Benthic phosphorus cycling in the Peruvian oxygen minimum zone

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Abstract. Oxygen minimum zones (OMZs) that impinge on continental margins favor the release of phosphorus (P) from the sediments to the water column, enhancing primary productivity and the maintenance or expansion of low-oxygen waters. A comprehensive field program in the Peruvian OMZ was undertaken to identify the sources of benthic P at six stations, including the analysis of particles from the water column, surface sediments, and pore fluids, as well as in situ benthic flux measurements. A major fraction of solid-phase P was bound as particulate inorganic P (PIP) both in the water column and in sediments. Sedimentary PIP increased with depth in the sediment at the expense of particulate organic P (POP). The ratio of particulate organic carbon (POC) to POP exceeded the Redfield ratio both in the water column (202 ± 29) and in surface sediments (303 ± 77). However, the POC to total particulate P (TPP = POP + PIP) ratio was close to Redfield in the water column (103 ± 9) and in sediment samples (102 ± 15). This suggests that the relative burial efficiencies of POC and TPP are similar under low-oxygen conditions and that the sediments underlying the anoxic waters on the Peru margin are not depleted in P compared to Redfield. Benthic fluxes of dissolved P were extremely high (up to 1.04 ± 0.31 mmol m⁻² d⁻¹), however, showing that a lack of oxygen promotes the intensified release of dissolved P from sediments, whilst preserving the POC / TPP burial ratio. Benthic dissolved P fluxes were always higher than the TPP rain rate to the seabed, which is proposed to be caused by transient P release by bacterial mats that had stored P during previous periods when bottom waters were less reducing. At one station located at the lower rim of the OMZ, dissolved P was taken up by the sediments, indicating ongoing phosphorite formation. This is further supported by decreasing porewater phosphate concentrations with sediment depth, whereas solid-phase P concentrations were comparatively high.

1 Introduction

Phosphorus is an essential nutrient; it serves as an energy carrier for all living species and is a limiting macronutrient for marine primary production on geological timescales (Ingall and Jahnke, 1994; Föllmi, 1996; McManus et al., 1997; Filippelli, 2002; Paytan and McLaughlin, 2007; Tsandev et al., 2012; Ruttenberg, 2014). Due to its impact on marine primary production, the oceanic phosphorus inventory modulates the atmospheric CO₂ level and Earth’s climate (Ganeshram et al., 2002; Wallmann, 2003; Ingall, 2010). Hence, it is crucial to understand feedback mechanisms of the P cycle to make future predictions.

Particulate and dissolved phosphorus in the ocean originates from terrestrial chemical weathering of the P containing mineral group of apatite (Filippelli, 2002). Only around 30 % of the P discharged to the oceans is potentially bioavailable (Compton et al., 2000) as dissolved P, inorganic P adsorbed to minerals, or associated with metal oxides and P bound within particulate organic matter. However, the largest fraction of the fluvial P is trapped in estuaries or buried in continental margin sediments and thereby removed from the P cycle before it reaches the open ocean (Compton et al., 2000). The delivery of P to the sediments in the open ocean is mainly composed of organic and inorganic P associated with the export of organic detritus and other particles from the photic zone. P adsorbed to minerals such as Mn and Fe (oxyhydr)oxides (Föllmi, 1996; Delaney, 1998; Faul et al., 2005) are further sources, as well as P input from fish debris.

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P cycling is strongly affected by redox-dependent processes. P can be scavenged by Fe (oxyhydr)oxides in oxic sediment and released across the sediment–water interface due to the reduction dissolution of Fe (oxyhydr)oxides in anoxic sediments (Sundby et al., 1986; Slomp et al., 1998). Furthermore, recent studies showed that sulfur bacteria found in surface sediments of anoxic environments can internally store and release P under oscillating redox conditions (Ingall and Jahnke, 1997, and references therein). Therefore, these organisms are a key player for the modulation of porewater P concentrations and benthic P release to the water column. Additionally, hypoxic or anoxic conditions favor the precipitation of P in the form of authigenic carbonate fluorapatite (CFA) (Froelich et al., 1988; Suess and von Huene, 1988; Goldhammer et al., 2010; Ingall, 2010; Schenau and De Lange, 2000). For non-upwelling areas, the required phosphate oversaturation in the porewaters has been attributed to the reductive dissolution of P bearing Fe (oxyhydr)oxides (e.g., Ruttenberg and Berner, 1993, Slomp et al., 1996). In contrast, the CFA formation in sediments of the Namibian upwelling area was linked to microbial P release into the porewaters (Schulz and Schulz, 2005). The resulting feedback on oceanic primary production and atmospheric O$_2$ and CO$_2$ levels triggered by changes in benthic P sequestration is still unclear. Presently, three opposing views have been raised: (1) intensified phosphate release from the sediments to the water column caused by an expansion of low-oxygen waters (Ingall and Jahnke, 1994; Stramma et al., 2008) could stimulate the primary production in the surface waters (Wallmann, 2003). This, in turn, may lead to a more intensified oxygen demand and a positive feedback with benthic P release (Slomp and Van Cappellen, 2007; Wallmann, 2010; Moffitt et al., 2015). (2) A negative feedback on P release has been postulated based on observations of CFA mineral precipitation found in the present-day oxygen-depleted upwelling areas (Schulz and Schulz, 2005; Arning et al., 2009a, b; Goldhammer et al., 2010; Cosmidis et al., 2013). Being a major sink for bioavailable P (Delaney, 1998; Ingall, 2010), it has been argued that the expansion of OMZs may increase the CFA precipitation in the sediments and thus mitigate the benthic phosphate release (Ganeshram et al., 2002; Goldhammer et al., 2010; Ingall, 2010). (3) A third scenario suggests that the formation of CFA is in balance with enhanced P release from anoxic sediments, implying that the dissolved oceanic P inventory is largely unaffected by oxygen concentrations (Delaney, 1998; Anderson et al., 2001; Roth et al., 2014). These conflicting scenarios show that there is further need to explore the benthic–pelagic P cycling in oxygen-deficient environments in order to enable improved predictions.

In this study, we explore P cycling in the Peruvian OMZ to identify and quantify P sources to the sediment and the return of inorganic dissolved P back to the water column. Our data set comprises samples of particulate matter from the water column as well as porewater, sediment samples and samples of filamentous sulfur bacteria. We present in situ benthic phosphate fluxes, particulate matter C / P ratios for water-column particles and surface sediments and P burial fluxes, and relative abundances of sulfur bacteria for six stations along the depth transect across the Peruvian shelf at 12°S. From a mass balance for P cycling in the sediments, we conclude that the benthic P sources and sinks were, in general, imbalanced during our sampling campaign.

2 Study area

The study area is located in the center of the Peruvian OMZ at 12° S covering the shallow shelf from ~70 m water depth to mid-slope depths of about ~400 m (Fig. 1). During our sampling campaign in January 2013, neutral or slightly negative El Niño–Southern Oscillation (ENSO) conditions dominated (http://www.cpc.ncep.noaa.gov), and the bottom water oxygen concentrations were below the detection limit of the Winkler titration (5 µmol L$^{-1}$) down to ~450 m water depth (Fig. 1, Table 1). Below the OMZ, oxygen concentrations increased to 19 and 53 µM at 770 and 1025 m water depth, respectively. Nitrate concentrations were below 12 µM from 128 to 407 m water depth (Table 1). During the measuring period, the bottom water at station I (74 m) was sulfidic and depleted in nitrate (Table 1; Sommer et al., 2016).

