxic, respectively, were incubated on board as described by Pozzato et al. (2013a, 2013b). In brief, particulate or dissolved organic matter from the diatom *Thalassiosira pseudonana* containing 20 and 18 % ¹³C, respectively, were added to the tops of core tubes of 10 cm internal diameter. Between 2 and 6 % of the added carbon was respired, resulting in a highly enriched ¹³C_{DIC} pool, enabling the tracing of autotrophic processes in addition to heterotrophic processes.

Eight cores are discussed here; these were incubated under oxic or suboxic conditions for 7 days (125 μ M, 6 μ M O₂, respectively; Table 1). At the end of incubation, cores were sliced in the intervals 0 – 2, 2 – 4, and 4 – 10 cm depth. They were then frozen and freeze dried for the isotopic analysis of the bacteriohopanepolyol lipids (BHPs), including bacteriohopanetetrol (BHT) and its stereoisomer and biomarker for anammox bacteria, BHT'. Both biomarkers have been studied previously in the Arabian Sea water column and sediments (Jaeschke et al., 2009; Sáenz et al., 2011) Furthermore, cores from 8 stations between 900 and 3000 m water depth were also collected, as described by Lengger et al. (2014, 2012). The top 0 – 0.5 cm were used for total organic carbon (TOC) and 13 C of organic carbon analysis. For the core from P900 (32 cm length), all depths were analysed in 0.5 cm – 4 cm resolution (Lengger, Hopmans, Reichart, et al., 2012).

2.2. Anammox enrichment cultures

To determine the δ^{13} C values for the two bacteriohopanetetrols in anammox bacteria (BHT and BHT', the latter being unique to anammox in the marine environment), an enrichment culture of 'Ca. Scalindua profunda' was analysed. It was grown in a sequencing batch reactor as described by van de Vossenberg et al. (2008). Analysis of this enrichment culture that showed 'Ca. S. profunda' comprised about 80% of the cells, while other bacteria belonging to the phyla Bacteriodetes (including Flavobacteriaceae) and Proteobacteria (including Alphaproteobacteria) accounted for the majority of the remaining populations (van de Vossenberg et al., 2008, 2013).

2.3. Extraction and purification

The freeze-dried subsamples of the unamended and incubated cores were ground, and the homogenised sediments and the culture were extracted by a modified Bligh-Dyer extraction method (Lengger et al., 2012). Briefly, they were extracted ultrasonically three times in a mixture of methanol/dichloromethane (DCM)/phosphate buffer (2:1:0.8, v:v:v) and centrifuged, and the solvent phases were combined. The solvent ratio was then adjusted to 1:1:0.9, v:v:v to separate the DCM phase. Liquid extraction was repeated two more times, the DCM fractions were combined, the solvent was evaporated and the larger particles were filtered out over glass wool. The extraction procedure was performed on the enrichment culture material and repeated on the sediment for analysis of BHPs. An aliquot of the extract was subjected to column chromatography using 5% aminopropyl solid phase extraction (SPE), eluting with hexane, DCM, and MeOH, which contained BHPs. For analysis by chromatographic techniques, the extract was derivatised in 0.5 mL of a 1:1 (v:v) mixture of acetic anhydride and pyridine at 50 °C for 1 h, then at room temperature overnight in the case of HPLC-MS analysis. Solvent was dried under a stream of N_2 on a 50°C heating block.



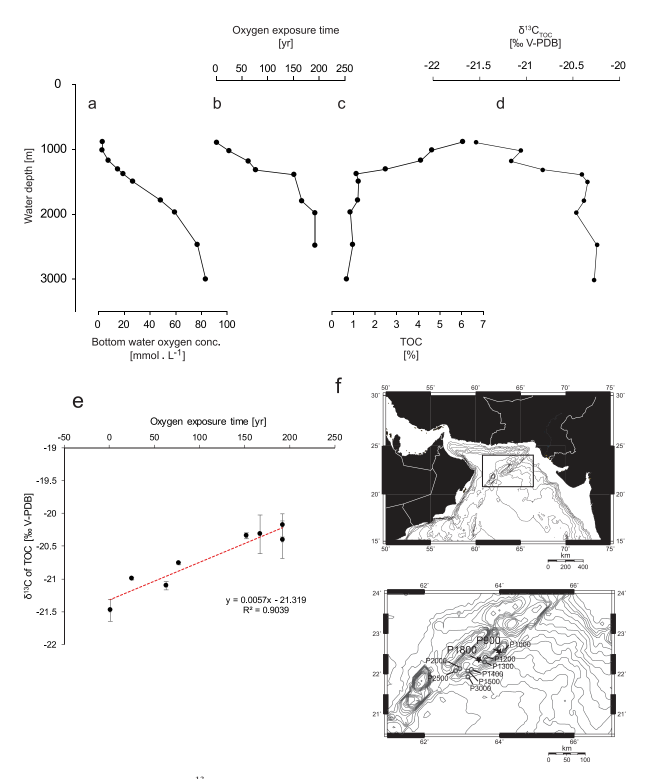


Figure 1. Arabian Sea depth gradients. Shown are $\delta^{13}C_{org}$ and % TOC values of core top sediments, and bottom water oxygenation plotted with depth (a-d), all but $\delta^{13}C_{org}$ replotted from Lengger et al. (2014), a scatter plot of oxygen exposure time versus $\delta^{13}C_{org}$ in Arabian Sea core tops along Murray Ridge (e), and a map of the sampling stations with the two main stations used for BHT analysis used here indicated with a star (f).

