Organic carbon production, mineralisation and preservation on the Peruvian margin

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Abstract. Carbon cycling in Peruvian margin sediments (11 and 12° S) was examined at 16 stations, from 74 m water depth on the middle shelf down to 1024 m, using a combination of in situ flux measurements, sedimentary geochemistry and modelling. Bottom water oxygen was below detection limit down to ca. 400 m and increased to 53 µM at the deepest station. Sediment accumulation rates decreased sharply seaward of the middle shelf and subsequently increased at the deep stations. The organic carbon burial efficiency (CBE) was unusually low on the middle shelf (< 20 %) when compared to an existing global database, for reasons which may be linked to episodic ventilation of the bottom waters by oceanographic anomalies. Deposition of reworked, degraded material originating from sites higher up on the slope is proposed to explain unusually high sedimentation rates and CBE (> 60 %) at the deep oxygenated sites. In line with other studies, CBE was elevated under oxygen-deficient waters in the mid-water oxygen minimum zone. Organic carbon rain rates calculated from the benthic fluxes alluded to efficient mineralisation of organic matter in the water column compared to other oxygen-deficient environments. The observations at the Peruvian margin suggest that a lack of oxygen does not greatly affect the degradation of organic matter in the water column but promotes the preservation of organic matter in sediments.

1 Introduction

The Peruvian upwelling forms part of the boundary current system of the Eastern Tropical South Pacific and is one of the most biologically productive regions in the world (Pennington et al., 2006). Respiration of organic matter in subsurface waters leads to the development of an extensive and perennial oxygen minimum zone (Walsh, 1981; Quiñones et al., 2010). Bottom water dissolved oxygen (O₂) concentrations have been measured to be below the analytical detection limit from the shelf down to 400 m (Bohlen et al., 2011). Sediments within this depth interval display organic carbon contents in excess of 15 % (Reimers and Suess, 1983a; Suess et al., 1987; Arthur et al., 1998); much higher than the average continental margin of < 2 % (Seiter et al., 2004). Oxygen-deficient margins like Peru have thus been proposed to be sites of enhanced carbon preservation and petroleum-source rock formation (Demaison and Moore, 1980).

An understanding of the factors that enhance carbon preservation and burial in marine sediments is critical to interpret the sedimentary record and constrain global carbon sources and sinks over geological timescales (Berner, 2004; Wallmann and Aloisi, 2012). Pioneering workers argued that carbon preservation is strongly driven either by the absence of O₂ in the bottom water (Demaison and Moore, 1980) or by higher primary production (Pedersen and Calvert, 1990).

Since then, much work on the biogeochemical characteristics of sediments has been undertaken to better disentangle these factors (Hedges and Keil, 1995; Arthur et al., 1998; Hedges et al., 1999; Keil and Cowie, 1999; Vanderwiele et al., 2009;
Zonneveld et al., 2010; and many others). These studies do broadly indicate that organic matter under oxic bottom waters is in a more advanced state of degradation compared to oxygen-deficient waters. Investigations in the water column have also shown that respiration of organic carbon is significantly reduced in oxygen-deficient waters, leading to elevated carbon fluxes to the sediments (Martin et al., 1987; Devol and Hartnett, 2001; Van Mooy et al., 2002).

Rates of carbon burial and mineralisation on the Peruvian margin have been studied as part of the Collaborative Research Center 754 (Sonderforschungsbereich, SFB 754, www.sfb754.de/en) “Climate-Biogeochemistry Interactions in the Tropical Ocean” (first phase 2008–2011 and second phase 2012–2015). The overall aim of the SFB 754 is to understand the physical and biogeochemical processes that lead to the development and existence of oxygen-deficient regions in the tropical oceans. In this paper, in situ benthic fluxes and sedimentary geochemical data collected during two campaigns to the Peruvian margin at 11 and 12° S are used to summarise our current understanding of carbon cycling in this setting. We address the following questions:

1. What is the rate of organic carbon mineralisation and burial in the sediments down through the oxygen minimum zone (OMZ)? Do these data point toward diminished rates of organic carbon mineralisation in the water column?

2. Which factors determine the carbon burial efficiency at Peru and is there any marked difference for stations underlying oxic and anoxic bottom waters?

2 Study area

Equatorward winds engender upwelling of nutrient-rich equatorial subsurface water along the Peruvian coast (Fiedler and Talley, 2006). Upwelling is most intense between 5 and 15° S where the shelf narrows (Quiñones et al., 2010). The sampling transects in this study at 11 and 12° S are located within the same upwelling cell (Suess et al., 1987). The highest rates of primary productivity (1.8–3.6 g C m^{-2} d^{-1}) are 6 months out of phase with upwelling intensity due to the deepening of the mixed layer during the upwelling period (Walsh, 1981; Echevin et al., 2008; Quiñones et al., 2010). Austral winter and spring is the main upwelling period, with interannual variability imposed by the El Niño–Southern Oscillation (ENSO) (Morales et al., 1999). The lower vertical limit of the OMZ is around 700 m water depth off Peru (O_2 < 20 µmol kg^{-1}; Fuenzalida et al. 2009). The mean depth of the upper boundary of the OMZ on the shelf at 11 and 12° S is around 50 m (Gutiérrez et al., 2008), but deepens to ca. 200 m or more during ENSO years (e.g. Levin et al., 2002). At these times, dissolved O_2 on the shelf can vary between 0 and 100 µM within a matter of days to weeks as opposed to several months during weaker ENSO events (Gutiérrez et al., 2008; Noffke et al., 2012).

Sediments at 11 and 12° S are generally diatomaceous, rapidly accumulating muds (Suess et al., 1987, and many others). Grain size analysis shows that clay/silt fractions are highest on the shelf and in mid-waters (> 80 %), whereas the sand content is highest (40 %) in deeper waters (Mosch et al., 2012). The sediments can thus be described as sandy mud to slightly sandy mud (Flemming, 2000). The distribution of sediment on the margin is influenced by resuspension, winnowing and lateral particle transport due to bottom currents and breaking of internal waves on the slope (Arthur et al., 1998; Levin et al., 2002; Mosch et al., 2012). Surface particulate organic carbon (POC) content is high in mid-waters (15 to 20 %) with lower values (5 to 10 %) on the shelf and in deep waters (Böning et al., 2004). δ^{13}C analysis and other geochemical indicators confirm that the organic matter at this latitude is almost entirely of marine origin (Arthur et al., 1998; Reimers and Suess, 1983b; Levin et al., 2002; Gutiérrez et al., 2009).

The sediments down to around 400 m are notably cohesive, ranging from dark olive green to black in colour with no surface-oxidised layer (Bohlen et al., 2011; Mosch et al., 2012). The surface is colonised by dense, centimetre-thick mats of gelatinous sheaths containing microbial filaments of the large sulfur oxidising bacteria Thioploca spp. (Hennrichs and Farrington, 1984; Arntz et al., 1991). These bacteria glide vertically through the sediments to access sulfide, which they oxidise using nitrate stored within intracellular vacuoles (Jørgensen and Gallardo, 2006). The bacterial density varies with time on the shelf, depending on the bottom water redox conditions (Gutiérrez et al., 2008). Spiionid polychaetes (ca. 2 cm length) have been observed in association with the mats (Mosch et al., 2012). The biomass of macrofauna generally tends to be highest in the OMZ but with low species richness, dominated by polychaetes and oligochaetes (Levin et al., 2002). At the lower boundary of the OMZ, high abundances of epibenthic megafauna such as ophiuroids as well as echinoderms, pennatulaceans, Porifera, crustaceans, gastropods and echinoderms have been observed (Levin et al., 2002; Mosch et al., 2012). Sediments here are olive green throughout with a thin upper oxidised layer light green/yellow in colour (Bohlen et al., 2011; Mosch et al., 2012).

For the purposes of this study, we divide the Peruvian margin into three zones broadly reflecting bottom water O_2 distributions and sedimentary POC content: (i) the middle and outer shelf (< ca. 200 m, POC 5 to 10 %, O_2 < detection limit (dl, 5 µM) at time of sampling) where non-steady conditions are occasionally driven by periodic intrusion of oxygenated bottom waters; (ii) the OMZ (ca. 200 to 450 m, POC 10 to 20 %, O_2 predominantly < dl); and (iii) the deep stations below the OMZ with oxygenated bottom water (POC ≤ ca. 5 % and O_2 > dl).
3 Material and methods

3.1 Flux measurements and sediment sampling

We present data from six stations along 11° S sampled during expedition M77 (cruise legs 1 and 2) in November/December 2008 and ten stations along 12° S during expedition M92 (leg 3) in January 2013 (Fig. 1). Both campaigns took place during austral summer, i.e. the low upwelling season, and under neutral or negative ENSO conditions (http://www.cpc.ncep.noaa.gov). With the exception of the particulate phases, the geochemical data and benthic modelling results from the 11° S transect have been published previously (Bohlen et al., 2011; Scholz et al., 2011; Mosch et al., 2012; Noffke et al., 2012). Data from 12° S are new to this study.