The oxygen-deficient waters off Peru belong to one of the world’s most prominent OMZs. Southeasterly trade winds...
that are driven by the Pacific subtropical anticyclone engender offshore transport of surface waters and upwelling of subsurface waters from the poleward-propagating Peru upwelling (PUC) (Strub et al., 1998). These water masses are oxygen depleted and rich in nutrients, favoring primary production of up to 3.6 g C m\(^{-2}\) d\(^{-1}\) in surface waters (Pennington et al., 2006). As a consequence, the intense oxygen consumption induced by the degradation of sinking particulate organic matter and a sluggish ventilation causes the development of a strong OMZ. Based on the definition that the oxycline of an OMZ is at \(\sim 22 \mu M\) (Fuenzalida et al., 2009), the Peruvian OMZ extends from approximately 50 to 700 m water depth. The greatest upwelling strength is reached during austral winter and spring between 5 and 15\(^{\circ}\) S (Strub et al., 1998). The phases of strong upwelling are followed by high rates of primary production in austral summer. The coastal area off Peru displays a highly variable hydrographic regime. Especially during positive ENSO periods, coastal trapped waves emerging from equatorial Kelvin waves in the equatorial eastern Pacific occur frequently (Gutiérrez et al., 2008, and references therein; Mosch et al., 2012). Consequently, the thermocline and the oxycline shift downwards by ca. 100 m and bottom-water oxygen concentrations can increase from practically zero to around 100 \(\mu M\) in days to weeks (Gutiérrez et al., 2008; Schunck et al., 2013; Graco et al., 2016). Seasonally, bottom waters of the shelf (75 m) can transition from oxic or hypoxic between austral winter/spring (low primary production) to anoxic throughout the rest of the year (Noffke et al., 2012). Furthermore, porewater uranium profiles at 11\(^{\circ}\) S indicate variable redox conditions at the upper rim of the OMZ (Scholz et al., 2011). The shelf area above 200 m water depth is therefore characterized by non-steady-state conditions, whereas the oxygen concentrations in the core OMZ (\(\sim 200\)–400 m water depth) are predominantly below the detection limit throughout the year.

The sediments of the Peruvian OMZ have POC contents ranging from 15 to 20 wt% within the OMZ and > 5 wt% below the OMZ and on the shelf (Dale et al., 2015). The fine-grained, diatomaceous mud lens between 11 and 15\(^{\circ}\) S accumulates under low PUC bottom-water velocities in 50 to 500 m water depth (Krissel et al., 1980). This favors high sedimentation rates, carbon preservation, and burial (Suess et al., 1987; Dale et al., 2015). Further down, at mid-slope depth, a high-energy regime favoring erosive settings leads to the formation of phosphorites (Reimers and Suess, 1983; Glenn and Arthur, 1988; Arning et al., 2009b; Mosch et al., 2012). Another interesting observation between 70 and \(\sim 300\) m of water depth is the occurrence of mat-forming filamentous sulfur bacteria (Mosch et al., 2012). Bacterial mats are not conspicuous below 300 to 400 m water depth, and instead foraminiferal sands are more common.

### 3 Methods

Sampling of water-column particulate matter and sediment cores as well as the deployment of the benthic landers BIGO I and II (Biogeochemical Observatories) were conducted along the 12\(^{\circ}\) S depth transect during RV Meteor cruise.
M92 in January 2013. The geographical position and water-column properties for the main stations are reported in Table 1. The data set on in situ phosphate fluxes comprised 10 stations from 74 to 989 m water depth. The water-column particle sampling was performed at six stations from 74 to 407 m water depth. These stations are considered as main stations and for consistency the stations are numbered according to the data set published in Dale et al. (2015). Hydrographic parameters and oxygen concentrations were obtained by deploying a CTD/rosette equipped with a Sea bird oxygen sensor (detection limit is 5 µM) calibrated by Winkler titration.

3.1 Water-column particles

Particulate matter was filtered using water from Niskin bottles from the CTD/rosette and analyzed for total particulate phosphorus (TPP), particulate inorganic phosphorus (PIP), and particulate organic carbon (POC) concentrations. Following Labry et al. (2013), we expect the PIP phase to be comprised of inorganic P phases originating from abiotic particulate P as well as inorganic P from biogenic particulate P. Abiotic PIP comprises detrital P associated with minerals from terrigenous sources. Biotic PIP is composed of orthophosphates, pyrophosphates, and polyphosphates within eukaryotic and prokaryotic cells.

Between three and six water depths were sampled per station. The water was filtered into 10 L PE containers rinsed beforehand with ultrapure water (MilliQ). The containers were shaken before filtration, which was performed within 24 h after sample retrieval. Approximately 2 to 4 L of seawater were filtered through a pre-weighed and combusted (450 °C, 5 h) 0.7 µM Whatman GF/F filter using a seawater vacuum pump and Duran bottle top filters. After filtration, all filters were immediately frozen at −20 °C. At the shore-based laboratory the GF/F filters were dried overnight at 45 °C and divided into three equally sized pieces using a scalpel. The total filtered water volume was divided by 3 to calculate elemental concentrations on each filter section assuming homogenous coverage of particles on the filters.

3.1.1 Total particulate phosphorus (TPP), particulate inorganic phosphorus (PIP) and particulate organic phosphorus (POP)

The determination of TPP and PIP concentrations by combustion and colorimetric methods has been described by Aspila et al. (1976); Loh and Bauer (2000); Asahi et al. (2014), and others. However, the combustion and acid dissolution (HTC/hydrolysis) that has been applied for PIP determination is limited by the fact that polyphosphates are only partly hydrolyzed, but provides the best compromise compared to other methods (Labry et al., 2013). Although polyphosphates are underestimated, it includes intracellular P that is often dominant in plankton and only small amounts of organic P. In the case of TPP we used the Aspila (1976) method without an oxidant (e.g., MgSO₄) during combustion, which underestimates the TPP concentration in water-column particles. However, for the sediments, the results following the Aspila (1976) method were compared to total digestions showing an average recovery of the Aspila method of approximately 102 %. In two samples we found an overestimation in total digestion of more than 100 % and, in one sample, an underestimation of 32 %, possibly due to heterogeneity of the samples caused by very small phosphorite granules.

Filter segments for TPP concentration were combusted at 550 °C for 90 min and afterwards soaked with 20 mL 1 N hydrochloric acid (HCl) and shaken for 24 h at room temperature. Then, the solution was filtered and 0.35 mL triple reagent (40 mL 9.8 N sulfuric acid, 12 mL ammonium molybdate, and 4 mL potassium antimonyl tartrate solution) and 0.175 mL ascorbic acid and 3 mL 1 N HCl were added to 3.75 mL of the sample solution. Then, 0.3 mL of 12.5 N sodium hydroxide (NaOH) was added to the solution before colorimetric measurement of phosphate at 880 nm using a Hitachi U-2001 photospectrometer. This was done because test runs with the standard series revealed that the slope of the calibration curve was not steep enough to measure the low phosphate concentrations expected. To resolve this issue, the pH of the solution was slightly increased using NaOH. Measurements were accompanied using a standard series consisting of eight standards ranging from 5 to 100 µM PO₄³⁻, prepared from a Merck phosphate stock solution. We used 0.75 mL of each standard for the standard series and treated each of them in the same manner as described above. The samples were measured undiluted due to low concentrations and the technical detection limit of a 1 cm cuvette. Hence, we used 3.75 mL of the filtered sample solution, added the reagents mentioned above, and divided the concentrations by a factor of 5 to adjust the results to those of the standard series. A factor of 0.02 was used to transform the concentration unit to µmol L⁻¹. The amount of filtered water (f) refers to 1/3 of the total filtered water volume (f is different for each sample):

$$\text{TPP or PIP [µmol L}^{-1}]=\left[\frac{\text{PO}_4^{3-}}{5 \cdot f}\right] \cdot 0.02 \cdot \frac{0.02}{f}.$$  

The same procedure was performed for PIP without the combustion step. The POP concentration was calculated by the difference of the measured (as phosphate) TPP and PIP concentrations.