2.4. Instrumental techniques

2.4.1. High temperature gas chromatography coupled to flame ionization detection (HTGC-FID)

GC analysis of acetylated BHPs was done using a HP-5890 Series II GC equipped with a flame ionization detector was fitted with a 0.25 mm \times 0.1 μ m VF5-ht capillary column (CP9045, CP9046, Agilent



Table 1 $\delta^{13}C$ values of analysed samples. s.d. = standard deviation of three repeat analyses, n.d. = not determined (single analysis)

Туре	Sample			δ ¹³ C value [‰ V-PDB]			
		внт	s.d.	BHT'	s.d.	C _{org}	
Biomass	Scalindua sp.	-47.8	1	-47.6	1		
Arabian Sea Natural abundance	P900-1 cm	-24.7	1.3	-39.1	1.3	-21.5	
	P900-3 cm	-27.0	1.4	-48.1	1.1		
	P900-7 cm	-26.1	0.4	-41.4	0.5		
	P1800-1 cm	-25.1	1.7	-48.1	1.6	-20.3	
	P1800-3 cm	-28.8	2.1	-46.6	4.1		
	P1800-7 cm	-27.1	0.6	-45.5	1.0		
Arabian Sea Incubations	P900, suboxic, DOM - 1 cm	-27.9	n.d.	-45.9	n.d.		
	P900, suboxic, DOM - 3 cm	-26.6	n.d.	-42.0	n.d.		
	P900, suboxic, POM - 1 cm	-41.6 ^a	n.d.	-75.2 ^a	n.d.		
	P900, suboxic, POM - 3 cm	-26.4	n.d.	-49.8	n.d.		
	P900, oxic, POM - 1 cm	-27.1	n.d.	-51.5	n.d.		
	P900, oxic, POM - 3 cm	-25.2	n.d.	-47.8	n.d.		
	P1800, oxic, DOM - 1 cm	-21.3	n.d.	-42.0	n.d.		
	P1800, oxic, DOM - 3 cm	-28.3	n.d.	-45.6	n.d.		
	P1800, suboxic, POM - 1 cm	-52.2 ^a	n.d.	-101.9 ^a	n.d.		
	P1800, suboxic, POM - 3 cm	-28.8	n.d.	-47.1	n.d.		
	P1800, oxic, POM - 1 cm	-30.2	n.d.	-51.3	n.d.		
	P1800, oxic, POM - 3 cm	-28.0	n.d.	-46.1	n.d.		

The values shown are natural abundance from unamended cores, and from incubations with ¹³C labelled DOM and POM, under oxic and suboxic conditions. ^aindicates outliers (Grubbs).

Technologies UK Ltd., Stockport, UK) of 30 m length (Lengger et al., 2018). An on-column injector was used. To the 30 m column, 1m of a 0.25 mm HT-deactivated silica tubing was attached as a guard column (Zebron Z-Guard, 7CG-G000-00GH0, Phenomenex, Macclesfield, UK). Analysis of bacteriohopanepolyols employed a constant flow of 2 ml/min He and a temperature ramp from 70°C (1 min hold) to 400°C at 7°C min⁻¹ (1 min hold).

2.4.2. High temperature gas chromatography coupled to mass spectrometric detection (HTGC-MS)

Analysis of acetylated BHPs, using HTGC-MS was performed using a Thermo Scientific Trace 1300 gas chromatograph coupled with an ISQ single quadrupole mass spectrometer. Diluted samples were introduced using a PTV injector in splitless mode onto a 0.53 mm fused silica pre-column connected to a 30 m \times 0.25 mm i.d. fused-silica capillary column coated with dimethyl polysiloxane stationary phase (ZB-5HT; film thickness, 0.1 μ m; 7HG-G015-02, Phenomenex, Macclesfield, UK). The initial injection port temperature was 70 °C with an evaporation phase of 0.05 min, followed by a transfer phase from 70 °C to 400 °C at 0.2 °C s $^{-1}$. The oven temperature was held isothermally for 1 min at 70 °C, increased at a rate of 7 °C min $^{-1}$ to 400 °C and held at 400 °C for 10 min. Helium was used as a carrier gas and maintained at a constant flow of 2.5 ml min $^{-1}$. The mass spectrometer was operated in the electron ionization (EI) mode (70 eV) with a GC interface temperature of 400 °C and a source temperature of 340 °C. The emission current was 50 μ A and the mass spectrometry set to acquire in the range of m/z 50–950 Daltons with 0.5 s dwell time. Data acquisition and processing were carried out using the Thermo XCalibur software (version 3.0.63). Due to the lack of authentic standards for BHT and BHT', only relative and not absolute values are reported, assuming similar ionization energies.

2.4.3. High temperature gas chromatography coupled to isotope ratio mass spectrometry (HTGC-IRMS)

The stable carbon isotopic composition (δ^{13} C) of BHPs were determined using HTGC-isotope ratio mass spectrometry. To this end, an Elementar visION IRMS with GC5 interface (Elementar UK Ltd., Cheadle, UK), and an Agilent 7890B GC were modified in-house and allowed us to achieve column temperatures of up to 400 °C, which resulted in baseline resolution of BHT and BHT' (Figure 2). 1 μ l of the derivatized samples dissolved in ethyl acetate were injected on a cool-on-column injector, into a Zebron Z-Guard Hi-Temp Guard Column (1 m × 0.25 mm, Zebron Z-Guard, 7CG-G000-00GH0, Phenomenex, Macclesfield, UK) and

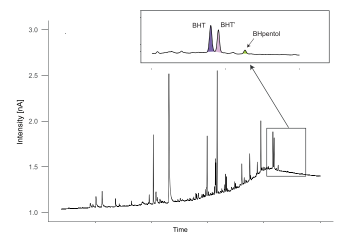


Figure 2. HTGC-IRMS chromatogram showing baseline separation between BHT, BHT', and BHpentol.

separated on a Zebron ZB-5HT analytical column (30 m \times 0.25 mm \times 0.1 μm, Phenomenex Ltd., Macclesfield, UK). He was used as a carrier gas at a flow rate of 1.5 ml min⁻¹ and the oven was programmed as follows: 1 min hold at 70 °C, increase by 7 °C min⁻¹ to 350 °C (10 min hold). Organic compounds were combusted to CO2 in a 0.7 mm ID quartz tube with CuO pellets at 850°C. Instrumentation performance was monitored using an n-alkane standard (B3, A. Schimmelmann, Indiana University, Bloomington, IN, USA; RMS 0.4 %), and results were calibrated using an in-house mixture of five fatty acid methyl esters, which was injected between every six sample analyses and analyzed using a He flow of 1 ml min⁻¹, with a slightly different temperature program (injection at 50 °C held for 1 min followed by an increase of 10°C min⁻¹ to 300 °C and a 10 min hold). This is the first time baseline resolution between BHT and BHT' has been achieved on a GC-IRMS, which allows the direct determination of the isotopic composition of both BHT and BHT' in sediment samples (Figure 2). The isotopic composition of the acetyl group used to derivatise the BHT and BHT' was determined by acetylation of myo-

inositol, and then subtracted from the values of BHT and BHT' in a mass balance correction (Angelis et al., 2012), as authentic standards for BHT or BHT' were not available.