In situ fluxes were measured using data collected from Biogeochemical Observatories, BIGO (Sommer et al., 2008; Dale et al., 2014). BIGO landers contained two circular flux chambers (internal diameter 28.8 cm, area 651.4 cm²). One lander at 11° S, BIGO–T, contained only one benthic chamber. Each chamber was equipped with an optode to monitor dissolved O₂ concentrations. A TV-guided launching system allowed smooth placement of the observatories on the seafloor. Two hours (11° S) and 4 h (12° S) after the landers were placed on the seafloor, the chamber(s) were slowly driven into the sediment (∼30 cm h⁻¹). During this initial period, the water inside the flux chamber was periodically replaced with ambient bottom water. After the chamber was driven into the sediment (∼10 cm), the chamber water was again replaced with ambient bottom water to flush out solutes that might have been released from the sediment during chamber insertion. The water volume enclosed by the benthic chamber ranged from 7.8 to 18.5 L and was mixed using a 5 cm stirrer bar at 140 rpm located 10–15 cm above the sediment surface. During the BIGO-T experiments, the chamber water was replaced with ambient bottom water halfway through the deployment period to restore outside conditions and then re-incubated.

Four (11° S) or eight (12° S) sequential water samples were removed periodically with glass syringes (volume of each syringe ∼47 mL) to determine fluxes of solutes across...
the sediment–water interface. For BIGO-T, four water samples were taken before and after replacement of the chamber water. The syringes were connected to the chamber using 1 m long Vygon tubes with an internal volume of 6.9 mL. Prior to deployment, these tubes were filled with distilled water, and great care was taken to avoid enclosure of air bubbles. Concentrations were corrected for dilution using measured chloride concentrations in the syringes and bottom water. Water samples for gas measurements (12\(^{24}\)S) were taken at four regular time intervals using 80 cm long glass tubes (internal volume ca. 15 mL). An additional syringe water sampler (four or eight sequential samples) was used to extract ambient bottom water samples from 30–40 cm above the seafloor. The benthic incubations were conducted for time periods ranging from 17.8 to 33 h. Immediately after retrieval of the observatories, the water samples were transferred to the onboard cool room set to the average bottom water temperature on the margin (8 °C) for further processing and sub-sampling. Benthic fluxes were estimated from linear regressions of the concentration-time data and corrected for the volume to surface area ratio of the chamber. The volume was estimated on board using the mean height of water above the sediments in the recovered chambers.

Sediment samples for analysis were taken using multiple-corners (MUC) deployed adjacent to the BIGO sites. Retrieved cores were immediately transferred to the cool room and processed within a few hours. Sub-sampling for redox sensitive constituents was performed under anoxic conditions using an argon-filled glove bag. Sediment samples for porosity analysis were transported to the onshore laboratory in air-tight containers at 8 °C. Samples for porewater extraction were centrifuged at 4500 rpm for 20 min. Prior to analysis, the supernatant was filtered with cellulose acetate Nuclepore® filters (0.2 µm) inside the glove bag. The centrifugation tubes with the remaining solid phase of the sediment were stored at −20 °C for further analysis on shore. Additional samples for bottom water analysis were taken from the water overlying the sediment cores.

3.2 Analytical details

Dissolved oxygen concentrations in the water column were measured using a Seabird SBE43 polarographic membrane oxygen sensor mounted on a SeaBird 911 CTD rosette system. The sensors were calibrated against discrete samples collected from the water column on each CTD cast and analysed on board using Winkler titration with a detection limit of ca. 5 µM. The optodes inside the benthic chambers were calibrated by vigorously bubbling unfiltered bottom seawater with air or argon for 20 min and calibrated using Winkler. We broadly define O\(_2\) concentrations below the detection limit of the Winkler analysis as anoxic, whilst noting that sub-micromolar levels have been measured in the OMZ using microsensors (Kalvelage et al., 2013).

Ammonium (NH\(_4^+\)) was measured on board using standard photometric techniques with a Hitachi U2800 photometer (Grashoff et al., 1999). The detection limit was 1 µM and the precision of the analyses was 5 µM. Total alkalinity (TA) was determined by direct titration of 1 mL porewater with 0.02 M HCl using a mixture of methyl red and methylene blue as an indicator and bubbling the titration vessel with argon gas to strip CO\(_2\) and hydrogen sulfide. The analysis was calibrated using IAPSO seawater standard, with a precision and detection limit of 0.05 meq L\(^{-1}\). Ion chromatography (Methrom 761) was used to determine sulfate (SO\(_4^{2-}\)) in the onshore laboratory with a detection limit of < 100 µM and precision of 200 µM. Major cations were determined by ICP-AES with a detection limit and precision as given by Haffert et al. (2013).

The partial pressure of CO\(_2\) (pCO\(_2\)) was analysed in the benthic chambers at 12\(^{24}\)S by passing the sample from the glass tubes (without air contact) through the membrane inlet of a quadrupole mass spectrometer (GAM200 IPI Instruments, Bremen). The samples were analysed sequentially, flushing with distilled water between samples. Standards of 300, 500, 1000 and 5000 ppm CO\(_2\) were prepared by sparging filtered seawater from the bottom water from each station using standard bottles of CO\(_2\) of known concentration at in situ temperature for 30 min. Calibration was performed before and after analysis of the samples from each site. The relative precision of the measurement was < 3 %.

Wet sediment samples for analysis of POC and particulate organic nitrogen (PON) were freeze-dried in the home laboratory and analysed using a Carlo-Erba element analyser (NA 1500). POC content was determined after acidifying the sample with HCl (0.25 N) to release the inorganic components as CO\(_2\). Weight percent of total carbon was determined using samples without acidification. Inorganic carbon was determined by weight difference between the total and organic carbon. The precision and detection limit of the POC analysis was 0.04 and 0.05 dry weight percent (% C), respectively. The precision and detection limit of the inorganic carbon analysis was 2 and 0.1 % C, respectively. Porosity was determined from the weight difference of the wet and freeze-dried sediment. Values were converted to porosity (water volume fraction of total sediment) assuming a dry sediment density of 2 g cm\(^{-3}\) (Böning et al., 2004) and seawater density of 1.023 g cm\(^{-3}\). The analysis of total aluminium (Al) concentrations in digestion solutions was carried out using an inductively coupled plasma optical emission spectrometer (ICP-OES, VARIAN 720-ES) following the procedure described by Scholz et al. (2011).

Additional samples from adjacent MUC liners taken from the same cast were used for the determination of downcore profiles of unsupported (excess) \(^{210}\)Pb\(_{ex}\) activity by gamma counting. This approach includes the monitoring of the main peaks of anthropogenic deposition of \(^{241}\)Am during the 1950s (test of nuclear weapons) as an independent time marker. Between 5 and 34 g of freeze-dried and ground
sediment, each averaging discrete 2 cm depth intervals, was embedded into a 2-phase epoxy resin (West System Inc.), all in the same counter-specific calibrated disc geometry (2 inch diameter). Following Mosch et al. (2012), a low-background coaxial Ge(Li) planar detector (LARI, University of Göttingen) was used to measure total $^{210}\text{Pb}$ via its gamma peak at 46.5 keV and $^{226}\text{Ra}$ via the grandaughter $^{214}\text{Pb}$ at 352 keV. Prior to analysis, $^{226}\text{Ra}$ and $^{214}\text{Pb}$ in the gas-tight embedded sediment were allowed to equilibrate for at least 3 weeks. To determine $^{210}\text{Pb}_{ss}$, the measured total $^{210}\text{Pb}$ activity of each sample was corrected by subtracting its individual $^{226}\text{Ra}$ activity, assuming post-burial closed-system behaviour. Uncertainty of the $^{210}\text{Pb}_{ss}$ data was calculated from the individual measurements of $^{210}\text{Pb}$ and $^{226}\text{Ra}$ activities using standard propagation rules. The relative error of the measurements ($2\sigma$) ranged between 8 and 58 %.

### 3.3 Calculation of dissolved inorganic carbon (DIC) fluxes

DIC concentrations in the benthic chambers at 12° S were calculated from the concentrations of TA and $p\text{CO}_2$ using the equations and equilibrium coefficients given by Zeebe and Wolf-Gladrow (2001). Since four samples for $p\text{CO}_2$ were taken using the glass tubes versus eight samples for TA analysis in the syringes, each successive pair of TA data were averaged for calculating DIC (see Supplement). A constant salinity (35 psu), total boron concentration (0.418 mM) and seawater density (1.025 kg L$^{-1}$) were assumed. For the shelf stations, where sulfide was released from the sediment (Sommer et al., unpub. data), corrections were made for the contribution of HS$^-$ to TA using the relevant equilibrium constants (Zeebe and Wolf-Gladrow, 2001). DIC fluxes were calculated from the concentrations as described above.