3.1.2 Organic carbon concentration

The filter sections for the analysis of POC concentration were fumed with 37 % HCl overnight to remove inorganic carbon, dried, and wrapped into tin caps. Samples were measured by flash combustion with a Carlo Erba elemental analyzer (NA1500). The analytical precision and detection limit were 0.04 dry wt %. The water-column POC concentrations are given in µmol L⁻¹.
3.2 Porewater and solid-phase analysis

Sediment cores were recovered using video-guided multiple corers (MUCs) equipped with PVC liners with an inner diameter of 10 cm. The porewater and solid-phase subsampling was performed immediately after recovery in an argon-filled glove bag at in situ seafloor temperature. The bottom water was siphoned with a plastic tube and filtered through cellulose acetate filters. Afterwards, the cores were sectioned into 0.5 cm intervals from 0 to 5 cm sediment depth and 1 cm intervals afterwards. The sediment samples were filled into centrifuge tubes and the porewater was separated from the sediments by centrifuging for 20 min at ∼ 3940 G. The supernatant porewater was filtered through cellulose acetate filters inside the glove bag. Samples were immediately analyzed for total dissolved phosphate (TPO$_4^-$) and dissolved ferrous iron (Fe$^{2+}$) after porewater extraction using a Hitachi U-2001 spectrophotometer. The analyses were performed according to the standard techniques described in Grasshoff et al. (1999). A sediment subsample was taken from each sediment depth and stored refrigerated in pre-weighed air-tight plastic cups to determine the water content, porosity, and total organic carbon (TOC) content. The residual sediments were stored frozen at −20 °C (sediment and filter samples) and the porewater samples were refrigerated at 4 °C for land-based analytics.

The TOC concentration (in µmol mg$^{-1}$) of freeze-dried and ground sediment samples was determined by flash combustion in a Carlo Erba Elemental Analyzer (NA 1500). For POC determination, samples were decalcified with 2.5 N HCl prior to the measurement. Solid-phase TPP and PIP concentrations were measured according to the method of Aspila et al. (1976) in a similar manner as described above. The 50 mg of freeze-dried and ground sediment were digested in 1 N HCl for a minimum of 24 h to dissolve the sedimentary PIP phase. Sediment portions analyzed for TPP were combusted at 550 °C for 90 min before adding 1 N HCl. The solutions were filtered and the reagents mentioned above were added prior to measurement. We used the sedimentary reference standards SDO-1 (Devonian Ohio Shale, USGS; Govindaraju, 1994) and MESS-3 (Marine Sediment Reference Material, Canadian Research Council) and replicate measurements of samples to ensure measurement accuracy. The standard series applied to the measurements covered a concentration range from 5 to 100 µM.

To determine the terrigenous P input to the sediments, and to calculate the TPP burial flux (for calculation, see Sect. 3.6 and Table 2), sediments were analyzed using total digestion. About 100 mg of freeze-dried and ground sediment were digested in hydrofluoric acid (40%, supra pure), nitric acid (65%, supra pure), and perchloric acid (60%, supra pure). For measurement accuracy the reference standards SDO-1 and MESS-3 as well as methodological blanks were included in the analysis. The aluminum concentration in the digestion solutions was measured using an inductively coupled plasma optical emission spectrometer (ICP-OES, Varian 720 ES). The relative standard deviation (RSD) for [Al] was found to be < 1 %.

The XRD data of core 107MUC23 from 407 m water depth were obtained from approximately 1 g of freeze-dried and ground sediment in the lab of the University of Bremen.

3.3 POC in relation to various fractions of P (POC / xP ratios)

The molar POC / xP ratios (where xP = TPP, PIP, or POP) of the water-column particles at stations I, IV, and V were calculated from measurements on two filter samples per water depth. For these samples a minimum and maximum value were calculated. For the other stations III, VI, and VIII, only one sample per water depth was available. Here, we assumed an average natural variability calculated from the duplicate measurements of stations I, IV, and V for each P species (Supplement). For sediment samples we calculated a standard deviation from repeated measurements of the sediment standards MESS-3 and SDO-I (Supplement).

3.4 Benthic lander fluxes

Benthic lander deployments were performed at 10 stations along the 12°S transect (I to X according to Dale et al., 2015). In situ benthic fluxes were obtained using the two BIGOs I and II (BIGO: Biogeochemical Observatory). They were equipped with two circular flux chambers (internal diameter 28.8 cm, area 651.4 cm$^2$) (Sommers et al., 2009). An online video-controlled launch system allowed precise placement of the BIGO at the seafloor directly located beneath the particle sampling stations in the water column and in proximity to the multi-corer stations. After a 4 h rest period at the seafloor during which surrounding bottom water was periodically flushed into the chamber, the chambers were slowly driven into the sediment. The BIGOs stayed for 28 h at the seafloor, while eight water samples per chamber were taken via glass syringes. In order to obtain bottom water background information, additional samples were taken every 8 h from the ambient bottom water. Phosphate concentrations in the syringe samples were measured on board using an auto analyzer. The standard series covered a concentration range from 0.05 to 3.5 µM. The fluxes were calculated from the slope of linear regression of all eight data points vs. the sampling time (Supplement) and corrected for the water volume in the chamber and the dead volume of the 1 m long Vygon tubes connecting the syringes with the flux chambers. The error caused by the dilution from the dead volume of these tubes was calculated from the chloride concentration measured in the syringe samples. Benthic lander TPO$_4^-$ fluxes for most sites are based on two replicate chamber measurements. The uncertainty given for the TPO$_4^-$ fluxes is the difference between the minimum and maximum fluxes from the average of the two benthic chambers. At two stations (IV and V),
3.5 Diffusive flux calculations

The diffusive fluxes of TPO\textsubscript{4} and Fe\textsuperscript{2+} from the sediment to the bottom water were calculated by applying Fick’s first law of diffusion:

\[ \frac{dF_{\text{TPO}_4/\text{Fe}^{2+}}}{dx} = -\dot{\phi} D_{\text{SED}} \frac{d[C]}{dx}, \]

where the term $d[C]/dx$ describes the concentration gradient between the bottom water and the first porewater sample of the surface sediment divided by the corresponding sediment depth (0.25 or 0.5 cm), and $\dot{\phi}$ is the porosity of the surface sediment. The diffusion coefficient, $D_{\text{SED}}$, for sediments was calculated accordingly:

\[ D_{\text{SED}} = D_{\text{SW}}/\theta^2. \]

The diffusion coefficients for TPO\textsubscript{4} and Fe\textsuperscript{2+} under standard conditions ($D_{\text{SW}}$: 298.15 K and 1 bar) were taken from Li and Gregory (1974) and corrected for the in situ temperature and pressure using the Stokes–Einstein equation. The tortuosity ($\theta^2$) was derived from the sediment porosity according to the modified Weissberg formulation (Boudreau, 1996):

\[ \theta^2 = 1 - \ln(\phi^2). \]

Uncertainty in the diffusive flux was calculated at station I as the difference between the fluxes calculated from two separate sediment cores. For the other stations, only one core was available to calculate the flux.

3.6 Mass balance of benthic phosphorus cycling

To investigate benthic P cycling quantitatively, a mass balance was developed considering P input, P burial, and P release. The equations for the P mass balance calculations are shown in Table 2. Under steady-state conditions, the total P rain rate should balance the P buried in the sediments and the benthic TPO\textsubscript{4} flux (Eq. 4 in Table 2). The rates were calculated using the POC/Al ratio of the water-column particles that were taken as close as possible to the seafloor at each station (2–5 m above ground) and the POC rain rate (RR\textsubscript{POC}) (Table 4 and the Supplement). RR\textsubscript{POC} for the same stations along the 12° S transect were previously calculated by Dale et al. (2015) as the sum of the measured benthic DIC flux and the POC accumulation rate. The terrigenous P input (RR\textsubscript{Pterr}) (Eq. 7 in Table 2) can be estimated by multiplying the solid-phase Al concentration of the first sediment sample by the mass accumulation rate (Dale et al., 2015) and a P / Al ratio of 0.02 that characterizes the P / Al ratio of riverine particles originating from the continent (Viers et al., 2009).