2.4.4. High performance liquid chromatography coupled to positive ion atmospheric pressure chemical ionization mass spectrometry (HPLC/APCI-MS)

To verify the GC-derived assignments, an aliquot of the acetylated BHP samples was dissolved in MeOH:propan-2-ol (3:2; v:v) and filtered on 0.2 μ m PTFE filters. BHPs were analysed by HPLC/APCI-MS, using a data-dependent scan mode (3 events) on an HPLC system equipped with an ion trap MS, as described in Talbot et al. (2007) and van Winden et al. (2012). Relative BHP concentrations were semi-quantitatively estimated based on the response factor of authentic standards (M. Rohmer; Strasbourg, France; Cooke et al., 2008), with a typical reproducibility of \pm 20%, according to Cooke et al. (2009).

2.4.5. Bulk sedimentary organic matter and suspended particulate matter

Freeze-dried core tops from 8 stations between 900 and 3000 m for sedimentary organic matter, and punches from 0.7 μ m GFF filters for suspended particulate organic matter, were decalcified with 2N HCl, washed, freeze-dried, and subjected to analysis via a Flash EA 1112 Series (Thermo Scientific) analyser, coupled via a Conflo II interface to a Finnigan Delta^{plus} mass spectrometer as described by Lengger et al. (2014; sediment) and Pitcher et al. (2011; filters). Standards for δ^{13} C analysis were acetanilide and benzoic acid and samples were analysed in duplicate.

3. Results

To quantify the provenance of sedimentary organic matter and the contribution of anammox from the OMZ, we analysed the isotopic composition of sedimentary organic matter deposited in close spatial proximity over a large depth gradient in the Arabian Sea, as well as the isotopic composition of biomarker lipids derived from anammox bacteria, chemoautotrophic microbes living in the OMZ. In order to determine whether these were water-column derived or sedimentary, we used stable isotope probing experiments on sediments retrieved from within and below the OMZ.

3.1. δ^{13} C values of C_{org}

 $\delta^{13}C$ values of C_{org} in surface sediments were low (-21.5 %) at P900 and increased with water depth to -20.2 % at P2500 (Figure 1d). $\delta^{13}C_{org}$ values correlated positively and linearly (Slope 0.0057, R^2 = 0.90, Figure 1e) with oxygen exposure times as calculated by Koho et al. (2013) and Lengger et al. (2014). Similarly, organic carbon content in the core tops was negatively correlated with oxygen exposure times (R^2 =0.93, from Lengger et al., 2014). The increase mirrors the decrease in % TOC – and thus progressing degradation – with increasing oxygen exposure time (Figure 1b,c, Lengger et al., 2014). At P900, where the whole depth of the core was analysed, values increased slightly with depth, from -21.5 % at the surface, to -20.9 % (Figure 3). $\delta^{13}C$ values of particulate organic carbon (suspended particulate organic matter) decreased throughout the

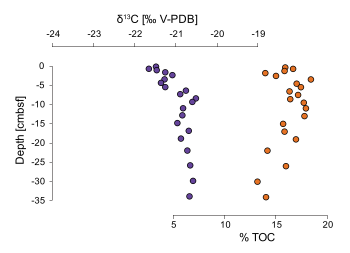


Figure 3. Depth profile of $\delta^{13}C_{\rm org}$ and % TOC in an unamended core at P900

water column from -19 to -21.8 ‰, though with a substantially ¹³C-depleted, yet unexplained, value at the very surface (20 m depth) of -22.9 ‰ (Figure S1).

3.2. Biomarkers

3.2.1. Bacteriohopanepolyols (BHPs)

Analysis of BHPs in the Arabian Sea cores using both HPLC-APCI-MS and HTGC-MS, all showed that BHT', specific for *Scalindua*, was abundant, and the fractional abundance of BHT's when compared to the sum of BHT and BHT' ranged from 0.4 to 0.6. Other, relatively non-source-specific, BHPs were also present: BHT, 35-aminobacteriohopane-32,33,34-triol (aminotriol), bacteriohopane-31,32,33,34,35-pentol (BHpentol), bacteriohopanetetrol cyclitol ether (BHT-CE) and anhydro-BHT (Figure S3). The relative concentrations of bacteriohopanetetrols were 75 to 96 % of total bacteriohopanepolyols, an order of magnitude higher than other BHPs (Figure S3). HTGC-MS was able to detect BHT, BHT' and BHpentol (Figure S2), as well as small amounts of anhydroBHT, a BHT degradation product, and BHP-570, which has been

tentatively identified by Sessions et al. (2013) as acetylated bacteriohopanediol, possibly also a degradation product of bacteriohopanetetrol. In the core from P900, the ratio of BHT over BHT increased with depth (Figure S4).

We also analysed the BHP content of sediment cores incubated with ¹³C-labelled organic matter at both sites under the different conditions detailed by Pozzato et al. (2013a, 2013b), which was respired and generated ¹³C-labelled DIC, allowing to trace autotrophic processes such as anammox. No changes were noted, except for in P900; in those, BHT abundance increased (i.e. BHT'/BHT decreased), indicating that some of the sedimentary BHT could have been produced in situ (Figure S4).

3.2.2. BHT and BHT' δ^{13} C values

To establish the isotopic difference of BHT and BHT' derived from anammox we analysed biomass obtained from a batch reactor. BHT and BHT' were the main biohopanoids in the biomass from 'Ca. S. profunda' detected by HTGC-MS (Figure S2b). In addition to being present in 'Ca Scalindua sp.' anammox bacteria, BHT is a ubiquitous lipid common to many bacteria, BHT', however, is specific to Ca Scalindua sp.' in marine environments (Rush et al., 2014). The δ^{13} C values of BHT and BHT' were identical within the error of analysis (-49 and -48 ‰, respectively; Table 1), indicating identical fractionation and thus biosynthetic pathways for both lipids.