### 3.4 Determination of sediment accumulation rates

Particle-bound $^{210}\text{Pb}_{ss}$ is subject to mixing in the upper sediment layers by the movement of benthic fauna. The distribution of $^{210}\text{Pb}_{ss}$ can thus be used to determine bioturbation coefficients as well as sedimentation rates using a reaction-transport model. We simulated the activity of $^{210}\text{Pb}_{ss}$ in Bq g$^{-1}$ using a steady-state numerical model that includes terms for sediment burial, mixing (bioturbation), compaction and radioactive decay:

\[
\frac{D}{Dx} \left( (1 - \varphi(x)) \times \rho \times \frac{d^{210}\text{Pb}_{ss}(x)}{dt} \right) = \\
\frac{D}{Dx} \left( (1 - \varphi(x)) \times \rho \times D_B(x) \times \frac{d^{210}\text{Pb}_{ss}(x)}{dx} \right) \\
- \frac{D}{Dx} \left( (1 - \varphi(x)) \times \rho \times v_s(x) \times 2^{10}\text{Pb}_{ss}(x) \right) \\
+ (1 - \varphi(x)) \times \rho \times \lambda^{210} \times \text{Pb}_{ss}(x). \quad (1)
\]

In this equation, $t$ (yr) is time, $x$ (cm) is depth below the sediment–water interface, $\varphi(x)$ (dimensionless) is porosity, $v_s(x)$ (cm yr$^{-1}$) is the burial velocity for solids, $D_B(x)$ (cm$^2$ yr$^{-1}$) is the bioturbation coefficient, $\lambda$ (0.03114 yr$^{-1}$) is the decay constant for $^{210}\text{Pb}_{ss}$ and $\rho$ (2.0 g cm$^{-3}$) is the bulk density of solid particles.

Porosity was described using an exponential function assuming steady-state compaction:

\[
\varphi(x) = \varphi(L) + (\varphi(0) - \varphi(L)) \times \exp \left( - \frac{x}{z_{por}} \right), \quad (2)
\]

where $\varphi(0)$ is the porosity at the sediment–water interface, $\varphi(L)$ is the porosity of compacted sediments and $z_{por}$ (cm) is the attenuation coefficient. These parameters were determined from the measured data at each station (Table S2 in Supplement).

Sediment compaction was considered by allowing the sediment burial velocity to decrease with sediment depth:

\[
v_s(x) = \frac{\omega_{acc} \times (1 - \varphi(L))}{1 - \varphi(x)}, \quad (3)
\]

where $\omega_{acc}$ corresponds to the sediment accumulation rate of compacted sediments.

The decrease in bioturbation intensity with depth was described with a Gaussian-type function (Christensen, 1982):

\[
D_B(x) = D_B(0) \times \exp \left( - \frac{v_s^2}{2 \times x_s^2} \right), \quad (4)
\]

where $D_B(0)$ (cm$^2$ yr$^{-1}$) is the bioturbation coefficient at the sediment–water interface and $x_s$ (cm) is the bioturbation halving depth.

The flux continuity at the surface serves as the upper boundary condition:

\[
F(0) = (1 - \varphi(0)) \times \rho \times \\
\left( v_s(0) \times 2^{10}\text{Pb}_{ss}(0) - D_B(0) \times \frac{d^{210}\text{Pb}_{ss}(x)}{dx}\right|_0 \right), \quad (5)
\]

where $F(0)$ is the steady-state flux of $^{210}\text{Pb}_{ss}$ to the sediment surface (Bq cm$^{-2}$ yr$^{-1}$). The influx of $^{210}\text{Pb}_{ss}$ was determined from the measured integrated activity of $^{210}\text{Pb}_{ss}$ multiplied by $\lambda$:

\[
F(0) = \lambda \times \rho \int_0^\infty 2^{10}\text{Pb}_{ss}(x) \times (1 - \varphi(x))dx. \quad (6)
\]

$^{210}\text{Pb}_{ss}$ was present down to the bottom of the core at the 74 m station (12° S), implying rapid burial rates. Here, $F(0)$ was adjusted until a fit to the data was obtained.

A zero gradient (Neumann) condition was imposed at the lower boundary at 50 cm (100 cm for the shallowest stations at 12° S). At this depth, all $^{210}\text{Pb}_{ss}$ will have decayed.

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for the burial rates encountered on the Peruvian margin. The model was initialised using low and constant values for $^{210}$Pb$_{xs}$ in the sediment column. Solutions were obtained using the numerical solver NDSolve in MATHEMATICA 9 with a mass conservation > 99%.

The adjustable parameters ($\omega_{acc}$, $D_{B}$, $x_{s}$) were constrained by fitting the $^{210}$Pb$_{xs}$ data. Unsupported $^{210}$Pb measurements were not made at the 101 and 244 m station (12° S) and sedimentation rates here were estimated from adjacent stations. Parameters and boundary conditions for simulating $^{210}$Pb$_{xs}$ at 12° S are given in Table S2 and in Bohlen et al. (2011) for 11° S. For some cores, the subsampling strategy revealed the detection of the anthropogenic enrichment peak of nuclide $^{241}$Am (co-analysed on 60 keV). This provides an independent time marker in the profiles and potential validation of the radiometric age model deduced from the $^{210}$Pb$_{xs}$-based sediment accumulation rates.

### 3.5 Diagenetic modelling of POC degradation

A steady-state 1–D numerical reaction-transport model was used to simulate the degradation of POC in surface sediments at all stations. The model developed for 12° S is based on that used to quantify benthic N fluxes at 11° S by Bohlen et al. (2011) with modifications to account for benthic denitrification by foraminifera (Glock et al., 2013).

The basic model framework follows Eq. (1). Solutes were transported by molecular diffusion, sediment accumulation (burial) and non-local transport by burrowing organisms (bioirrigation) in oxygenated sediments below the OMZ. Solid transport by burial and bioturbation was parameterised using the results of the $^{210}$Pb$_{xs}$ model. Model sensitivity analysis based on solute fluxes showed that bioirrigation rates were very low.

The model includes a comprehensive set of redox reactions that are ultimately driven by POC mineralisation. POC was degraded by aerobic respiration, denitrification, iron oxide reduction, sulfate reduction and methanogenesis. Manganese oxide reduction was not considered due to negligible total manganese in the sediment (Scholz et al., 2011). The rate of each carbon degradation pathway was determined using Michaelis–Menten kinetics based on traditional approaches (e.g. Boudreau, 1996). DIC is produced by POC degradation only, that is, carbonate dissolution or precipitation are not included (see Results).

The total rate of POC degradation was constrained using a nitrogen-centric approach based on the relative rates of transport and reactions that produce/consume NH$_4$+. The procedure follows a set of guidelines that is outlined fully in Bohlen et al. (2011). The modelled POC mineralisation rates for 11° S were constrained using both porewater concentration data and in situ flux measurements of NO$_3$, NO$_2$ and NH$_4$+. Dissolved O$_2$ flux data were used as an additional constraint at the deeper stations. The POC degradation rates at 12° S were further constrained from the measured DIC fluxes. The model output includes concentration profiles, benthic fluxes and reaction rates, which are assumed to be in steady state. Note, however, that the bottom waters on the middle shelf at 12° S were temporarily depleted in NO$_3$ and NO$_2$ at the time of sampling. Although this leads to uncertainties in the rate of nitrate uptake by Thiothrix, POC degradation rates remain well-constrained from the DIC fluxes.

The model was solved in the same way as described for $^{210}$Pb$_{xs}$. The sediment depth ranged from 50 to 100 cm depending on the station (Boudreau, 1996). Measured solute concentrations and known or estimated particulate fluxes to the seafloor served as upper boundary conditions (Bohlen et al., 2011). At the lower boundary, a Neumann (zero flux) boundary was generally implemented. A steady-state solution was obtained (invariant concentrations with time and sediment depth) with a mass conservation > 99%.

### 3.6 Pelagic modelling of primary production

Primary production was estimated using the high-resolution physical-biogeochemical model (ROMS-BioEBUS) in a configuration developed for the Eastern Tropical Pacific (Montes et al., 2014). It consists of the hydrodynamic model ROMS (Regional Ocean Model System; Shchepetkin and McWilliams, 2003) coupled with the BIOgeochemical model developed for the Eastern Boundary Upwelling Systems (BioEBUS, Gutknecht et al., 2013). BioEBUS describes the pelagic distribution of O$_2$ and the N cycle under a range of redox conditions with twelve compartments: phytoplankton and zooplankton split into small (flagellates and ciliates, respectively) and large organisms (diatoms and copepods, respectively); detritus; NO$_3$; NO$_2$; NH$_4$; dissolved organic N; and a parameterisation to determine nitrous oxide (N$_2$O) production (Sutharalingam et al., 2000, 2012).

The model configuration covers the region between 4° N and 20° S and from 90° W to the west coast of South America. The model horizontal resolution is 1/9° (ca. 12 km) and has 32 vertical levels that are elongated toward the surface to provide a better representation of shelf processes. The model was forced by heat and freshwater fluxes derived from COADS ocean surface monthly climatology (Da Silva et al., 1994) and by the monthly wind stress climatology computed from QuikSCAT satellite scatterometer data (Liu et al., 1998). The three open boundary conditions (northern, western and southern) for the dynamic variables (temperature, salinity and velocity fields) were extracted from the Simple Ocean Data Assimilation (SODA) reanalysis (Carton and Giese, 2008). Initial and boundary conditions for biogeochemical variables were extracted from the CSIRO Atlas of Regional Seas (CARS 2009; for NO$_3$ and O$_2$) and SeaWiFS (O’Reilly et al., 2000; for chlorophyll $a$). Other biogeochemical variables were computed following Gutknecht et al. (2013) and Montes et al. (2014). Monthly chlorophyll climatology from SeaWiFS was used to generate phytoplank-
ton concentrations, which were then extrapolated vertically from the surface values using the parameterisation of Morel and Berthon (1989). Based on Koné et al. (2005), a cross-shore profile following in situ observations was applied to zooplankton, with higher concentrations near the coast.