The P burial flux ($F_{\text{Pbur}}$) (Eq. 8 in Table 2) was calculated by multiplying the mass accumulation rate (MAR) and the average solid-phase P concentration of the first 11 sediment
Figure 2. (a) Concentration profiles of TPP, PIP, POP, and POC of the water-column particles and the surface sediments along the 12° S transect. Water-column particle concentrations (upper panel) are given in μmol L⁻¹ and surface sediment concentrations (lower panel) are shown in μmol mg⁻¹. Note that the water-column particle concentrations shown for station VI (244 m) at 10 m water depth are ~5 times higher than at the other stations. (b) Ratios of POC to TPP, PIP, and POP (POC/xP) along the 12° S depth transect of water-column particles and surface sediments (0–5.5 cm depth) of station I to VIII (74 to 407 m).

centimeters (P$_{11}$) (11 cm is due to our sampling resolution). This approach was chosen according to Dale et al. (2015), who also calculated the POC accumulation rate for the OMZ stations (i.e., not on the shelf or below the OMZ) using the elemental average concentration. MAR (Eq. 9 in Table 2) was calculated from the sedimentation rate ($\omega$acc in cm yr$^{-1}$), dry bulk density ($\rho$dry; in g cm$^{-3}$), and the average porosity of the sediments at the lower core end. Sedimentation accumulation rates were determined from particle-bound$^{210}$Pb$_{a}$ measurements using a modeling approach. A detailed method description and the values used for this work are published in Dale et al. (2015). The error derived from modeling the sedimentation rate was given as 20 % and propagates to all subsequent calculations where it was used.

3.7 Freeze–thaw experiments

In order to determine the amount of polyphosphate stored in sulfide-oxidizing bacteria, foraminifera, and other bacteria, we conducted additional sediment experiments at all transect stations except station IV. Sediments from MUC corers were sliced into 1 cm thick slices from the surface sediment to 10 cm sediment depth. Before phosphate analysis, sediment slices were repeatedly frozen at $-80$°C and defrosted in order to burst microbial cells and release the internally stored P to the porewater.

3.8 Molecular quantification of filamentous bacteria

In order to quantify the abundance of filamentous microorganisms at the benthic boundary layer, we used a molecular approach. Nucleic acid purification was performed on 0.5 g sediment following established protocols (Bertics et al., 2013). DNA was quality checked on an agarose gel and quantified using a NanoDrop spectrophotometer (Pepqlab, Erlangen, Germany). The 16S rDNA fragments were taken from a previously generated metagenome from this region (GenBank Bioproject PRJNA280940) and the respective sequence countings were deposited at GenBank (ID KU312264-KU312267). Sequencing was carried out in the Institute of Clinical Molecular Biology at Kiel University. Sequences were analyzed using a Clustal W alignment tool on Mega 6 (Tamura et al., 2013). A qPCR primer and probe set was established using the Primer Express software (Life Technologies, Carlsbad, USA), a 2.5 µmol µL$^{-1}$ TaqMan probe, 5 pmol µL$^{-1}$ of each primer, 400 ng µL$^{-1}$ bovine serum albumin (to avoid PCR inhibition without affecting standard curves or detection limits), 3 µL PCR water, and 5 µL of either standard DNA or environmental sample. A plasmid containing the target sequence was used to generate a standard dilution series for absolute quantification. The melting temperature was set to 50°C. A theoretical detection limit of four copies per PCR reaction was calculated. The results of the analysis are given in copies g$^{-1}$ of 16S rDNA sequences of sulfur bacteria that are related to *Marithioploca*.

4 Results

4.1 P composition of water-column particulate matter and surface sediments

4.1.1 TPP, PIP, POP and POC concentrations

The TPP concentrations ranged from 0.02 to 0.2 µmol L$^{-1}$ in the water-column particles and from 0.04 to 2.37 µmol mg$^{-1}$ in the surface sediments (Supplement). Overall, the profiles along the transect show no significant trends through the water column (Fig. 2a). The highest concentrations occurred in the surface waters around 10 m of water depth. At stations I, V, and VIII (74, 195, and 407 m), the TPP concentrations slightly increased close to the seafloor, whereas at the other stations no such trend was observed (Fig. 2a and the Supplement). In the sediments the TPP concentrations slightly decreased with sediment depth, except at station VIII (407 m). Here, the highest sedimentary TPP concentrations across the transect were found at 2.25 cm sediment depth (17 µmol mg$^{-1}$). Below that depth the concentrations decreased, but remained high compared to the other stations.

The PIP and POP concentrations (water column and sediments) generally followed the trends of the TPP profiles and contributed roughly equally to TPP. The PIP fraction accounted for 21 to 74 % of TPP in the water-column particles (Fig. 3), similar to that reported in previous studies (Paytan et al., 2003; Faul et al., 2005; Benitez-Nelson et al., 2007; Lyons et al., 2011; Sekula-Wood et al., 2012). At stations I, VI, and VIII (74, 244, and 407 m), the PIP concentrations were larger than POP. The opposite occurred at station V (195 m), where the POP fraction was clearly larger than the PIP fraction throughout the entire water column. In comparison to the water-column particles, the sedimentary PIP fraction was larger than POP in most samples, reaching between 48 and 98 % of TPP. However, the strongest deviation between PIP and POP concentrations was found in the sediments of station VIII (407 m), where the PIP concentration was up to a factor of $\sim$ 50 larger than the POP concentration and comprised up to 98 % of TPP. XRD data from that station revealed that 7 to 16 wt % of the sediments consisted of apatite and other P-bearing minerals (data not shown).
Figure 3. Average distribution of POP and PIP (%) per station in the water-column particles and in the top 5.5 cm of the sediments.

The particulate organic carbon concentrations ranged from 2.2 to 17.6 µmol L$^{-1}$ in the water-column particles and from 2.8 to 13.4 µmol mg$^{-1}$ in the surface sediments. Within the water column the highest concentrations occurred in the surface water samples, while the concentration measured at station VI (244 m) is about 5 times higher compared to other samples from the same water depth. Below the surface water concentration peak, the concentrations were on average 7 µmol L$^{-1}$ with distinct depth trends. The sedimentary POC concentrations were on average 7.7 µmol mg$^{-1}$ for the whole transect. Overall, the POC concentrations showed little variability with sediment depth. At stations III, VI, and VIII (128, 244, and 407 m), the concentrations increased slightly with sediment depth. At station I (74 m) the concentration decreased slightly with sediment depth.

4.2 Particulate organic carbon to phosphorus (POP, PIP, TPP) ratios

4.2.1 POC / POP ratios

The molar POC / POP ratios of the water-column particles and of the surface sediments were consistently higher than the Redfield ratio of 106 (Redfield, 1963) at all stations (Fig. 2b). The average POC / POP ratio of the water-column particles was 202 ± 29. There was no clear trend through the water column, except for slightly increasing ratios between the deepest water-column samples and the surface sediment samples. In the surface sediments, the ratios increased within the upper 6 cm with an average POC / POP ratio of 303 ± 77. Station VIII (407 m) is an exception, and here the ratio decreased to 81 within the first ~2 cm of sediment and then strongly increased to ≥ 600.

4.2.2 POC / PIP ratios

The average POC / PIP ratio of the water-column particles was 248 ± 34. Similar to the POC / POP ratios, there was no significant trend through the water column. At the majority of the stations, the ratios decreased close to the seabed. The average POC / PIP ratio of the surface sediments was 184 ± 34 and almost invariable with water depth. In the sediments, the ratios showed no significant down-core trend. At station VIII (407 m) the ratio in the sediments showed a dramatic decrease compared to the water column, with an average of 7, similar to the POC / TPP ratios (below).

4.2.3 POC / TPP ratios

The POC / TPP ratios of the water-column particles and surface sediments consistently varied around the Redfield ratio. The exceptions are station I (74 m) and the deepest station (station VIII, 407 m). At station I, the sediments showed significantly lower than Redfield ratios, with an average of 69 in the surface sediments. The average POC / TPP ratio of the surface sediments at station VIII (407 m) was 7. Between the deepest water-column sample and the first sediment sample, the POC / TPP ratios were rather constant without a consistent trend, again with the exception of station VIII (407 m), where the ratios decreased sharply.