In the Arabian Sea sediments (all unamended cores), BHT was markedly enriched in 13 C relative to BHT', with values ranging from -24.7 to -28.8 % and -39.1 to -48.1 %, respectively (Figure 4b-4c). At P1800 (below the OMZ), BHT and BHT' were slightly more depleted in 13 C, with BHT at -27 \pm 3 % and BHT' at -47 \pm 4 %, as compared to -26 \pm 1 % and -43 \pm 5 % for BHT and BHT' at P900 (in the OMZ). However, the difference between P900 and P1800 was not statistically significant for either BHT or BHT'. Moreover, even though the proportion of BHT' increased with depth in the anoxic core, the δ^{13} C values did not change. We also analysed BHT and BHT' in the cores that had been incubated with 13 C-labeled POM and DOM, and these showed no indication of 13 C-enrichment in BHT'. Excluding outliers (defined by a Grubbs test at 99% confidence level and indicated in Table 1), BHT' values were on average -48 \pm 4 % and -46 \pm 2 % at P1800, and P900, respectively. BHT was slightly enriched compared to the unamended incubations at 3 cm depth ($\Delta\delta^{13}$ C = 4.6 \pm 0.7 %), but not at 1 cm depth. BHpentol concentrations were too low to allow reliable isotopic determination, and anhydroBHT and BHP-570 co-eluted with other compounds, also precluding their isotopic characterization.

4. Discussion

4.1. Origins of BHT and BHT' - a biomarker for anammox

BHT', a biomarker likely unique for annamox in marine environments (Rush et al., 2014,), is highly abundant in the sediment and occurred throughout both sediment cores (Figure 4a), suggesting a significant contribution of anammox biomass to sedimentary organic matter. In the anoxic core (P900), BHT and BHT'

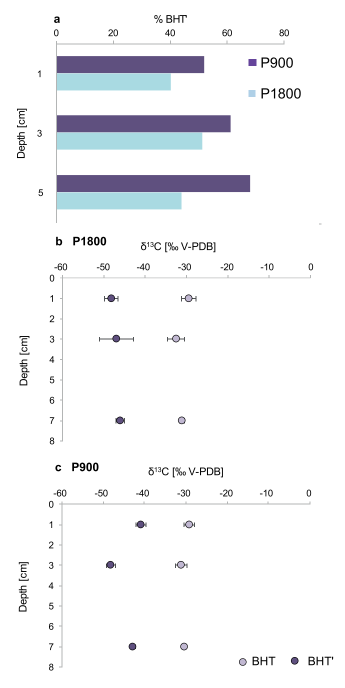


Figure 4. Anammox biomarkers in unamended Arabian Sea sediment. Panel (a) shows the proportion of BHT' relative to BHT+BHT', panels (b) and (c) show the δ^{13} C values of BHT and BHT' in the unamended oxic (P1800) and anoxic core (P900), respectively.

were present in concentrations at least an order of magnitude higher than other BHPs (Figure S2), and BHT' was the most abundant of the two stereoisomers (Figure 4a). The high fractional abundance of BHT' over the sum of BHT and BHT' (0.4 – 0.6) is contrary to our expectations, as BHT is a ubiquitous lipid and presumed to derive from both anammox and non-anammox sources such as cyano- and many other bacteria (Pearson & Rusch, 2009); it is, therefore, expected to be abundant in most depositional contexts. Lower proportions of BHT were reported previously in nearby core tops (0.22-0.30) by Sáenz et al. (2011); this could be caused by a difference in settings, or in BHP extraction protocol. However, the high proportions of BHT' observed in the Arabian Sea are not unprecedented: they are slightly lower than BHT' proportions in sediments underlying the Humboldt Current System OMZ offshore Peru (0.45 – 0.69 in surface sediments, Matys et al., 2017).

Supporting evidence for the unique source of BHT' comes from its 13 C values of -40 to -50‰. Anammox bacteria are known to fractionate strongly against δ^{13} C, with up to 26 ‰ fractionation observed for biomass in cultures and sediment, with even more strongly depleted BHT and BHT' (Brocadia sp. by 47 ‰ and Scalindua sp. by 49 ‰ against DIC; Schouten et al., 2004). Anammox bacteria within the oxygen minimum zone and using dissolved inorganic carbon, which is up to 2 ‰ lighter within the OMZ than at the surface (Kroopnick, 1985), are likely to produce such 13 C-depleted lipids. In line with an anammox origin, BHT' here is decidedly more depleted than other biomarkers in the Arabian Sea (cf. Wakeham & McNichol, 2014), though distinctly depleted highly branched isoprenoids (HBIs, -37 ‰) have been found in Arabian Sea cores from the Holocene (Schouten et al., 2000).

BHT, however, is much more enriched in ¹³C in the Arabian Sea sediments, with values of -29 to -30 % (Figure 4b,c). This is in contrast to anammox cultures, where BHT and BHT' are produced with identical isotope values (Figure S4, Table 1). This strongly supports the idea that BHT has a mixed origin. The sources for BHT could be varied: cyanobacteria or heterotrophic bacteria thriving in the OMZ or the sediment (Pearson & Rusch, 2009), and possibly including nitrite-oxidising bacteria (Kharbush et al., 2018). The δ^{13} C values of BHT derived from these alternative sources are poorly constrained (Hayes, 2001; Kharbush et al., 2018; Pearson, 2010; Sakata et al., 1997; Schouten et al., 1998). However, for heterotrophic bacteria, values similar to the consumed OM minus the depletion associated with polyisoprenoids of 6-8 % are expected (Pancost & Sinninghe Damsté, 2003). Similar values would be expected for cyanobacterial lipids, too, as those are depleted by 22 - 30 % compared to dissolved CO2, which varies from 0 to 2 ‰ in the photic zone. Methane cycling could also lead to 13C depleted OM, but is not important in the Arabian Sea (Lüke et al., 2016), and there is no biomarker evidence for

it, such as 13 C depleted archaeal lipids, or methylated BHPs. Thus, a BHT δ^{13} C value of -30 ‰ suggests heterotrophic and other bacterial sources, possibly mixed with an anammox source.