The simulation period was 18 years: the first 13 years considered the hydrodynamics only, and then the biogeochemical model was coupled for the following 5 years. The coupled model reached a statistical equilibrium after 4 years. The data presented here correspond to the final simulation year. Details of model configuration and validation are described by Montes et al. (2014).

Primary production (PP) was computed as the sum of the production supported by NO$_3^-$ and NO$_2^-$ uptake and regenerated production of NH$_4^+$ uptake by nano- and microphytoplankton (Gutknecht et al., 2013). Rates (in N units) were calculated for the station locations listed in Table 1 by integrating over the euphotic zone. The atomic Redfield C:N ratio (106/16). Redfield et al. (1963) was used to convert PP into carbon units.

4 Results

4.1 Sediment appearance

Bottom sediments at 12° S were very similar to those at 11° S (see Sect. 2). The sediments down to ca. 300 m were cohesive, dark-olive anoxic mud (Gutiérrez et al., 2009; Bohlen et al., 2011; Mosch et al., 2012). Porosity was high on the shelf and in the OMZ (> 0.9) decreasing to < 0.8 at the deepest stations (Fig. S1 in Supplement and Table 1). Porewater had a strong sulfidic odour, especially in the deeper layers. Shelf and OMZ sediments were colonised by mats of large filamentous bacteria, presumably Thioploca spp. (Gallardo, 1977; Henrichs and Farrington, 1984). Surface coverage by bacterial mats was 100% on the shelf and decreased to roughly 40% by 300 m where the bacteria formed patches several decimetres in diameter. Mat density was much lower at 11° S, not exceeding 10% coverage (Mosch et al., 2012). Thioploca trichomes extended 2 cm into the overlying water to access bottom water NO$_3^-$ (cf. Huettel et al., 1996) and were visible down to a depth of ca. 20 cm at the mat stations. Polychaetes and oligochaetes were also present on the shelf, but not at the deeper stations within the OMZ. Despite anoxic bottom waters, no mats were visible at St. 8 (409 m, 12° S). Sediments here consisted of hard grey clay underlying a 2–3 cm porous surface layer that was interspersed with cm-sized phosphorite nodules. The upper layer contained large numbers of live foraminifera that were visible to the naked eye (J. Cardich et al., unpublished data). Similar foraminiferal “sands” containing phosphorite granules were noted at 11° S, in particular below the OMZ (Mosch et al., 2012). Phosphorite sands on the Peruvian margin were found in areas of enhanced sediment reworking by bottom currents and the breaking of internal waves on the seafloor (Suess, 1981; Glenn and Arthur, 1988; Mosch et al., 2012). Below the OMZ, macrofauna were more prevalent and included harpacticoids, amphipods, oligochaetes and large polychaetes.
4.2 Sediment mixing and accumulation rates

At most stations, $^{210}$Pb, distributions decreased quasi-
exponentially and showed little evidence of intense, deep mixing by bioturbation (Fig. 2 and Bohlen et al., 2011); a feature that is supported by the lack of large bioturbating organisms in and below the OMZ. The highest bioturbation co-

efficient determined by the model was $4 \text{ cm}^2 \text{ yr}^{-1}$ for St. 3 at $12^\circ \text{ S}$ (Table S2).

Mass accumulation rates (MAR) derived from $^{210}$Pb, modelling (Fig. 3a) were similar to values reported previously (Reimers and Suess, 1983c). Rates were extremely high at the shallow stations (1200 and 1800 g m$^{-2}$ yr$^{-1}$ at 11 and $12^\circ \text{ S}$, respectively). These are a factor of 2–3 times higher than measured elsewhere on the transects and
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Table 2. Measured and modelled carbon fluxes and burial efficiencies.

<table>
<thead>
<tr>
<th>12°S Transect</th>
<th>Inner Shelf</th>
<th>Outer Shelf</th>
<th>OMZ</th>
<th>Below OMZ</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>St. 1</td>
<td>St. 2</td>
<td>St. 3</td>
<td>St. 4</td>
</tr>
<tr>
<td>Water depth, m</td>
<td>74</td>
<td>101</td>
<td>128</td>
<td>142</td>
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<tr>
<td>Sediment accumulation rate (o&lt;sub&gt;sh&lt;/sub&gt;), cm yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.45</td>
<td>0.32</td>
<td>0.2</td>
<td>0.04</td>
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<td>Mass accumulation rate (MAR), g m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>1800</td>
<td>768</td>
<td>600</td>
<td>128</td>
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<td>POC content at 10 cm (POC&lt;sub&gt;10&lt;/sub&gt;), %&lt;sup&gt;c&lt;/sup&gt;</td>
<td>3.3</td>
<td>3.8</td>
<td>7.2</td>
<td>8.6</td>
</tr>
<tr>
<td>POC accumulation rate at 10 cm (POC&lt;sub&gt;AR10&lt;/sub&gt;), g C m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>60</td>
<td>29</td>
<td>34</td>
<td>11</td>
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<tr>
<td>Benthic DIC flux (J&lt;sub&gt;DIC&lt;/sub&gt;), mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>65.9 ± 21</td>
<td>27.9 ± 4.2</td>
<td>20.4 ± 7</td>
<td>8.0 ± 0.4</td>
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<tr>
<td>POC rain rate (RRPOC), mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>79.5 ± 33</td>
<td>34.2 ± 11</td>
<td>28.2 ± 12</td>
<td>10.5 ± 3</td>
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<tr>
<td>Carbon burial efficiency at 10 cm (CBE), %&lt;sup&gt;δ&lt;/sup&gt;</td>
<td>17 ± 7</td>
<td>19 ± 6</td>
<td>28 ± 12</td>
<td>24 ± 7</td>
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<table>
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<th>11°S Transect</th>
<th>St. 1</th>
<th>St. 2</th>
<th>St. 3</th>
<th>St. 4</th>
<th>St. 5</th>
<th>St. 6</th>
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<tr>
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<tr>
<td>Sediment accumulation rate (o&lt;sub&gt;sh&lt;/sub&gt;), cm yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.3</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Mass accumulation rate (MAR), g m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>1200</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>POC content at 10 cm (POC&lt;sub&gt;10&lt;/sub&gt;), %&lt;sup&gt;n&lt;/sup&gt;</td>
<td>2.4</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>POC accumulation rate at 10 cm (POC&lt;sub&gt;AR10&lt;/sub&gt;), g C m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>29</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benthic DIC flux (J&lt;sub&gt;DIC&lt;/sub&gt;), mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>31</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>POC rain rate, mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>8.2</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Carbon burial efficiency at 10 cm, %&lt;sup&gt;k&lt;/sup&gt;</td>
<td>18</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Primary production from ROMS, mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>122</td>
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</tbody>
</table>

a Determined from <sup>210</sup>Pb (see Table S2 in the Supplement). Sedimentation rates at St. 2 (101 m) and St. 6 (250 m) were not measured and instead estimated from the neighbouring stations. o<sub>sh</sub> has a 20 % uncertainty. b Calculated as o<sub>sh</sub> × (1 − ρ) × ρ× 10000 (ρ = dry solid density, 2 g cm<sup>-3</sup>). c For St. 8 at 12°S (409 m) the content at 3 cm was taken since the underlying sediment is old, non-accumulating clay. For the OMZ stations the mean POC content in the upper 10 cm was used instead. This was approximated as follows: POC<sub>10</sub> (%) = ∫<sub>0</sub><sup>10</sup> POC(x) dx where POC(x) is in %. d Calculated analogously to footnote c. e Mean fluxes calculated from the in situ TA and pCO<sub>2</sub> measurements in two benthic chambers. No pCO<sub>2</sub> measurements were made at 11°S. Errors represent 50 % of the difference of the two fluxes. f Calculated as POC<sub>AR10</sub> (in mmol m<sup>-2</sup> d<sup>-1</sup>)+J<sub>DIC</sub>. g Calculated as POC<sub>AR10</sub> (in mmol m<sup>-2</sup> d<sup>-1</sup>)/RRPOC × 100 %. g Errors were calculated using standard error propagation rules assuming a 20 % uncertainty in o<sub>sh</sub> and POC<sub>10</sub>. h Calculated analogously to footnote g using modelled data. i Calculated analogously to footnote f using modelled data. j Calculated analogously to footnote g using modelled data. k Determined from <sup>210</sup>Pb modelling (see Bohlen et al., 2011). l For the OMZ stations, the mean POC content in the upper 10 cm was used (see footnote c). m Calculated as the depth-integrated POC degradation rate (Bohlen et al., 2011).
3–4 times higher than the global shelf average of 500 g m\(^{-2}\) yr\(^{-1}\) (Burwicz et al., 2011). They corresponded to sedimentation rates (\(\omega_{\text{acc}}\)) of 0.45 and 0.3 cm yr\(^{-1}\) (Table 2). Beyond the middle shelf, MAR decreased sharply to 132 and 44 g m\(^{-2}\) yr\(^{-1}\) at St. 2 (11° S) and St. 8 (12° S), respectively. These latter values are associated with measurable \(^{210}\text{Pb}_{\text{xs}}\) in the upper 3 cm only and thus indicative of sediment winnowing or resuspension, as mentioned above. A relatively low MAR of 128 g m\(^{-2}\) yr\(^{-1}\) was also determined for St. 4 at 12° S compared to the neighbouring stations. MAR and \(\omega_{\text{acc}}\) tended to be higher at the deep oxygenated stations compared to the OMZ stations, with \(\omega_{\text{acc}}\) of 0.06 cm yr\(^{-1}\) at St. 10 (12° S) and 0.05 cm yr\(^{-1}\) at St. 5 (11° S). Aluminium accumulation showed similar trends to MAR, with highest values on the shelf and a pronounced increase below the OMZ (Fig. 3b). For some cores, peaks in \(^{210}\text{Pb}\) activities could be detected, which provide independent validation of \(\omega_{\text{acc}}\) derived from the \(^{210}\text{Pb}_{\text{xs}}\) systematics (Fig. 2 and Appendix A).