4.3 In situ benthic chamber fluxes

The benthic lander TPO$_4^-$ fluxes ($F_{TPO_4}$) are presented in Table 3 and Fig. 4a. Positive fluxes are defined as directed from the sediments into the water column. The highest TPO$_4^-$ flux along the depth transect of 1.04 ± 0.31 mmol m$^{-2}$ d$^{-1}$ occurred at station I (74 m). Below 74 m water depth, fluxes decreased by at least a factor of 3 to 0.2–0.3 mmol m$^{-2}$ d$^{-1}$ at 144 m water depth. Measurements at station V (195 m) showed a slightly increased TPO$_4^-$ flux of 0.44 ± 0.07 mmol m$^{-2}$ d$^{-1}$ at 144 m water depth. Measurements at station V (195 m) showed a slightly increased TPO$_4^-$ flux of 0.44 ± 0.07 mmol m$^{-2}$ d$^{-1}$, while the fluxes measured at 244 m (station VI) and 306 m (station VII) decreased to the levels mentioned before. At 407 m water depth (station VIII), the TPO$_4^-$ flux was negative, indicating a phosphate uptake by the sediment. Below the OMZ, the fluxes increased to slightly positive values, but remained low at 0.06 and 0.02 ± 0.02 mmol m$^{-2}$ d$^{-1}$.
Table 3. In situ benthic chamber TPO$_4$ fluxes in mmol m$^{-2}$ d$^{-1}$ along the 12$^\circ$S transect. The numbers are shown as an average and standard deviation calculated from the minimum and maximum fluxes determined from two benthic chambers.

<table>
<thead>
<tr>
<th>Station</th>
<th>Water depth (m)</th>
<th>$F_{\text{TPO}_4}$ (mmol m$^{-2}$ d$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I BIGO1_2</td>
<td>74</td>
<td>1.04 ± 0.31</td>
</tr>
<tr>
<td>II BIGO1_5</td>
<td>101</td>
<td>0.35 ± 0.01</td>
</tr>
<tr>
<td>III BIGO2_4</td>
<td>128</td>
<td>0.30 ± 0.05</td>
</tr>
<tr>
<td>IV BIGO1_1</td>
<td>141</td>
<td>0.23*</td>
</tr>
<tr>
<td>V BIGO1_4</td>
<td>195</td>
<td>0.12*</td>
</tr>
<tr>
<td>VI BIGO2_2</td>
<td>243</td>
<td>0.44 ± 0.07</td>
</tr>
<tr>
<td>VII BIGO2_1</td>
<td>306</td>
<td>0.26 ± 0.04</td>
</tr>
<tr>
<td>VIII BIGO2_5</td>
<td>409</td>
<td>−0.07*</td>
</tr>
<tr>
<td>IX BIGO2_3</td>
<td>756</td>
<td>0.06*</td>
</tr>
<tr>
<td>X BIGO1_3</td>
<td>989</td>
<td>0.02 ± 0.02</td>
</tr>
</tbody>
</table>

* Only one benthic flux was measured.

4.3.1 Comparison of benthic chamber TPO$_4$ fluxes and diffusive TPO$_4$ fluxes

The measured benthic chamber TPO$_4$ fluxes and the calculated diffusive TPO$_4$ fluxes showed large discrepancies. The calculated diffusive fluxes were consistently higher than the benthic fluxes (Table 4, Fig. 4b). In contrast to the in situ measured benthic chamber TPO$_4$ release rates, the calculation of diffusive TPO$_4$ fluxes relies on bottom water and porewater PO$_4^{3-}$ concentrations. A subsurface PO$_4^{3-}$ peak occurred at all stations in the uppermost depth interval at 0–0.25 cm, causing a large concentration gradient between the bottom water and the porewater PO$_4^{3-}$ concentrations (Fig. 7). Even though the measured benthic TPO$_4$ fluxes exceeded the fluxes that could be generated by TPP degradation by a factor of approximately 6, the diffusive TPO$_4$ fluxes were still higher (Table 4). Hence, the diffusive TPO$_4$ flux will be referred to as potential TPO$_4$ flux in the following.

4.4 TPP burial fluxes and TPP burial efficiency

The P burial fluxes decreased with increasing water depth (Table 4). Station I (74 m) showed by far the highest P burial flux, with 0.23 mmol m$^{-2}$ d$^{-1}$. In contrast, the P burial efficiency at this station (Eq. 10) was comparatively low, reaching only approximately 26%. At station VIII (407 m), the TPP burial flux was 13 mmol m$^{-2}$ d$^{-1}$, and the P burial efficiency exceeded 100% due to the uptake of dissolved P from ambient bottom waters.

4.5 Molecular analysis and relative abundance of filamentous sulfur bacteria

Molecular analysis indicated the presence of Marithioploca-related bacteria (Salman et al., 2011) in the surface sediments (0–5 cm). Our analysis may, however, underestimate the absolute abundance of these organisms due to a self-splicing mechanism of the 16S rDNA gene (Salman et al., 2012) and is therefore only indicative for their relative abundance.

The relative abundance of Marithioploca-related bacteria decreased with increasing water depth (Table 4). The highest relative abundance with more than 4000 copies g$^{-1}$ sediment was found at station I (74 m), decreasing by more than a factor of 20 to 190 copies g$^{-1}$ sediment at station VIII (407 m).
Table 4. Measured and calculated parameters for the P mass balance along the 12° S transect. The numbers in front of key parameters correspond to equations in Table 2.

<table>
<thead>
<tr>
<th>12° S</th>
<th>Transect section I</th>
<th>Transect section II</th>
<th>Phosphorite formation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Station I</td>
<td>Station III</td>
<td>Station IV</td>
</tr>
<tr>
<td>74 m</td>
<td>128 m</td>
<td>141 m</td>
<td>195 m</td>
</tr>
<tr>
<td>Benthic chamber TPO flux (FP_{TPO}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>1.10 ± 0.4</td>
<td>1.07 ± 0.6</td>
<td>1.05 ± 0.5</td>
</tr>
<tr>
<td>Benthic diffusive TPO flux (potential flux) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.99 ± 0.3</td>
<td>0.96 ± 0.3</td>
<td>0.95 ± 0.4</td>
</tr>
<tr>
<td>Relative abundance of Marithiola-related bacteria copies g&lt;sup&gt;-1&lt;/sup&gt; (0.5 cm sediment depth)</td>
<td>4159</td>
<td>1687</td>
<td>3072</td>
</tr>
<tr>
<td>Benthic chamber DIC flux (FP_{DIC}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>68.0 ± 21</td>
<td>42.0 ± 7</td>
<td>8.0 ± 0</td>
</tr>
<tr>
<td>POC rain rate (RR_{POC}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>79.5 ± 33</td>
<td>28.2 ± 12</td>
<td>10.5 ± 3</td>
</tr>
<tr>
<td>Sediment accumulation rate (s_{sed}) cm yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.45 ± 0.09</td>
<td>0.2 ± 0.04</td>
<td>0.04 ± 0.008</td>
</tr>
<tr>
<td>Mass accumulation rate (MAR&lt;sub&gt;2&lt;/sub&gt;) g m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>1800 ± 160</td>
<td>600 ± 120</td>
<td>128 ± 26</td>
</tr>
<tr>
<td>Ratios for particulate matter from the water column (2 to 5 m above the sea floor):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>POC / TPP&lt;sup&gt;1&lt;/sup&gt;</td>
<td>56 ± 4</td>
<td>88 ± 10</td>
<td>94 ± 10</td>
</tr>
<tr>
<td>POC / POP&lt;sup&gt;2&lt;/sup&gt;</td>
<td>197 ± 17</td>
<td>125 ± 34</td>
<td>291 ± 79</td>
</tr>
<tr>
<td>TPP rain rate (RR_{TPP}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>1.00 ± 0.31</td>
<td>0.40 ± 0.09</td>
<td>0.11 ± 0.02</td>
</tr>
<tr>
<td>TPP ratio (FP_{TPP}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.39 ± 0.14</td>
<td>0.22 ± 0.04</td>
<td>0.04 ± 0.01</td>
</tr>
<tr>
<td>POP rain rate (RR_{POP}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.61 ± 0.18</td>
<td>0.18 ± 0.05</td>
<td>0.07 ± 0.02</td>
</tr>
<tr>
<td>POP ratio (FP_{POP}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>126 ± 17</td>
<td>149 ± 29</td>
<td>142 ± 3</td>
</tr>
<tr>
<td>(7) Terrigenous P input (RR_{terr})</td>
<td>0.10 ± 0.02</td>
<td>0.01</td>
<td>0.02</td>
</tr>
<tr>
<td>(8) Burial flux (FP_{burial}) at 11 cm sediment depth mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.23 ± 0.09</td>
<td>0.02</td>
<td>0.08</td>
</tr>
<tr>
<td>Avg. Al conc. (0.1 cm sediment) (Al&lt;sub&gt;c&lt;/sub&gt;) mmol g&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.99 ± 0.70</td>
<td>1.10</td>
<td>0.97</td>
</tr>
<tr>
<td>Avg. P conc. (0.1 cm sediment) (P&lt;sub&gt;c&lt;/sub&gt;) mmol g&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.05 ± 0.05</td>
<td>0.07</td>
<td>0.09</td>
</tr>
<tr>
<td>(9) P burial efficiency (PBE) at 11 cm sediment depth</td>
<td>26 ± 4</td>
<td>23 ± 4</td>
<td>23 ± 5</td>
</tr>
<tr>
<td>(10) P release from POP degradation according to Redfield (FP_{Redfield}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.62 ± 0.02</td>
<td>0.19 ± 0.06</td>
<td>0.08 ± 0.01</td>
</tr>
<tr>
<td>(11) P release from POP degradation (FP_{PP}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.52 ± 0.16</td>
<td>0.14 ± 0.05</td>
<td>0.06 ± 0.01</td>
</tr>
<tr>
<td>(12) P release from TPP degradation (FP_{TPP}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.87 ± 0.17</td>
<td>0.3 ± 0.1</td>
<td>0.09 ± 0.01</td>
</tr>
<tr>
<td>Diffusive Fe&lt;sup&gt;2+&lt;/sup&gt; flux (FP_{Fe}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.04 ± 0.02</td>
<td>0.01</td>
<td>0.02</td>
</tr>
<tr>
<td>(13) P-deficit to outbalance the P budget (FP_{deficit}) mmol m&lt;sup&gt;-2&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.04 ± 0.002</td>
<td>0.001</td>
<td>0.002</td>
</tr>
</tbody>
</table>