Anammox activity has been shown to occur both within the OMZ water column (Jensen et al., 2011; Lüke et al., 2016; Pitcher et al., 2011; Villanueva et al., 2014) as well as in sediments (Devol, 2015; Sokoll et al., 2012). To test whether the anammox-derived biomarkers are formed in the sediment, the labelled cores were incubated for 7 days with particulate and dissolved organic matter (Table 1), which resulted in the substantial incorporation of 13 C into bacterial fatty acids, as well as generating 13 C-enriched CO₂ due to heterotrophic respiration (up to 14 % of the added C was respired, Pozzato, et al., 2013a, 2013b). Anammox



bacteria are autotrophic, and it is therefore likely that an active sedimentary community would result in some uptake of this 13 C labelled CO $_2$ formed by respiration. However, no significant labelling was observed in BHT' or BHT (Table 1, Table S1), suggesting that most of this pool is water-column derived. Further, BHT and BHT' are also present in the surface of the oxic sediments (P1800), in proportions similar to the anoxic sediments at P900. Even if anammox growth was too slow for labelling to take effect, substantial sedimentary production would have resulted in a decreasing δ^{13} C value of BHT' in the unamended cores, as DIC gradually becomes more 13 C depleted with sediment depth (Fernandes et al., 2018), but this is not observed (Figure 4b and 4c). Collectively, these data support the idea that the vast majority of BHT' is derived from anammox bacteria living in the OMZ of the water column, and that BHT also has a predominant pelagic origin with limited sedimentary production.

The δ^{13} C values of geohopanoids (i.e. the geological degradation product of biohopanoids such as BHT and BHT') can exhibit dramatic variability, and often pronounced depletion in terrestrial (Pancost et al., 2007) and marine settings (Köster et al., 1998). These are commonly attributed to aerobic methane oxidising bacteria, and thus regarded as evidence for a significant contribution of methane oxidisers to sedimentary organic matter. However, our data show that hopanes of -35 to -50 % could also indicate the presence of a significant amount of anammox bacteria in an anoxic water column. Several factors could attenuate the anammox signal. The decrease of biohopanoids in structural complexity upon degradation means that the anammox signal would be diluted by mixing with aerobically and anaerobically produced hopanoids such as those of Geobacter (Fischer et al., 2005; Härtner et al., 2005). Chemoautotrophs operating in euxinic settings employ biochemical pathways resulting in ¹³C-enriched biomass and lipids compared to DIC (van Breugel, Schouten, Paetzel, Ossebaar, & Sinninghe Damsté, 2005). However, in anoxic basins such as the Black Sea or anoxic fjords, remineralization of organic matter also results in distinctly depleted $\delta^{13}C_{\rm DIC}$ below the chemocline (-12 %; Fry et al., 1991; Volkov, 2000). This may explain why, in some marine anoxic settings where we might expect to see an anammox signature, the ¹³C depletion of hopanes parallels that of algal biomarkers (Sinninghe Damsté et al., 2008; van Breugel et al., 2005). Nonetheless, we suggest that potential anammox contributions to the sedimentary hopanoid pool should be considered when interpreting their abundances, distributions and isotopic compositions.

4.2. Origin of sedimentary organic matter (SOM)

The Murray Ridge represents an open ocean setting, and the selected coring sites were in close proximity to each other, with no substantial terrigenous contribution (Koho et al., 2013; Lengger et al., 2014; Lengger, Hopmans, Reichart, et al., 2012; Nierop et al., 2017; Figure 1d). Despite this, δ^{13} C values of sedimentary organic matter of a purely marine origin varied: The value at the shallowest location (-21.5 %; P900), within the OMZ, was 2.5 % more depleted than the estimated value for surface water-derived OM -19.8 %(Fontugne & Duplessy, 1986), and the δ^{13} C values of sedimentary C_{org} increased with increasing oxygen bottom water concentrations/oxygen exposure time (Figure 1e). These observations agree with earlier studies from this setting. Cowie et al. (2009, 1999) detected similar trends in surficial sediments across different settings in the Arabian Sea, with values of -21 % within and -19 % above and below the OMZ. Organic matter in sediment traps collected in the north western Arabian Sea (i.e. sinking POC) had a δ^{13} C value of -22.4 ‰ (composite of the OMZ between 500 and 900 m depth). The corresponding sedimentary $\delta^{13}C_{opp}$ value, from oxygenated bottom waters at 1445 m depth, was, however, more enriched (-20.8%; Wakeham & McNichol, 2014). Fernandes et al. (2018) detected similar, though less pronounced, trends in sediments collected from the Pakistan margin. An increase in $\delta^{13}C_{org}$ with enhanced degradation, i.e. ^{13}C -enriched sediments vs. depleted POM, was also observed in the South China Sea (Liu et al., 2007), and in the Eastern Tropical North Pacific (Jeffrey et al., 1983).

Despite its common occurrence in OMZ settings, this trend is unusual and, at present, not explained: degradation of organic carbon in marine environments usually preferentially removes isotopically heavy carbon (Hatch & Leventhal, 1997), causing a depletion in δ^{13} C with increased degradation of the sediment. This can be due to the removal of the more labile marine carbon, and subsequent relative enrichment of terrigenous organic material of a lower initial reactivity and lower δ^{13} C values (Huguet et al., 2008; Middelburg et al., 1993). However, progressive depletion also occurs in areas with purely marine input; this is due to preferential loss of 13 C-enriched carbohydrates over the more 13 C-depleted lipids, and preferential degradation of easily accessible material over biopolymers (Spiker & Hatcher, 1987), and polymerization and elimination