### 4.3 Geochemistry

Dissolved \(\text{O}_2\) concentrations in the water column reveal the vertical extent of the OMZ and the presence of oxygen-deficient water overlying the upper slope sediments at both latitudes (Fig. 1). Qualitatively, geochemical solute profiles in the sediments from 11 and 12° S are typical for continental margin settings (Bohlen et al., 2011; Fig. 4). The model was able to accurately simulate the geochemical profiles along both transects. Sediment porewater concentrations of \(\text{NH}_4^+\) and alkalinity were highest on the shelf and decreased with water depth. Conversely, \(\text{SO}_4^{2-}\) depletion was more extensive at the shallower stations. \(\text{SO}_4^{2-}\) also showed a much stronger depletion on the shelf at 12° S compared to 11° S, leading to the formation of a methanogenic layer below 65 cm. These trends confirm general expectations that less reactive organic material reaches the seafloor as water depth increases (e.g. Suess, 1980; Levin et al., 2002).

### 4.4 Organic carbon distributions and burial rates

Surface POC content at 12° S was lowest (ca. 5 %) on the middle shelf and below the OMZ (Fig. 4). At these stations, POC decreased in the upper 10 cm and reached asymptotic values at around 10 cm, where refractory components dominated (Reimers and Suess, 1983a). PON showed the same qualitative trends. Maximal POC contents of ca. 17 % were measured inside the OMZ and are typical for the Peruvian margin (Suess, 1981). Here, POC showed a marked change at around 15 to 20 cm depth (Fig. 4). This may reflect the regime shift in the Peruvian OMZ during the Little Ice Age circa 1820 AD, caused by a northward displacement of the Intertropical Convergence Zone (Gutiérrez et al., 2009). These features were also present at St. 4 and 5 on the outer shelf. The steady-state model does not capture centennial changes in the OMZ conditions suggested by the POC profiles. Very similar trends were observed at 11° S, implying that organic matter distributions are qualitatively and quantitatively driven by the same first-order processes at both latitudes.

POC accumulation rates for the middle shelf and deep stations were calculated from the mass accumulation rates and POC content at 10 cm (Table 2). Due to recent variations in POC content for the OMZ stations, the average POC content in the upper 10 cm was used. For St. 8 at 12° S (409 m), POC accumulation was calculated at 3 cm, since the underlying sediment was old, non-accumulating clay. The highest carbon accumulation rates were calculated for the middle shelf at 12° S (60 g m\(^{-2}\) yr\(^{-1}\); Table 2). POC accumulation generally decreased with increasing water depth at 12° S with relatively low values at St. 4 (11 g m\(^{-2}\) yr\(^{-1}\)) and St. 8 (2 g m\(^{-2}\) yr\(^{-1}\)). Accumulation rates were more variable for the 11° S transect.
Table 3. Mean rates of organic carbon accumulation and primary production on the Peruvian margin from this study compared to global averages by Burdige (2007) and Sarmiento and Gruber (2006). Units: mmol m$^{-2}$ d$^{-1}$.

<table>
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<tr>
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<tbody>
<tr>
<td>POC accumulation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shelf (0–200 m)</td>
<td>13.8</td>
<td>9.9</td>
<td>4.0</td>
<td>–</td>
</tr>
<tr>
<td>Upper slope (200–1000 m)</td>
<td>7.2</td>
<td>2.8</td>
<td>1.0</td>
<td>–</td>
</tr>
<tr>
<td>Total margin (0–1000 m)</td>
<td>6.8</td>
<td>6.1</td>
<td>2.9 (1.2)</td>
<td>1.4</td>
</tr>
<tr>
<td>Primary production (0–1000 m)</td>
<td>250–400$^c$</td>
<td>–</td>
<td>71</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ From Tables 4 and 5 in Burdige (2007) based on a large number of independent studies. The number in parentheses considers low POC burial rates in sandy sediments. $^b$ From Table 6.5.1 in that study. $^c$ Range based on previous reports (Walsh, 1981; Quiñones et al., 2010) and references therein.

Cumulative POC burial rates across the margin in Fig. 5b were calculated by integrating the measured POC accumulation at each station over the distance between stations in Fig. 5a. POC burial increased sharply on the middle shelf to ca. 100 m at both latitudes, at which point the rates diverged to give higher burial at 12° S down to the lower edge of the OMZ. Burial at 11° S amounted to 254 kmol C m$^{-1}$ yr$^{-1}$ (per metre of coastline). A lower value of 181 kmol C m$^{-1}$ yr$^{-1}$ was calculated for 12° S, possibly due to the steeper slope down to ca. 1000 m. Mean POC burial on the margin, calculated by dividing the total cumulative burial by the transect length, was 6.8 (11° S) and 6.1 mmol C m$^{-2}$ d$^{-1}$ (12° S). This compares to a range of 1.2 to 2.9 mmol C m$^{-2}$ d$^{-1}$ for the average continental margin (Table 3).

4.5 DIC fluxes

Measured DIC fluxes were high on the middle shelf (65.9 mmol m$^{-2}$ d$^{-1}$) and decreased quasi-exponentially with depth (Table 2). DIC fluxes were low in the OMZ at 12° S (2.2–4.7 mmol m$^{-2}$ d$^{-1}$) and similar to the deep sites (1.2–2.8 mmol m$^{-2}$ d$^{-1}$). Measured DIC in the benthic chambers was assumed to originate entirely from POC mineralisation. There was no clear increase or decrease in Ca$^{2+}$ and Mg$^{2+}$ concentration in the benthic chambers that would indicate an important role for carbonate precipitation/dissolution (data not shown). This is also inferred from porewater gradients of Ca$^{2+}$ and Mg$^{2+}$ (Fig. S3). Ca$^{2+}$ and Mg$^{2+}$ fluxes show that the potential contribution of carbonate precipitation was < 5% of the DIC flux across all stations. This is well within the error of the DIC flux (Table 2), such that carbonate precipitation can be ignored for all practical purposes.

Modelled DIC concentrations inside the benthic chambers at 12° S showed good agreement with those calculated from measured TA and pCO$_2$ concentrations (Fig. S2). Measured and modelled fluxes agreed to within ±50%, but most stations were simulated to within ±20% or better (Fig. 6a). It should be remembered that the model is not only constrained by the DIC fluxes but also by porewater distributions and benthic DIN and O$_2$ fluxes (Bohlen et al., 2011). Thus, whilst the modelled DIC fluxes could be improved, they form only one aspect of the overall goodness-of-fit to the observed database. In general, the agreement between the modelled and measured DIC fluxes affords confidence in the modelled DIC fluxes at 11° S where in situ pCO$_2$ measurements were not made (Table 2). The simulated DIC fluxes at 11° S showed the same trends as 12° S, although the flux on the middle shelf (8.2 mmol m$^{-2}$ d$^{-1}$) was a factor of 8 smaller.

4.6 Organic carbon burial efficiency (CBE)

At each station, CBE (%) was calculated as POC accumulation rate ÷ (POC accumulation rate + DIC flux) × 100% (Table 2). Measured and modelled CBE at 12° S showed very good agreement (Fig. 6b). Low CBEs of 19 ± 6% to 28 ± 12% were derived for St. 1 to 4 on the shelf. This contrasts with the elevated POC accumulation rates, but is in agreement with the high DIC fluxes (Fig. 6a). Low CBE of 19 ± 6% were also observed at the 409 m site where winnowing is suspected to occur. Relatively high CBEs were calculated at St. 5 to 7 in mid-water depth range of ca. 200 to
300 m (55 ± 23 % to 74 ± 37 %) and at the deep oxygenated sites (46 ± 48 % to 64 ± 19 %). At 11° S, model-derived CBE showed broadly similar trends, although the CBE at the shelf station was higher (47 %). The highest CBE of 81 % was calculated for the 695 m station (11° S) in oxygen-deficient waters (O₂ < 20 µM) below the OMZ. A detailed discussion of the uncertainties in the CBE estimates is provided in Appendix A.