<sup>1</sup> This study; <sup>2</sup> published data from Dale et al. (2015)

5 Discussion

5.1 POC / xP ratios in water-column particles and sediments

In order to characterize the fate of P in oxygen-deficient waters and sediments, we determined POC / xP ratios from both environments. Previous studies focused either on the water column or on the sediments (Anderson et al., 2001, and references therein; Benitez-Nelson et al., 2004; Faul et al., 2005; Jilbert et al., 2011; Lyons et al., 2011; Sekula-Wood et al., 2012). The present data set provides a more complete insight into compositional changes and allows us to more rigorously constrain the sedimentary P mass balance compared to earlier studies (Ingall and Jahnke, 1994; Mort et al., 2010; Kraal et al., 2012; Noffke, 2014).

Both water-column particles and the surface sediments from the Peruvian OMZ displayed POC / POP ratios above Redfield, indicating depletion of organic P relative to organic C. Similar observations have been reported before from this and other regions of the ocean (Loh and Bauer, 2000; Benitez-Nelson et al., 2004; Faul et al., 2005; Franz et al., 2012, and others). Preferential remineralization of P phases from sinking particles should lead to increasing POC / POP ratios with increasing water depth, as observed in oxygenated areas of the ocean (reviewed by Ruttenberg, 2014). However, in the anoxic Cariaco Basin, no such preferential P mineralization was noted (Benitez-Nelson et al., 2004). Our results also showed no clear decrease in the POC / POP ratio in the water column, with the possible exception of stations I and VIII. Higher than Redfield POC / POP ratios have been observed previously in Peru, and may instead be driven by the C-to-P composition of the diatomaceous phytoplankton communities (Franz et al., 2012) rather than preferential P dissolution or other controls such as the input of terrestrial plant material with high POC / POP ratios.

Preferential POP over POC remineralization in anoxic sediments remains controversial (Ingall and Van Cappellen, 1990; Ingall and Jahnke, 1994; Colman et al., 1997; Anderson et al., 2001; McManus et al., 1997; Ruttenberg, 2003; Jilbert et al., 2011). Our results showed no clear trend across all stations. At stations I, IV, and VI (74, 141, and 244 m), the POC / POP increased with sediment depth, indicating preferential POP over POC remineralization. At station III (128 m) this trend occurred only in the upper cm, and at station V (195 m) no preferential POP over POC remineralization was found. POC / POP ratios at station VIII (407 m) showed a P enrichment over the upper 2 cm of the sediment. Below this depth, there was a sudden increase in POC / POP ratios, which is likely due to intense POP remineralization and subsequent authigenic formation of phosphorites (sink-switching; see Sect. 5.2.3).

Another interesting finding pointing to sink-switching can be found in the POC / POP and POP / PIP ratios of the deepest water-column particle sample and the first sediment sample at stations I, IV, VI, and VIII (74, 141, 244, and 407 m).
Increasing POC / POP and decreasing POC / PIP ratios indicate that POP is converted into PIP while the TPP content of the sediment is conserved (Fig. 2a and b). In agreement with similar findings in anoxic sediments (Ruttenberg and Berner, 1993; Faul et al., 2005, and others), we assume that the observed POP to PIP transformation is due to the formation of CFA. However, our results emphasize that sink-switching is obviously occurring at the interface between bottom waters and sediments.

Interestingly, the water-column POC / TPP ratios were close to the Redfield ratio. This could be an effect of surface adsorption of P on phytoplankton as previously described by Sanudo-Wilhelmy et al. (2004). Those authors investigated different species of *Trichodesmium* from the Atlantic Ocean and found that the intracellular P pool was strongly depleted relative to C, whereas the combination of the intracellular plus surface adsorbed P was close to the Redfield ratio. Although we did not exactly differentiate between internal and external P pools, and considering that additional P sources like terrigenous P are negligible (see Sect. 5.2.1), our results generally support the findings of Sanudo-Wilhelmy et al. (2004). However, future studies are required to substantiate this hypothesis.

Sedimentary POC / TPP ratios were also close to Redfield, except at stations I and VIII (74 and 407 m). Thus, on the one hand, the sink switching mechanism operates efficiently under low-oxygen conditions and, on the other hand, the relative degradation of TPP and POC do not change from the water column to the sediments (Fig. 2b). This is an important finding because at first sight this observation seems to be inconsistent with the long-standing paradigm that low-oxygen conditions promote the enhanced release of dissolved P from sediments (Van Cappellen and Ingall, 1996). However, it should be noted that marine sediments covered by oxygenated bottom waters display molar POC / TPP ratios ranging from about 10 to 50 (Baturin, 2007). These ratios are much lower than Redfield because P is retained in sediments via adsorption, authigenic mineral precipitation such as Fe (oxyhydr)oxides and terrigenous P was of minor importance for the total sedimentary P inventory. However, authigenic Ca–P with a high amount of fish-P accounted for up to 47 % of the total P in sediments down to 300 m water depth and for up to 70 % in sediments below 300 m water depth. Consequently, Noffke (2014) suggested that authigenic Ca–P phases are an additional major P source besides organic P for benthic P release in the Peruvian OMZ.

Their mass balance approach was solely based on benthic work and has large uncertainties regarding the particulate P input from the water column. Our approach includes the particulate organic and inorganic P input from the water column, benthic P flux measurements, and the P burial fluxes, and is based on the steady-state assumption that the P input has to be sufficient to maintain the benthic P flux and P burial flux. The P mass balance calculations (Table 4) illustrate the variability in TPO₄ release and burial as well as in the magnitude of particulate organic (Fig. 5a) and particulate inorganic P input (Fig. 5b) across the transect. Following the general assumption that POP is the major P phase delivered to the sediments (Delaney, 1998; Filippelli, 2008), we first calculated whether the rain rate of POP (RRPOP) to the seafloor can account for the measured benthic P fluxes (F₅TPO₄) and P burial fluxes (F₅Pbur) (Fig. 5a, Table 4). However, as already suggested by Noffke et al. (2012) and Noffke (2014), the POP fraction is far too small to balance the measured benthic fluxes in the Peruvian OMZ. The POP rain rates calculated along the transect can account for only 25 to 48 % of the measured TPO₄ fluxes (Fig. 5a), suggesting the likely presence of an additional inorganic source of dissolved phosphate (Noffke, 2014). Similar to previous studies (Paytan et al., 2003; Faul et al., 2005; Benitez-Nelson et al., 2007; Lyons et al., 2011; Sekula-Wood et al., 2012), we found that
the PIP fraction in water-column particles ranging from 75 to 407 m water depth comprises between 21 and 74 % of TPP (Fig. 3). In the sediments, the average PIP fraction rises to 48–98 % of TPP (Fig. 3). Furthermore, POC and PIP were correlated \( (r^2 = 0.74) \) in the water-column particles, indicating highly reactive material.