of functional groups (Balabane et al., 1987; Galimov, 1988). Conversely, sulfurization, a process observed in euxinic settings appears to preferentially preserve 13C-enriched material such as carbohydrates (Van Kaam-Peters et al., 1998); however, this process is not expected to occur here - the Arabian Sea is anoxic but not sulfidic (Ulloa et al., 2012) and has not experienced euxinia for the past 120 ka (Schenau et al., 2002). Nonetheless, in Arabian Sea sediment, $\delta^{13}C_{org}$ values increased by 1.8 % with increasing oxygen exposure time and thus increasing degradation; at the same time, organic carbon contents decreased from 60 to 10 mg g dw⁻¹, indicating progressing remineralization (Figure 1c, Lengger et al., 2014). Cowie (2005) attributed this to the contribution of – potentially – organic matter from the facultatively autotrophic, chemosynthetic sulfur-bacterium Thioploca sp., which has been observed in the Arabian Sea (Schmaljohann et al., 2001). However, Thioploca sp. has only been reported for shelf and upper slope sediments in the Arabian Sea (above and upper part of OMZ), and it is unlikely that the sulfur-dependent Thioploca sp. could have caused this significant depletion by chemoautotrophy, as sulfide concentrations are negligible within the OMZ (Kraal et al., 2012), and there is no evidence for the production of severely ¹³C-depleted biomass by filamentous sulfur bacteria (Zhang et al., 2005). Further, the depletion is not only observed in sediments, but also in particle fluxes (Wakeham & McNichol, 2014), which strongly suggests the exclusion of sedimentary sources for 13 C-depleted organic matter. This is also supported by the δ^{13} C_{org} values of the suspended particulate matter (Figure S1), which becomes gradually more depleted with depth.

An active nitrogen-cycle in the water column within the Arabian Sea OMZ is undisputedly present, with heterotrophic denitrifying bacteria, nitrifying archaea, nitrite-oxidising and anammox bacteria present in high abundances (Lüke et al., 2016; Villanueva et al., 2014). The ¹³C fractionation for the carbon fixation pathways employed by nitrifying archaea (3-Hydroxypropionate/4-Hydroxybutyrate, 3-HP/4-HB) is similar to that of phytoplankton using Rubisco (Könneke et al., 2012). The biomass of heterotrophic denitrifiers is close to the value of the source organic matter (1 ‰ more enriched; Hayes, 2001), and the biochemical pathway employed by nitrite oxidisers (reverse TCA cycle) produces ¹³C-enriched biomass (Pearson, 2010). Anammox bacteria, however, are abundant and active in the Arabian Sea (Jensen et al., 2011) and known to produce highly ¹³C-depleted biomass by inorganic carbon fixation (Schouten et al., 2004). Anammox bacteria, and other, yet undescribed chemoautotrophs, could thus present a potential pathway for addition of ¹³C-depleted organic matter to sinking organic matter, a hypothesis we explore further below.

The water-column derived 13 C-depleted BHT', in combination with the unusual δ^{13} Corg trends, suggests that there may be a substantial contribution of 13 C depleted organic carbon, produced by anammox bacteria, to the sedimentary organic matter. Based on BHT' δ^{13} C values determined in this study, and the fractionation factor associated with anammox lipid biosynthesis of $16\,\%$ (lipid versus total biomass; Schouten et al., 2004), we can estimate a δ^{13} C value for anammox biomass: of ca.-28.6 \pm 6 %, which is similar to the expected value calculated from $\epsilon_{biomass-DIC}$ (22-26 %; Schouten et al., 2004) and the generally observed DIC value in the Arabian Sea OMZ at depth, 0 % (Moos, 2000). This is significantly depleted compared to phytoplankton biomass in the Arabian Sea (-19.8 %, Fontugne & Duplessy, 1978), and would add depleted organic carbon to the sinking POM.

We modelled the contribution of annamox biomass ($\chi_{anammox}$) to SOM using an isotopic mass balance approach (using IsoError; Phillips et al., 2005), which employs uncertainty propagation and error estimates and allows the determination of contributions of likely sources of organic matter to the sediment. The end member value of -19.8 \pm 0.5 % was used for phytoplankton-derived SOM (Fontugne & Duplessy, 1986; Ziegler et al., 2008). This value is slightly lower than the -19 % observed from the surface particulate carbon, but is representative of sedimentary organic matter, as it accounts for the decreased δ^{13} C values of POM caused by degradation of 13 C enriched compounds such as carbohydrates (Close, 2019; Ziegler et al., 2008). More recent values for phytoplankton δ^{13} C are, to the best of our knowledge, not available. Planktonic, sinking organic matter, and organic material produced by heterotrophs also contribute to sedimentary OM. However, this OM value would likely be similar to the organic matter assimilated (Blair et al., 1985; Hayes, 2001). -28.6 \pm 6 % was used for anammox-derived organic carbon, as derived from the δ^{13} C value of BHT' and $\varepsilon_{\text{bm/lipid}}$ of 16 ± 4 % (Schouten et al., 2004; Table 1; uncertainty represents combined standard deviations of δ^{13} C $_{\text{BHT}}$ and $\varepsilon_{\text{lipid/bm}}$). The δ^{13} C value -21.5% of surface sediment C $_{\text{org}}$ in the OMZ was used as the value of the mixture (Table 1):

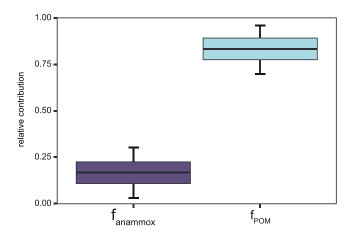


Figure 5. Results from the isotope mass balance model, showing the calculated means, standard error, and confidence intervals for contribution from anammox, and planktonic-derived OM (POM) to the surface sediment at P900.

$$\chi_{anammox} \cdot \delta^{13} C_{anammox} + \chi_{PP} \cdot \delta^{13} C_{PP} = -21.5\% \tag{1}$$

The modelling, assuming the above end member contributions and their statistical uncertainties, yields a proportion of anammox with a mean of approximately 17% (Figure 5), and confirms that, with 95% confidence, anammox contribution to the sedimentary organic matter is between 3 and 30% among the different cores. This suggests that some of the sedimentary organic matter present at P900 is anammox-derived. However, other chemoautotrophic bacteria also present, or suspected to be present, in the Arabian Sea OMZ (e.g. ammonia-oxidizing archaea, Pitcher et al., 2011) possess metabolisms which would lead to different δ^{13} C values, and could therefore be diluting this signal (Hayes, 2001; Pearson, 2010). These results suggest that the contribution of prokaryotic organic material produced in the OMZ to SOM is larger than estimated in this simple, twocomponent mass balance. A three-source model resolving heterotrophic bacteria or degraded OM, anammox bacteria and phytoplankton was solved with SIAR (Parnell et al., 2010) and yielded similar results (not shown).