4.7 Primary production and organic carbon rain rate

Primary production (PP) estimates from the ROMS-BioEBUS model for 11 and 12° S are shown in Fig. 7. The data represent the annual mean ± s.d. for the locations close to where BIOE landers were deployed. Diatoms dominated the PP at both latitudes, with rates decreasing offshore from ca. 110 mmol m⁻² d⁻¹ at the shallowest site to ca. 80 mmol m⁻² d⁻¹ at the deepest site. The model revealed a much larger intra-annual variability ranging from ca. 70 to 170 mmol m⁻² d⁻¹ with highest values in austral summer (see Fig. S4).

Organic carbon rain rates to the seafloor (RRPOC) were calculated as the sum of the benthic carbon oxidation rate (i.e. DIC flux) and POC accumulation (Table 2). For the 11 and 12° S transects, the modelled and measured DIC fluxes were used, respectively. RRPOC showed a rapid decrease on the shelf stations at 12° S with a more attenuated decrease with depth (Fig. 7a). Station 8 (409 m) at 12° S is again an exception due to the low POC accumulation there. At 11° S the trends were not so obvious due to fewer sampling stations (Fig. 7b). The fraction of PP reaching the sediment was highest at the shallowest station at 12° S (65 %), decreasing to < 12 % at the OMZ and deep stations (Fig. 7c).

5 Discussion

5.1 Spatial patterns of organic carbon preservation

Fine-grained continental margin sediments in the modern ocean (< 1000 m) account for 70–85 % of global POC burial (Hedges and Keil, 1995; Burdige, 2007). The mean global POC burial flux on the margin (1.2 to 2.9 mmol C m⁻² d⁻¹) is equivalent to around 3 % of primary production (Table 3). Although the POC burial flux on the Peruvian margin is far higher, it accounts for a similar fraction of primary production, thus implying a lack of preferential carbon burial compared to the average continental margin. However, the CBE reveals interesting spatial trends in carbon preservation that are not discernible by comparison of mean burial fluxes.

Previously published CBEs for a range of marine environments show a positive dependency on sedimentation accumulation rate (Fig. 8). This arises because sediment accumulation is intrinsically linked to carbon burial flux, which itself is strongly tied to rain rate (Müller and Suess, 1979). In addition, sediments underlying oxygen-deficient waters (O₂ < 20 µM) appear to have a higher CBE than those deposited under oxygenated bottom waters. Data from Peru seaward of the shelf agree with this idea (red stars, Fig. 8). However, the CBE above and below the OMZ is inconsistent with the existing database. Firstly, CBEs for the anoxic shelf (blue stars) plot within the range for normal oxic conditions instead of > 50 % expected for the oxygen-depleted condi-
Figure 7. (a) Mean annual primary production (±s.d.) for the 12° S transect calculated by the ROMS-BioEBUS model (diamonds). Rain rates to the seafloor (squares) and POC accumulation rates at 10 cm (stars) were estimated using the benthic measurements (Table 2). The solid line is a power law regression through the PP data. (b) As (a) for 11° S, except that rain rate is the modelled data. (c) Fraction of PP that reaches the seafloor calculated by dividing the rain rate by the PP calculated from the regression curve. The solid line is the predicted fraction at 12° S according to the empirical function of Betzer et al. (1984). The fraction for 11° S differs by a few percent only (omitted for clarity). The grey shade highlights the OMZ stations (ca. 200 to 450 m).

Figure 8. Carbon burial efficiency versus bulk sediment accumulation rate in contemporary ocean sediments. Open and filled circles represent sites from studies with bottom water O2 > 20 and < 20 µM, respectively; the former enclosed by the solid line (Canfield, 1993, 1994; Burdige, 2007; Hartnett and Devol, 2003; Reimers et al., 1992). Euxinic settings are also indicated. Stars are data from this study for 11° S (modelled data) and 12° S (measured data). Blue stars indicate sites on the shelf, which mostly have lower-than-expected CBE compared to the existing database. The two green stars correspond to the deep oxygenated sites (O2 > 20 µM) with higher-than-expected CBE. Red show the expected elevated CBEs for low oxygen environments.

5.1.1 Low CBE on the shelf

Our low calculated shelf CBEs imply that POC is preferentially degraded there, relative to deeper sites. Trends in sediment grain size and POC content support this basic idea. The fine-grained sediment fraction (clay plus silt) at 11° S is > 80% on the shelf and decreases to ca. 60% below the OMZ due to increasing fractions of coarser particles (Mosch et al., 2012). Given that POC content correlates inversely with grain size (Mayer, 1994; Bergamaschi et al., 1997),
higher contents would be predicted on the shelf relative to the OMZ. Yet, POC is < 5 % in shallower waters and > 15 % in the OMZ (Fig. 4). We can therefore assume that the ratio of organic carbon to particle surface area (OC / SA) is likely to be lower on the shelf compared to the OMZ. Low OC / SA ratios indicate organic matter in a more advanced state of diagenetic alteration (Keil et al., 1994; Mayer, 1994; Hedges et al., 1999). We can be confident that the OC / SA ratio is largely unaffected by differences in the structural composition and reactivity of POC, since organic matter on the margin is almost entirely marine-derived (Arthur et al., 1998; Reimers and Suess, 1983b). The evidence suggests that benthic mineralisation is more complete on the shelf compared to the OMZ.

Results from biochemical studies dovetail with this idea. Spatial differences in the quality of organic matter on the shelf and slope have been described (Levin et al., 2002; Niggemann and Schubert, 2006; Lomstein et al., 2009). These workers found the highest amino acid content of sedimentary organic matter (i.e. “freshener” material) on the outer shelf and within the OMZ (ca. 130 – 360 mol %). Relatively low values were reported for the middle shelf, indicating more degraded material. Sediments from the Arabian Sea OMZ are also enriched in high-quality hydrolysable amino acids (Koho et al., 2013). Similarly, more extensive diagenetic alteration of organic matter on the Peru shelf relative to the OMZ has been inferred from the amino acid degradation index (Lomstein et al., 2009). Enhanced carbon preservation in the OMZ has been noted previously, and detected as the formation of insoluble humin termed “proto-kerogen” (Reimers and Suess, 1983a, b). Considering these results collectively, the low POC content and CBE on the shelf seem to be rooted in the quality of buried material.

Prolonged exposure of sediments to O2 strongly modulates amino acid and aldose distributions in sediments, thereby decreasing the quality and quantity of buried organic matter (Hedges et al., 1999). Furthermore, OC / SA ratios and CBE are inversely correlated with the oxygen exposure time (OET) of organic matter (Hartnett et al., 1998; Hedges et al., 1999). Whilst the significance of bottom water O2 on carbon preservation remains controversial (Demaison and Moore, 1980; Pedersen and Calvert, 1990; Canfield, 1994; Hedges and Keil, 1995; Hulthe et al., 1998; Burdige, 2007), the presence of O2 could explain the low OC / SA ratios and CBEs on the Peruvian shelf in a similar way as described for the Arabian Sea OMZ (Keil and Cowie, 1999). Although shelf bottom waters were anoxic at the time of sampling, periodic intrusions of oxygenated water occasionally lead to a deepening of the upper edge of the OMZ to 200 m or more (e.g. Levin et al., 2002). The frequency and duration of these episodes can be several months, driven by seasonal and sub-seasonal anomalies in oceanographic conditions such as positive ENSO periods and the passage of coastal trapped waves (Gutiérrez et al., 2008). During these events, bottom water O2 concentrations can vary by several tens of µM within days or weeks (Gutiérrez et al., 2008), leading to an increase in the OET of organic matter.

Oxygenation is followed by an increase in macrozoobenthos species richness, density, bioturbation potential, and the construction of sediment burrows and galleries that could further increase the OET of organic matter (Tarazona et al., 1988; Gutiérrez et al., 2008). This has been observed offshore Chile, where prolonged ventilation leads to a switch in the polychaete assemblage from tube-dwelling, interface feeders to burrowing, deposit feeders (Gutiérrez et al., 2008; Sellanes et al., 2007). This sequence of events likely explains the non-zero bioturbation coefficients on the shelf (Table S2). It also fits with the previous biochemical data, since sediments with episodic exposure to O2 tend to display biogeochemical characteristics of permanently oxic settings (Aller, 1994). Redox oscillations caused by the arrival of burrowing animals may further enhance mineralisation of refractive organic compounds via priming or co-oxidation pathways, leading to a further reduction in CBE (Aller, 1998; Canfield, 1994; Hulthe et al., 1998). On the basis of the information currently available, therefore, we argue that occasional bathing of shelf sediments with oxygenated bottom waters is a determining, if not dominating, factor driving the low CBE observed there.