The mass balance approach including the PIP rain rate to the seafloor (Fig. 5b) allows the depth transect to be divided into two sections. The transect section I (station I, 74 m, and III, 128 m) is characterized by high P input and release rates. The calculations on the P budget show a balance between the particulate P input, the benthic P fluxes, and the P burial fluxes within the error margin (±20 %). In transect section II (stations IV, 141 m, V, 195 m, and VI, 244 m), the P input decreases drastically (Fig. 5b, Table 4), whereas the benthic P fluxes are still comparatively high. The distinct mismatch in P input and P output prevails only 37 to 53 % of the measured TPO\(_4\) fluxes and calculated burial fluxes. This leads to the question: what drives the excess TPO\(_4\) release in the core of the Peruvian OMZ?

5.2.1 Additional P input

Besides the particulate matter raining to the sediments, other potential P sources can be considered. Firstly, riverine transported material from the continent may be an additional source of P to the sediments. Due to fast sinking speed and laterally dominated transport it is possible that this P fraction is at least underrepresented on the filter samples. In order to provide a maximum estimate for the contribution of the terrigenous P input to the sediments, this fraction was calculated using the mass accumulation rate of Al in the first centimeter of sediment and an average molar P / Al ratio (Table 2) of 0.02 for riverine suspended particles (Viers et al., 2009). The resulting terrigenous P flux accounted for 5–19 % of the total P input, which is insufficient to explain the observed discrepancies in the P budget of transect section II (Table 4, Fig. 4c, 5b).

Laterally transported particles enriched in P from the very shallow shelf could be an additional P source (e.g., Jahnke, 1990). However, the particles would need to be strongly enriched in P, which is not the case. In addition, this would have to be reflected in the POC / TPP ratios of the surface sediments in transect section II (Fig. 2b). The ratios are not, or are only slightly enriched in TPP compared to the water-column particles, which leads to the conclusion that lateral transport of P-enriched particles to the sediments is an unlikely candidate for the missing P source.

Another alternative is the existence of an additional PIP phase supplied by fast-sinking material (e.g., P containing fish scales; Suess, 1981) that was not sampled during CTD casts, and hence is underrepresented on our filter samples. Díaz-Ochoa et al. (2009) showed that fish P can make up to 20 % of the total sedimentary P inventory in the shelf sediments of the Peruvian OMZ. Fish P input should be depicted in low sedimentary POC / TPP ratios compared to the water-column particles. Since this difference is not observed, fish debris can be ruled out for closing the P budget during the sampling campaign. Theoretically, sediments need to be composed of particles having POC / TPP ratios between 11 ± 1 and 25 ± 12 (Table 4) to maintain the measured P release rates in transect section II. It seems unlikely that the mismatch in the P mass balance is caused by additional particles deposited at the seabed since their POC / TPP ratio would need to be much lower than any value observed in our data set.

5.2.2 Non-steady-state scenarios – internal sedimentary P pools

Besides an additional P input to the sediments from the water column, episodic dissolution of particulate P within the sediment could contribute to the excess P release (Nof-fke et al., 2012). This could include P solubilized from Fe (oxyhydr)oxides or the degradation of internally stored polyphosphates within sulfide-oxidizing bacteria. Driving factors could include the temporal variability in bottom-water oxygen and nutrient concentrations induced by the passage of internal Kelvin waves and/or interannual variability related to El Niño and La Niña (Guitérrez et al., 2008).

It is well recognized that the sedimentary cycles of Fe and P are strongly linked (e.g., Sundby et al., 1986). Fe (oxyhydr)oxides are expected to be important carriers for phosphorus from the water column to the sediments. Following dissolution of solid Fe minerals in the sediments, the adsorbed P is released into the porewaters. However, in the Peruvian OMZ, oxygen concentrations in the water column are generally below the detection limit. Consequently, Fe (oxyhydr)oxides are likely already dissolved in the water column and not such an important P source to the sediments. An estimate of phosphate released during the reduction of Fe (oxyhydr)oxides can be calculated from the diffusive Fe\(^{2+}\) fluxes and the molar Fe / P ratio typically found in Fe (oxyhydr)oxides. The diffusive Fe\(^{2+}\) fluxes were in the range of 0 to 0.03 mmol m\(^{-2}\) d\(^{-1}\) during the M92 cruise (Table 4), and the molar Fe / P ratio of Fe (oxyhydr)oxides in sediment is around 10 (Slomp et al., 1996). The calculation of the TPO\(_4\) release rates from Fe (oxyhydr)oxides (Eq. 14 in Table 2) results in a flux of 0.003 mmol PO\(_4^{3-}\) m\(^{-2}\) d\(^{-1}\) (Table 4, Fig. 4c, and 5b), which is equivalent to less than 5 % of the benthic P flux and burial flux.

An additional internally activated P pool is phosphate released from large sulfur-oxidizing bacteria, e.g., Beggiatoa (Sannigrahi and Ingall, 2005; Brock and Schulz-Vogt, 2011). These microorganisms store P in the form of intracellular polyphosphate granules when terminal electron acceptors for sulfide oxidation are available (oxic and nitrogenous conditions) and release dissolved P during periods when these oxidants are scarce (anoxic conditions). Hence, it is generally
Figure 5. Mass balance calculations and measured benthic TPO$_4$ fluxes for stations I to VIII (74 to 407 m). All fluxes are in mmol m$^{-2}$ d$^{-1}$.
(a) POP rain rates, TPO$_4$ fluxes, and P burial rates only. The number in percent denotes missing P needed to sustain the benthic TPO$_4$ fluxes.
(b) Mass balance calculations including the POP and PIP rain rates, the terrigenous P input, P release from the reductive dissolution of Fe (oxyhydr)oxides and the benthic TPO$_4$ fluxes into the bottom waters, as well as the P burial rates.

assumed that they strongly affect benthic TPO$_4$ fluxes in a system with frequently changing bottom water redox conditions (Ingall and Jahnke, 1994; Sannigrahi and Ingall, 2005; Schulz and Jorgensen, 2005; Brock and Schulz-Vogt, 2011; Dale et al., 2013). Polyphosphates have been shown to be an important P pool in the sediments of highly productive upwelling areas (Kraal et al., 2015). Along the depth transect at Peru, dense mats of sulfur bacteria related to *Marithioploca* were observed on the sediments during video-launched MUC and benthic lander deployments down to 300 m water depth (Dale et al., 2015). Similar findings that extended distributions of microbial mats along a latitudinal transect at 11° S were also previously described by Mosch et al. (2012) and Gutiérrez et al. (2008). Although these organisms are not known to store polyphosphates like their close relatives, *Beggiaota* spp. and *Thiomargarita* spp. (Høgslund et al., 2009; Holmkvist et al., 2010), our findings provide circumstantial indications of P uptake and release by *Marithioploca*-related bacteria.