This estimate, at a first glance, appears large. However, we can convert experimental rates of anammox-mediated ammonia oxidation of 27 - 38 nmol L^{-1} d⁻¹ as determined in the Arabian Sea (Jensen et al., 2011) and alternative rates of 0.24-4.32 nmol L^{-1} d⁻¹ estimated by Ward et al. (2009), to carbon fixed using stoichiometric rates determined for anammox bacteria of 0.07 mol C mol N_2^{-1} (Jetten et al., 2001; Strous et al., 1999; van de Vossenberg et al., 2008). Assuming a production depth of 300 to 900 m water depth, with a maximum at 600 m (Pitcher et al., 2011), anammox production (P) over the whole depth could be estimated using Equation (2) (see Figure S5),

$$P = \frac{(\mathbf{k} \cdot 600)}{2} \tag{2}$$

in which k corresponds to maximum rate at maximum production depth calculated from aforementioned published anammox rates (k equals 138.7 – 9855 μ mol m⁻³ yr⁻¹, Jensen et al., 2011; 87.6 – 1576.8 μ mol m⁻³ yr⁻¹, Ward et al., 2009; Figure S5).

Using this equation, we calculated that water-column anammox bacteria produce up to 3.5 g organic C m⁻² yr⁻¹. These estimates dwarf sedimentary anammox rates, with sedimentary C-fixation by anammox determined to be at 64 pg organic C m⁻² yr⁻¹ in the Arabian Sea (Sokoll et al., 2012), indicating that most of the anammox carbon in the sediment is water-column derived. Given organic carbon accumulation rates (Lengger, Hopmans, Reichart, et al., 2012) of 3 to 5 g C m⁻² yr⁻¹ at P900 and assuming a 17 % anammox contribution to sedimentary organic matter, a maximum of 24 % of annually produced, water-column anammox biomass is preserved in this anoxic setting. Within the sediment from the OMZ, as seen from a sedimentary depth profile at P900 in the OMZ (Figure 3), $\delta^{13}C_{org}$ also shifts to higher values with depth, to a maximum of -20.8 ‰ at 8 cmbsf. However, contrary to oxic degradation, no decrease in SOM contents occurs over the first 14 cm (Figure 3). In this case, the increase in $\delta^{13}C_{org}$ likely reflects input by other sedimentary autotrophs, producing enriched organic carbon. This suggests that the ^{13}C -depleted SOM signal from the OMZ is not completely preserved.

Other chemoautotrophs, where specific biomarkers are not available, could also contribute to the SOM: at depth, the increased availability of CO_2 derived from heterotrophic respiration of the POM pool would result in greater discrimination against ^{13}C (Freeman, 2001; Freeman et al., 1994). An example are lipids of ammonia-oxidising archaea, which were more depleted when deposited in the OMZ than below the OMZ (Table S2), suggesting that some of those were produced in the produced in the OMZ and transported to the sediment (Lengger et al., 2014; Lengger, Hopmans, Reichart, et al., 2012; Schouten et al., 2012). However, as these archaeal lipids present a mixed pelagic and sedimentary signal, quantitative estimates are not possible.



Hence, chemoautotrophic carbon fixation could explain the low δ¹³C values of sedimentary OM observed within the OMZ. As the newly produced organic carbon is more labile than surface-produced organic matter, it would degrade more quickly upon exposure to oxic bottom waters. This poorer preservation of anammox biomass results in a shift back towards the ¹³C signature of the primary photosynthetic production, which is observed as the enrichment in sedimentary $\delta^{13}C_{org}$ with increasing oxygen exposure times (Figure 1e). A likely explanation for this lability of anammox biomass is that, due to the lack of zooplankton in the OMZ, this organic matter is not fecal-pellet packaged or adsorbed to inorganic particles and thus not matrix protected (cf. Burdige, 2007). Wakeham et al. (2002) analysed the biomarker fluxes in sinking particles in the Arabian Sea, and they also found that surface-produced lipids such as alkenones (which would be fecal pellet packaged) were exceptionally well-preserved compared to the total organic carbon of the SPM. Gong and Hollander (1997) also noted an enhanced contribution of bacterial biomass to sediment deposited under anoxic conditions in the Santa Monica Basin, when compared to sediment located in nearby oxygenated bottom waters. Studies examining carbon fluxes in the Arabian Sea (Keil et al., 2016) and the Cariaco basin (Taylor et al., 2001) invoked the addition of chemoautotrophically derived carbon from a midwater source in order to explain the enhanced carbon fluxes observed, even if oxygen depletion and other factors such as the lack of zooplankton were taken into account.

4.3. Quantifying the biological pump and remineralization rates

Models of the biological pump consider primary production, particle fluxes, oxygen concentrations, and respiratory rates. Sinking organic matter is traditionally regarded as a reflection of exported organic material from the photic zone, and export models consider remineralization and C-loss (Cabré et al., 2015; Schlitzer, 2002), based on claims that bacterial contributions do not sink (Buesseler et al., 2007). However, relying on these parameters, estimates of particulate organic carbon flux in the Arabian Sea are generally too low to sustain experimentally observed denitrification, bacterial production, and oxygen deficiency in the Arabian Sea (Anderson & Ryabchenko, 2009; Andrews et al., 2017; Naqvi & Shailaja, 1993; Rixen & Ittekkot, 2005). New mechanisms have been discovered that can explain downward transport of smaller particles and bacteria, such as particle aggregation (Burd & Jackson, 2009), and mixed layer transport mechanisms (Bol et al., 2018), which enable the consideration of the contribution of anammox biomass to sinking OM.