5.1.2 High CBE below the OMZ

Oceanographic conditions below the OMZ are relatively pristine and more conducive to steady-state diagenesis (Scholz et al., 2011). The OET of organic matter in sediments at the deepest sites was calculated by dividing the O2 penetration depths by the 210Pb-δx-derived sediment accumulation rates. Penetration depths calculated using the relation of Cai and Sayles (1996) were 4 mm at 11°S (St. 6) and 7 mm at 12°S (St. 10). The corresponding OETs are 8 and 12 years (respectively). An empirical relationship between CBE and OET derived for a range of marine settings predicts a CBE of around 20 % for these OETs, although there is considerable uncertainty associated with this trend (Hartnett et al., 1998). This is similar to the normal oxic settings in Fig. 8 and much lower than the determined value of > 60 %. Hence, factors other than the OET of carbon in the sediment must influence the CBE at the deep sites.

Our CBE calculations at the deep sites may be influenced by lateral particle transport and sediment mobilisation by current-induced resuspension. The increase in sediment mass and aluminium accumulation below the OMZ suggests that terrestrial lithogenic material is preferentially deposited there. The shelf area between 6 and 10°S is a major area of sediment reworking and winnowing by the poleward-flowing Peru Undercurrent with average near-bottom velocities of 8–9 cm s−1 (Suess et al., 1987; Chaigneau et al., 2013). Fine-grained biogenic debris from this area may undergo multiple deposition/resuspension cycles as it is redistributed down the slope in the benthic boundary layer in a southerly direction.
5.2 Carbon mineralisation in the water column

A comparison of the rain rate and PP estimates from the ROMS-BioEBUS model shows that only a minor fraction of PP reaches the seafloor beyond the middle shelf (Fig. 7c). The true fraction may be even lower since modelled PP is 2 to 3 times below the range of 250 to 400 mmol m$^{-2}$ d$^{-1}$ reported previously (Walsh, 1981; Quiñones et al., 2010 and references therein). Robust PP estimates close to the coastline are impeded by the relatively coarse spatial resolution (1/9°) in the model which cannot accurately resolve nearshore processes. The model also represents climatological conditions (i.e. interannual steady state), whereas PP during neutral or cold (La Niña) ENSO phases (this study) may deviate from the mean value (Ryan et al., 2006). Nonetheless, based on the data in Fig. 7c, it appears that the Peruvian margin broadly behaves as an open ocean setting when compared to existing empirical relationships relating PP to rain rate (e.g. Betzer et al., 1984).

The rate at which organic matter is respired during transit through the water column is usually calculated using sediment trap data (Martin et al., 1987). The unrespired fraction of export production has been widely described using the following function:

$$F(z)/EP = (z/100)^{-b},$$

where EP is the export production, typically defined at 100 m, $F(z)$ is the sinking flux at depth, $z$ (m), below 100 m, and $b$ is the dimensionless attenuation coefficient (Martin et al., 1987). Low $b$ indicates slow degradation in the water column and a high fraction of export production reaching the sediment and vice-versa. A mean $b$ of 0.86 for oxic open-ocean waters was derived by Martin et al. (1987), which is similar to the 0.82 ± 0.16 quoted by Berelson (2001) for a range of marine settings. Primeau (2006) proposed a lower $b$ of 0.70 ± 0.08 based on a reanalysis of Berelson’s data. A recent study in the North Atlantic using free-drifting sediment traps also reported a $b$ of 0.70 (Giering et al., 2014).
We estimated $b$ by fitting the calculated rain rates at 11 and 12° S to the following function analogous to Eq. (7):

$$RRPOC(z)/EP = (z/101)^{-b}.$$  

(8)

The EP was determined at each station individually by multiplying the PP estimate from the pelagic model by the ratio of particle export to primary production ($pe$ ratio, Dunne et al., 2005). The $pe$ ratio was assumed to be constant across the margin and was calculated as PP / RRPOC using data from St. 2 (12° S). This station was taken as the reference point since the water depth (101 m) is approximately 100 m. The derived $pe$ ratio was 0.26; consistent with observations that most PP in the Humboldt system is mineralised in the surface mixed layer under non-El-Niño conditions (Quiñones et al. 2010 and references therein). The two shallowest stations with water depth < 101 m were excluded from the analysis in addition to St. 8 since, as mentioned, we suspect that sediment resuspension by the bottom currents is important there. To offset the potential bias in our RRPOC estimates due to seasonal variability in primary production, the PP was averaged over the period corresponding to the cruise and 1 month previously. (See Appendix.)

The best fit to Eq. (8) shown in Fig. 9 was obtained with $b = 0.54 ± 0.14$ (NonlinearModelFit function in MATHEMATICA). The $b$ value considering data from 12° S only is 0.85 ± 0.25 (the result for 11° S is not statistically significant at the 95% level). Our derived $b$ coefficients thus tend toward the higher open-ocean composite estimates of Martin (1987) and Primeau (2006). Yet, previous studies indicate that respiration of organic matter is significantly reduced in oxygen-deficient water columns (Martin et al., 1987; Devol and Hartnett, 2001; Van Mooy et al., 2002). For example, a $b$ of 0.32 was determined for the Peruvian OMZ at 15° S using sediment trap data (Martin et al., 1987). Devol and Hartnett (2001) calculated a value of 0.36 for the Mexican margin using benthic carbon oxidation rates and burial fluxes. This agrees with 0.40 derived using moored sediment trap data from the same location (Van Mooy et al., 2002). These workers supported their fieldwork with laboratory experiments showing that natural particulate material collected at the base of the euphotic zone was degraded less efficiently under anoxic versus oxic conditions. A notable exception to these studies is the $b$ value of 0.79 derived for the Arabian Sea OMZ using sediment traps (Berelson, 2001). Nonetheless, based on these findings, a much lower $b$ coefficient for our composite data set may have been expected.

We propose that the high $b$ determined in this study is attributed to the multiple resuspension/deposition of slope sediments by internal waves discussed above. Resuspension enhances the particle residence time in the water column and shifts the site of labile POC mineralisation away from the sediments. This will reduce the benthic DIC flux and thus lower the calculated RRPOC. This explanation is consistent with unusually high CBE below the OMZ. The low rain rate calculated for St. 8 at 12° S further exemplifies the importance of sediment reworking on the calculated rain rates (Fig. 8). By implication, particle transport on the Mexican margin and at the deep offshore station at Peru (15° S) is more controlled by vertical settling, resulting in shorter residence times and enhanced carbon preservation through the water column. The good agreement between the $b$ coefficients determined independently by traps and benthic data from Mexico supports this idea (Devol and Hartnett, 2001; Van Mooy et al., 2002). In contrast, the latitudinal differences in $b$ at Peru (this study; Martin et al., 1987) could simply reflect bias from comparing benthic- and pelagic-derived rain rates from a complex margin system displaying high temporal variability in productivity (Walsh et al., 1981).

6 Conclusions

Fieldwork undertaken on the fascinating Peruvian margin has improved our understanding of POC cycling in anoxic margin sediments and expanded the existing CBE database. Our key findings are as follows: (1) low CBE in shelf sediments is driven by the episodic intrusion of oxygenated waters. This is consistent with the existing CBE database if the shelf can be geochemically classified as a “normal” setting despite fluctuating bottom water O$_2$ levels. Previous biochemical analyses suggest that this is the case. (2) High CBE in oxygenated waters (O$_2$ > 20 µM) below the OMZ is caused by the deposition of reworked, degraded material originating from sites higher up on the slope. They are, therefore, not representative of oxygenated sediments. (3) Low oxygen concentrations in the OMZ below 200 m are associated with the preservation of POC in sediments, although the redistribution and accumulation of reworked sediments at these depths needs to be more clearly examined. (4) POC rain rates do not appear to be overly enhanced by low oxygen concentrations compared to other oxygen-deficient environments.

Our conclusions regarding this last point require further validation with particle flux measurements from sediment traps. Trap data may reveal lower rates of POC mineralisation in the water column, as observed in other low-oxygen settings. If this turns out to be the case, it would imply that POC fluxes are either subject to high temporal variability, or that a fraction of labile material resuspended from the sea bed is degraded in the benthic boundary layer and exported down the slope. The increase in sedimentation rates and high CBE below the OMZ supports the latter alternative. These investigations would benefit from an improved understanding of particle disaggregation by zooplankton and subsequent stimulation of the microbial loop (Giering et al., 2014).

A major outstanding barrier for deriving annual carbon and nutrient budgets on the Peruvian margin is the uncertainty associated with non-steady-state conditions. Inter- and intra-annual oceanographic anomalies dictate the depth and intensity of oxygen-deficiency, nutrient levels and *Thioploca*...
spp. biomass (Gutiérrez et al., 2008). Consequently, the contribution of the sediments to primary production on short (e.g. days, weeks) and long (e.g. months, years) timescales is completely unknown. This is a topic that could be tackled by incorporating benthic processes into the pelagic model ROMS-BioEBUS. Model predictions would be strengthened with benthic flux data collected during the low productivity season (austral winter).
Appendix A: Uncertainties in CBE calculations

The burial efficiency of carbon is the basic quantitative metric that allows comparisons of carbon preservation at different locations. Quantification of the CBE requires at least two of the following three pieces of information: (i) the rain rate of organic carbon to the sediment; (ii) the rate of carbon burial at the sediment depth where POC content remains constant; and (iii) the depth-integrated rate of organic carbon mineralisation (Buridge, 2007). That is, at steady-state, \( \omega_c = \omega_1 + \omega_2 \). They are typically quantified using (in the same order) (i) sediment trap particle fluxes; (ii) sedimentation rates (using radioactive isotopes) combined with carbon content measurements; and (iii) stoichiometric mass balances, diagenetic models or benthic fluxes. Each of these approaches integrates the carbon flux over very different timescales; from days to weeks for traps to hundreds or thousands of years for burial (Buridge, 2007). This temporal decoupling creates uncertainty that may affect CBE estimates.