Firstly, we found that the relative abundance of copies of *Marithioploca*-related bacteria per g$^{-1}$ sediment and the measured benthic TPO$_4$ release rates correlate linearly ($r^2 = 0.92$, Fig. 6). This finding supports the suggestion that bacteria exert an important control on benthic P fluxes. Secondly, a comparison of the in situ measured benthic P fluxes and the diffusive P fluxes calculated from the difference of the TPO$_4$ bottom water concentration and the TPO$_4$ porewater concentration of the surface sediments revealed large differences (Fig. 4b). Such a difference could be explained by the lysis of bacterial cells during sample retrieval followed by the release of the internally stored polyphosphate pool into the porewater. Following this argument, the diffusive P fluxes cannot be taken as real fluxes, but as a measure for potential maximum release rates of P by *Marithioploca*-related bacteria. It should be noted that the potential fluxes are more than sufficient to compensate for the missing P fraction in transect section II (ranging from 0.5 to 1.6 mmol m$^{-2}$ d$^{-1}$; Table 4, Fig. 4b).

In addition to the established porewater extraction procedure, we carried out freeze–thaw experiments to quantify the amount of P stored in sulfide-oxidizing bacteria (see the method in Sect. 3.7). The released polyphosphates from the
Marithioploca release timescales by otherwise and awaits further study. P storage by foraminifera has not been demonstrated previously, the foraminifera observed at station VIII (407 m), are tentatively, the foraminifera recovered from the Peruvian OMZ, including a station at 12° S from the same water depth (~400 m) close to sampling station VIII. The suboxic bottom waters and low sedimentation rates in that area seem to be favorable for phosphorite formation close to the sediment–water interface (Arning et al., 2009b). Cosmidis et al. (2013) suggested three mechanisms for how high porewater phosphate concentrations that are essential for the phosphogenesis can be generated in the sediments: (1) remineralization of organic matter mainly through bacterial sulfate reduction releasing phosphate to the porewaters, (2) reductive dissolution of Fe (oxyhydr)oxides and the release of adsorbed P, and (3) synthesis of internally stored polyphosphates by large sulfide-oxidizing bacteria. Using the same mass balance approach as presented before, we calculate a P accumulation rate of 33 ± 4 mmol m⁻² yr⁻¹ at station VIII, where most of the P is derived from ambient bottom waters (26 mmol m⁻² yr⁻¹). Hence, our data suggest that the phosphorite nodules at this station contain phosphate that originates predominantly from ambient bottom waters. Additionally, as already mentioned, sediments at station VIII were covered by benthic foraminifera instead of mat-forming sulfur bacteria. The release of phosphate from frozen samples from this site may indicate that these foraminifera are a source of polyphosphates (see Sect. 3.7 and Fig. 7). Our observations suggest that benthic foraminifera rather than bacterial mats might facilitate the uptake of bottom water phosphate and the formation of phosphorites at this station. However, this remains an open question and should be addressed in future field campaigns.

The P uptake rate of ~26 mmol P m⁻² yr⁻¹ derived from our lander measurements may be compared to previous estimates on phosphorite growth rates in the area. Dating of phosphatic laminites yields a P uptake rate of only ~3 mmol P m⁻² yr⁻¹ for a ca. 1 Ma old nodule (Arning et al., 2008). Hence, our data suggest that sulfur bacteria make an important contribution to this missing P source.

5.2.3 Indications for active phosphorite formation

In contrast to the stations between 74 and 244 m water depth characterized by P release, data from station VIII at 407 m water depth indicate the uptake of phosphate from the bottom water. To our knowledge, this is the first time that a downward flux of dissolved phosphate from bottom waters into phosphorite-bearing surface sediments has been documented by in situ benthic flux measurements. Furthermore, the PIP concentrations in the surface sediments of station VIII (Fig. 2a) were 10 to 60 times higher compared to the shallower stations where P was released from the sediments. Taken together, these observations indicate that a PIP phase, likely phosphorite, is precipitating from the porewater phosphate at the time of sampling. This is also reflected in decreasing porewater phosphate concentrations (Fig. 7).

Arning et al. (2008) presented investigations on phosphorites recovered from the Peruvian OMZ, including a station at 12° S from the same water depth (~400 m) close to sampling station VIII. The suboxic bottom waters and low sedimentation rates in that area seem to be favorable for phosphorite formation close to the sediment–water interface (Arning et al., 2009b). Cosmidis et al. (2013) suggested three mechanisms for how high porewater phosphate concentrations that are essential for the phosphogenesis can be generated in the sediments: (1) remineralization of organic matter mainly through bacterial sulfate reduction releasing phosphate to the porewaters, (2) reductive dissolution of Fe (oxyhydr)oxides and the release of adsorbed P, and (3) synthesis of internally stored polyphosphates by large sulfide-oxidizing bacteria. Using the same mass balance approach as presented before, we calculate a P accumulation rate of 33 ± 4 mmol m⁻² yr⁻¹ at station VIII, where most of the P is derived from ambient bottom waters (26 mmol m⁻² yr⁻¹). Hence, our data suggest that the phosphorite nodules at this station contain phosphate that originates predominantly from ambient bottom waters. Additionally, as already mentioned, sediments at station VIII were covered by benthic foraminifera instead of mat-forming sulfur bacteria. The release of phosphate from frozen samples from this site may indicate that these foraminifera are a source of polyphosphates (see Sect. 3.7 and Fig. 7). Our observations suggest that benthic foraminifera rather than bacterial mats might facilitate the uptake of bottom water phosphate and the formation of phosphorites at this station. However, this remains an open question and should be addressed in future field campaigns.

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2009a). These different fluxes may be at least partly explained by the methodological difference (present flux measurement vs. long-term average). However, growth rates determined on modern nodules are broadly consistent with our flux measurements (Burnett et al., 1982).

6 Conclusions

This study aimed to identify the P sources of benthic P release in the Peruvian OMZ. We determined the rain rates of particulate organic phosphorus and particulate inorganic phosphorus as well as benthic P release rates and P burial fluxes.

Our calculations revealed that within the core OMZ particulate phosphorus rain rates cannot account for measured benthic P fluxes and burial fluxes. From systematic analysis of potential P sources, we conclude that periodic P release from sulfur bacteria that store and release P under oscillation redox conditions could strongly modulate benthic P fluxes, and hence explain the missing P source. We visited the area during austral summer when oxygen and nitrate levels were depleted by high export production and respiration. It is possible that the Peruvian OMZ was less reducing prior to our sampling period due to lower respiration rates and/or better ventilation. Thus, we propose that the bacterial mats on the Peruvian margin act as phosphorus capacitors, being discharged during austral summer and recharged during other periods of the year when bottom waters are less reducing, as previously proposed in Dale et al. (2013). This hypothesis could be tested by studying the seasonality of benthic fluxes in the Peruvian upwelling system and P dynamics within the bacterial community.

In addition, measurements at one station (407 m water depth) showed clear indications for the uptake of dissolved phosphate by the sediments facilitating phosphorite formation. Our data imply that most of the P accumulating in these authigenic minerals originates from ambient bottom waters. Since this site was marked by a high abundance of benthic foraminifera, we speculate that phosphate uptake and phosphorite formation may be linked to the presence of these organisms. This requires further study.

There was no clear preferential mineralization of POP relative to POC with depth in the water column. POC / TPP ratios in both water-column particles and sediments were close to Redfield at most sites in the Peruvian OMZ. This observation strongly suggests that the relative burial efficiencies of POC and TPP are similar under low-oxygen conditions. Importantly, it further shows that the sediments underlying the anoxic waters on the Peru margin are not depleted in P compared to Redfield. Rather, they are depleted relative to sediments underlying oxic waters, which show POC / TPP ratios well below Redfield (Wallmann, 2010). Thus, in Peru, a lack of oxygen promotes the intensified release of dissolved P from sediments, whilst preserving a POC / TPP burial ratio that is similar to Redfield.

Our data support the hypothesis that benthic P release is linked via a positive feedback loop to intensified primary production in the surface water and oxygen demand in the water column during periods where bottom water redox conditions promote the release of P from the sediments. However, this positive feedback is limited by the formation of authigenic inorganic P phases that maintains the long-term average POC / TPP burial ratio close to Redfield.
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References


Filippelli, G. M.: The global phosphorus cycle, in Phosphates: Geochemical, Geobiological, and Materials Importance, edited by:
Kohn, M., Rakovan, J., and Hughes, J., Reviews in Mineralogy & Geochemistry, 391–425, 2002.