In previous work, the deficit in modelled and observed C-utilization estimated from denitrification was as high as 70 g C m $^{-2}$ y $^{-1}$ (Naqvi & Shailaja, 1993). Including anammox in models of the nitrogen cycle can account for at least 30% or more of the nitrogen losses, as well as accounting for organic carbon production, thereby reconciling some of this budget discrepancy. More recently, the combined inclusion of mesopelagic zooplankton and heterotrophic bacteria into 3D models resulted in much lower bacterial productivity (4.7 g C m $^{-2}$ y $^{-1}$) than observational estimates of 10.4 g C m $^{-2}$ y $^{-1}$ (Anderson & Ryabchenko, 2009). Other recent models, combined with experimental observations of particle fluxes, suggest that at some stations, the lower oxycline is associated with 0.4 to 1.8 g C m $^{-2}$ y $^{-1}$ increases in net carbon fluxes (Roullier et al., 2014). Horizontal transport, also from nepheloid layers, novel transport mechanisms from the mixed layer, and DOM, have previously been invoked as the missing carbon supply. Here, we estimate that an additional 3.5 g m $^{-2}$ y $^{-1}$ of organic carbon are produced by anammox from DIC, with other chemoautotrophic processes possibly adding to this estimate. This also affects estimates of nitrogen losses and has direct implications for the forecasting of OMZs in a warming world.

In paleoceanography, the δ^{13} C values of total organic carbon are commonly used in combination with organic carbon contents in sediment cores to interpret changes in sea surface biogeochemistry and to discern the causes for ocean deoxygenation (Jenkyns, 2010; Meyers, 2014). Negative δ^{13} C excursions are associated with either atmospheric injection of depleted CO_2 and/or increased CO_2 availability (Pagani, 2005; Pancost & Pagani, 2005), or a higher input of terrigenous organic matter. Positive excursions are inferred to reflect increased burial rates of organic carbon, i.e. the removal of 'light' carbon from the ocean-atmosphere reservoir (e.g. Jenkyns, 1985; Kuypers et al., 2002). An increase in TOC coinciding with a negative carbon isotope excursion has been observed in the sedimentary record numerous times. They are a prominent feature in OAE 1a (Leckie et al., 2002), during the PETM (Zachos et al., 2005), and in the Eastern Mediterranean during Sapropel deposition (Meyers & Arnaboldi, 2008). Our results show that if non-sulfidic OMZs expand to impinge on the sediment, a significant amount of ¹³C-depleted organic carbon can be preserved, changing the carbon isotope composition of organic matter by up to – 1.6 ‰ in the case of Arabian Sea surface



sediments, and by -0.5 to -1 ‰ in deeper sediments. When evaluating past events based on bulk δ^{13} C values, the incorporation of chemoautotrophically produced carbon must thus be considered. This also further supports claims that calculations of pCO₂ based on differences between δ^{13} C of organic and inorganic carbon should not rely on bulk organic carbon that includes bacterial contributions, but instead use compound specific δ^{13} C values derived from algae, consistent with previous investigations (Pancost et al., 1999; Pancost et al., 2013; reviewed by Witkowski et al., 2018). By extension, a deviation from the relationship between δ^{13} C bulk OM and δ^{13} C phytane could be an indicator for the chemoautotrophic contribution to sedimentary OM.

Chemoautotrophic metabolism in euxinic water columns, where often very low $\delta^{13}C_{DIC}$ values are observed (van Breugel, Schouten, Paetzel, Nordeide, & Sinninghe Damsté, 2005, van Breugel, Schouten, Paetzel, Ossebaar, & Sinninghe Damsté, 2005, Fry et al., 1991; Volkov, 2000), has previously been invoked as contributing to organic matter in these settings. Anammox, which has been shown to be an important process in the anoxic water column overlying euxinia (Kuypers et al., 2003), also needs to be considered. Anammox also plays an important role in the nitrogen cycle of Mediterranean sapropel events, even in sapropels where euxinia appears to have occurred (Rush et al., 2019), potentially causing a significant contribution of anammox to the depleted $\delta^{13}C_{org}$ values of these sapropels. Previously, a conceptual model invoking chemoautotrophy has been invoked to explain e.g. the End-Permian (Luo et al., 2014) and the Lomagundi isotope excursion (Bekker et al., 2008).

Settings similar to the Arabian Sea, with $\leq 4.5 \,\mu \text{mol kg}^{-1} \, O_2$, occur in the East Pacific and those settings in total occupy an area of 8.45 to 15×10^{12} m² (Karstensen et al., 2008; Hattori, 1983; Paulmier & Ruiz-Pino, 2009), or a volume of 0.46×10^{15} m³ with 1.148×10^{12} m² in contact with the sediment (Helly & Levin, 2004). OMZs are expanding in size and in volume (Queste et al., 2018; Stramma et al., 2010), and anammox bacteria play a key role in these settings, particularly because they tolerate higher oxygen concentrations than denitrifiers (Dalsgaard et al., 2014). The labile organic matter added to the sinking carbon can fuel heterotrophic processes, further exacerbating oxygen depletion, and nitrogen loss by denitrification. For accurate future forecasting, it is imperative to reconcile experimental and model-based estimates. Novel, detailed observations explaining the vertical transport of bacterial biomass, and evidence from the isotopic composition of TOC and biomarkers for chemoautotrophic processes, such as shown here for anammox, will enable the further unravelling of the biogeochemical mechanisms underpinning present, past, and future OMZs.

5. Conclusions

Combining the $\delta^{13}C$ values of BHT', a biomarker from anammox bacteria, and the $\delta^{13}C$ values of sedimentary organic matter allowed us to estimate the contribution of anammox to SOM in the Arabian Sea. This can be as much as 30 % of organic matter deposited within the OMZ. In the sediments underlying the Arabian Sea, $\delta^{13}C$ values of total organic matter can shift by 1 to 2 % due to these bacterial contributions, but it remains unclear how well and under what conditions this signature is preserved. Our results suggest that chemoautotrophs (e.g. anammox bacteria) contribute more than previously believed to the burial of carbon in oxygen deficient zones, and remineralization rates are potentially higher than inferred from organic matter decreases. This implies that, when past occurrences of OMZs are evaluated based on $\delta^{13}C$ values of SOM, a ^{13}C -depleted contribution of bacteria needs to be considered. Chemoautotrophic carbon fixation thus represents a mechanism of CO_2 removal from the pelagic water column, and contributes to fluxes of sinking organic carbon. It explains some of the mismatches in carbon budgets when experimental and modelling estimates are compared - and it should therefore be included in biogeochemical models predicting feedbacks to a warming world.

Data statement

All supporting data is available in Pangaea at https://doi.pangaea.de/10.1594/PANGAEA.908445.

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