Our calculated CBE rely on the use of benthic data only (i.e. (ii) and (iii) above) as opposed to sediment trap fluxes. It is well known that sediment trap data have multiple sources of uncertainty (Buesseler et al., 2007). Traps only capture the settling flux on timescales of deployment, typically days to weeks, which may be insufficient to capture temporal variability (Haake et al., 1992). Traps may also underestimate the particle flux in water depths < 1500 m due to high lateral current velocities (Yu et al., 2001), potentially leading to an overestimate of fluxes further offshore. This effect is exacerbated over the shelf and upper slope where particles are transported in the benthic boundary layer (Jahne et al., 1990). These dynamics may be particularly pertinent for the Peru margin where sediments are frequently resuspended by bottom currents and transported down the slope (Reimers and Suess, 1983c). Nonetheless, sedimentation rate and carbon oxidation rate calculations are not free from their own uncertainties, as discussed in the subsequent paragraphs.

Benthic chambers are arguably the best tools to estimate the exchange of solutes between the sediments and the water column (Tengberg et al., 2005). Yet, as for traps, DIC flux may display short-term (sub-annual) variability via changes in productivity and thus not truly reflect the long-term mean degradation rate. Mean PP estimates using the pelagic model do reveal substantial intra-annual variability on the Peru margin (Fig. S4). Both campaigns at 11 and 12°S took place toward the end of austral summer, that is, the high productivity season, which suggests a bias in our DIC fluxes towards higher values (Echevin et al., 2008). However, because benthic carbon mineralisation rates do not respond quantitatively and synchronously to changes in rain rate, it is not possible to accurately ascertain if the fieldwork timing adversely affects our assumption that the measured DIC flux accurately represents the mean flux. The timescale of the bulk organic carbon mineralisation is determined by the reactivity of the individual biomolecules, which vary over many orders of magnitude (Middelburg, 1989). Experiments with fresh phytoplankton have shown that approximately 50 % of organic carbon degrades rapidly within 1 month of deposition on the seafloor, with the other 50 % requiring many months or years to degrade (Westrich and Berner, 1984). This implies that variability in the measured benthic DIC fluxes will largely reflect the previous month of deposition of the labile POC fraction. In that regard, the benthic DIC flux provides a more attenuated estimate of changes in rain rate than sediment trap data. Based on the modelled primary production rates during the sampling months, which were around 25 % higher than the annual mean (Fig. S4), we tentatively estimate that the measured DIC fluxes at 12°S are roughly 25 % higher than the long-term average. The analytical error on the DIC flux calculation from TA and \( pCO_2 \) measurements is negligible by comparison. Correcting the DIC flux for seasonal variability would increase our CBE estimates by around 4 to 5 % on the shelf and 5 to 7 % in deeper waters. Strictly, though, the mean DIC flux for the last ca. 100 years would be required to be consistent with the burial data.

Artifacts in flux estimates due to enclosure of bottom water by benthic chambers have been discussed extensively in the literature (e.g. Hammond et al., 1996; Tengberg et al., 2005). Uncertainty in chamber volume, leakage and improper alteration of the hydrodynamic regime inside the chambers all influence our confidence in any particular concentration measurement or flux estimate. We can be reasonably confident that leakage through the bottom of the chambers in contact with the sediments did not occur at the sampling stations. Silicate (assumed conservative tracer) and \( O_2 \) (optode) concentration gradients inside the chambers were linear and showed no evidence of infiltration of chamber water by the outside bottom water (Sommer et al., unpublished data). Furthermore, the surface sediments were cohesive and porous along the whole margin (Table 1), allowing penetration of the benthic chambers to 8 to 10 cm below the sediment surface and rendering leakage unlikely. Yet, we cannot rule out the possibility that the measured DIC fluxes deviated from the true fluxes due to changes in water flow across the sediment–water interface. This artifact is mostly prominent in sandy sediments where current-driven advection through the surface layers contributes significantly to solute exchange between sediments and seawater (Jahne et al., 2000). The cohesive and fine-grained sediments (sandy mud to slightly sandy mud) at Peru typically have low permeability where molecular diffusion dictates the benthic flux. Thus, error in the DIC flux due to chamber artifacts as well as seasonal variability is likely to be much smaller than the difference in DIC flux measured in the two chambers during each lander deployment, that is, due to seafloor heterogeneity (Table 2).

POC accumulation rates have a relative error determined by the uncertainty in \( \omega_{a/c} \) and POC content. The uncertainty in POC content was assumed to be ±20 %, mainly due to data scatter. Whilst this is realistic for most of the sites investigated here, productivity and sediment POC content have
increased on the Peruvian margin in the last 200 years, coinciding with the expansion of the OMZ driven by a northward displacement of the Intertropical Convergence Zone (Gutiérrez et al., 2009). This is clearly seen in the upper 10 cm of the outer shelf and OMZ sediments (Fig. 4) and undermines our ability to accurately define the depth where the POC content shows little change with sediment depth. The regime shift is not recorded at the middle shelf sites because the high sedimentation rates restrict the observable archive to the last 100 years. Similarly, the deep stations below the OMZ are presumably beyond the sphere of influence of short-term climatic variations. These sites show the expected decrease in POC for sediments undergoing steady-state mineralisation with very little mixing by bioturbation. For the stations affected by the regime shift (St. 4 to 7), we calculated the CBE based on the average POC content in this recent layer (upper ∼ 10 cm) where our 210Pbxs measurements were made. A 50% decrease in POC content at these stations would lower the CBE by 20 to 40%. Note that higher POC mineralisation rates associated with the increase in POC content over recent centennial time scales should be accurately reflected in the DIC flux. Diffusive path lengths calculations (Lasaga and Holland, 1976) show that DIC concentrations in the upper 10 cm will adjust to higher POC fluxes within a few years only.

Sediment accumulation rates (ωacc) were constrained using 210Pbxs data (half-life of 22 yr) using a model that utilises a widely used empirical description of bioturbation intensity with sediment depth (Christensen, 1982). We estimated that ωacc are accurate to within ±20% for the derived bioturbation coefficients (Table S2). Continuously supplied tracers such as the 210Pbxs radioisotope have a characteristic timescale equal to about 5 half-lives, that is, ca. 110 yr. Therefore, our accumulation rates correspond to recent sediments deposited since the Little Ice Age and are of the same order as those published previously for the same area (Reimers and Suess, 1983c; Kim and Burnett, 1988; Levin et al., 2002; Gutiérrez et al., 2009). They are not applicable to the older underlying sediment, where mass accumulation rates are lower and have undergone hiatuses in the past (Reimers and Suess, 1983b; Gutiérrez et al., 2009; Schönfeld et al., 2014). However, sedimentation and bioturbation rates inferred from radioisotopes with vastly different half-lives often differ by an order-of-magnitude or more (e.g. Smith et al., 1993). Bioturbation coefficients of ca. 100 cm² yr⁻¹ were derived from 234Th (half-life 24 d) distributions below the OMZ on the Peru margin (Levin et al., 2002), which is four orders-of-magnitude higher than we determined using 210Pbxs. Given the low sedimentation and bioturbation rates at the deeper stations, the 210Pbxs distributions may thus indicate higher rates of sediment mixing and lower sedimentation rates than derived with the model.

We used the anthropogenic enrichment peak of the nuclide 241Am as an independent time marker to validate the lead chronology (Fig. 2). 241Am originates from nuclear tests in the Southern Hemisphere beginning in the early 1950s. At four stations, ωacc was estimated by dividing the depth of the 241Am peak by 60 years without correcting for compaction, resulting in rates of 0.15 cm yr⁻¹ at St. 5, and 0.05 cm yr⁻¹ at St. 4, 7, and 9. No 241Am was detectable at the other stations. These rates agree with the 210Pbxs values to within 10 to 50% (Table 2). Although ωacc estimated this way are only approximate, they confirm the order-of-magnitude values determined by the 210Pbxs model. Furthermore, it is obvious that the 234Th-derived bioturbation coefficient of 100 cm² yr⁻¹ does not capture the activity of the 210Pbxs data (red curve, Fig. 2). The high 234Th-derived coefficient is very likely a model artifact caused by the use of short-lived radioisotopes to infer mixing rates in weakly mixed sediments (Lecroart et al., 2010).

Clearly, there are multiple potential sources of uncertainty in CBE estimates based on DIC fluxes and POC burial fluxes. Yet, whilst concerns remain regarding both short- and long-term variability in fluxes, we believe our derived CBEs are robust on decadal timescales. The mean relative error in CBE across all stations is around 40% (at 12° S); a fair representation of the combined uncertainties.
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