



Major Bottom Water Ventilation Events Do Not Significantly Reduce Basin-Wide Benthic N and P Release in the Eastern Gotland Basin (Baltic Sea)

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Redox-sensitive mobilization of nutrients from sediments strongly affects the eutrophic state of the central Baltic Sea; a region associated with the spread of hypoxia and almost permanently anoxic and sulfidic conditions in the deeper basins. Ventilation of these basins depends on renewal by inflow of water enriched in oxygen (O₂) from the North Sea, occurring roughly once per decade. Benthic fluxes and water column distributions of dissolved inorganic nitrogen species, phosphate (PO₄³⁻), dissolved inorganic carbon (DIC), sulfide (HS⁻), and total oxygen uptake (TOU) were measured along a depth gradient in the Eastern Gotland Basin (EGB). Campaigns were conducted during euxinic conditions of the deep basin in Aug./Sept. 2013 and after two inflow events in July/Aug. 2015 and March 2016 when O₂ concentrations in deep waters reached 60 μM. The intrusion of O₂-rich North Sea water into the EGB led to an approximate 33 and 10% reduction of the seabed PO₄³⁻ and ammonium (NH₄⁺) release from deep basin sediments. Post-inflow, the deep basin sediment was rapidly colonized by HS⁻ oxidizing bacteria tentatively assigned to the family *Beggiatoaceae*, and HS⁻ release was completely suppressed. The presence of a hypoxic transition zone (HTZ) between 80 and 120 m water depth was confirmed not only for euxinic deep-water conditions during 2013 but also for post-inflow conditions. Because deep-water renewal did not ventilate the HTZ, where PO₄³⁻ and NH₄⁺ fluxes were highest, high seabed nutrient release there was relatively unchanged. Extrapolation of the in situ nutrient fluxes indicated that, overall, the reduction in PO₄³⁻ and NH₄⁺ release in response to deep-water renewal can be considered as minor, reducing the internal nutrient load by 2 and 12% only, respectively. Infrequent inflow events thus have a limited capacity to sustainably reduce internal nutrient loading in the EGB and mitigate eutrophication.

Keywords: major baltic inflows, benthic nutrient fluxes, euxinia, hypoxia, ventilation, sulfur bacteria, Gotland basin, Baltic Sea

INTRODUCTION

The Baltic Sea is a landlocked marginal sea with a narrow connection to the North Sea through the Kattegat. It consists of a series of basins separated by shallow sills and narrow channels. Restricted water exchange with the North Sea and freshwater input from river run-off maintain a strong surface salinity gradient from around 3 in the Bothnian Bay at the northern end to 20 in the Kattegat (Samuelsson, 1996). Density differences result in a strong stratification of the central basins, with a stable halocline located at water depths of 60–80 m (HELCOM, 2009b). Consequently, hypoxia ($O_2 < 63 \mu\text{M}$) has occurred naturally in the deep basins of the Baltic Sea since its formation at about 8000 year BP (Zillén et al., 2008; Conley et al., 2009). With increased terrestrial nutrient inputs, however, the spatial extent and intensity of hypoxia and degree of eutrophication is increasing (Conley et al., 2009; HELCOM, 2009a). Intense efforts backed by the Helsinki Commission have so far failed to significantly reduce eutrophication there (HELCOM, 2009b).

Ongoing chronic hypoxia in the Baltic Sea is partly due to a rapid turnover of phosphorus (P) from hypoxic and anoxic sediments (Conley et al., 2002; Savchuk, 2005; Stigebrandt et al., 2014). Rapid internal P cycling is superimposed on the slow long-term sink of P removal by burial in the sediments (Mort et al., 2010; Viktorsson et al., 2013a; Noffke et al., 2016). As hypothesized by Vahtera et al. (2007), this internal nutrient release delays recovery of the Baltic Proper from eutrophication despite major efforts to reduce the external nutrient load. For this reason, there have been calls to artificially ventilate the deep basins to help permanently sequester P in the sediment as iron-bound minerals and other forms (Stigebrandt and Gustafsson, 2007), although such large scale engineering solutions are not without complications (Conley et al., 2009). Pilot studies in an anoxic fjord have demonstrated the potential for major alterations to benthic nutrient cycles before and after forced bottom water oxygenation (Viktorsson et al., 2013b; Brabandere et al., 2015).

Natural ventilation of the deep central basins of the Baltic Sea exclusively depends on episodic inflow events from the North Sea (Matthäus and Franck, 1992; Stigebrandt, 2003). The physics of these overflows has been investigated intensively (see reviews by e.g., Meier et al., 2006; Reissmann et al., 2009; Omstedt et al., 2014; Mohrholz et al., 2015; and references therein). Saline inflows can be of baroclinic or barotropic type. Barotropic inflows are driven by wind and air-pressure induced sea level differences between the Kattegat and the central Baltic Sea. They mainly occur during autumn and winter when wind forcing is highest. Baroclinic inflows are driven by a salinity gradient between the Kattegat and the Baltic and typically occur during summer under calm wind conditions. Summer inflows usually contribute less to the ventilation of the deep basins. Small inflows are soon diluted on their pathway toward the central Baltic (Mohrholz et al., 2015). Flushing that is sufficiently dense (saline) to reach the central Baltic basins is termed a Major Baltic Inflow event (MBI). Specific weather conditions are a prerequisite for the formation of MBIs. Long lasting easterly winds depressing the sea level by about 10 cm to normal, followed by strong westerly winds push

North Sea water through the Belt Sea to the entrance area of the Baltic proper (Reissmann et al., 2009 and references therein). The strength of inflow events is related to the mass of imported salt, where strong events range between 2 and 3 Gt of salt and moderate inflows between 1 and 2 Gt (Reissmann et al., 2009). Overflows over the Belt Sea sills transport saline water into the entrance areas of the Baltic Sea where they form gravity-driven dense bottom currents. Those are subjected to entrainment, interleaving and boundary mixing strongly affecting dilution of solutes and geochemical processes (Reissmann et al., 2009; cf. their Figure 2). Due to volume conservation, the deep inflows lead to a compensating uplift of water masses in the central Baltic Sea. Stagnation periods in between inflow events cause strong O_2 depletion in the basin bottom waters due to respiration of organic carbon exported from the surface mixed layer. This often leads to fully anoxic conditions below the pycnocline and the build-up of elevated HS^- concentrations (Schincke and Matthäus, 1998).

Before ca. 1980, inflow events were relatively frequent and could be observed on average once a year (Matthäus and Franck, 1992). Since then the frequency of MBIs decreased strongly, and strong events were only recorded in 1993 and 2003 (Matthäus et al., 2008; Mohrholz et al., 2015, cf. their Figure 16). In December 2014, a MBI occurred with a total volume of $\sim 198 \text{ km}^3$ and terminated a long stagnation period following the last MBI event in 2003. This ventilated the bottom water that had been euxinic since 2005 (Nausch et al., 2012). In comparison to previous MBI events, this was ranked the third strongest since 1880 (Mohrholz et al., 2015). The strongest inflow was recorded in 1951 with an estimated volume of 225 km^3 (Mohrholz et al., 2015). The spreading velocity of MBIs into the Baltic proper depends on bathymetry as well as on its density, such that 4–5 months are required for the inflow to reach the Gotland basin (Nehring and Franke, 1981), which is the largest basin in the Baltic proper and the second deepest basin (249 m) in the Baltic Sea. The MBI of December 2014 reached the Eastern Gotland Basin (EGB) in March 2015 and replaced the anoxic deep water (Mohrholz et al., 2016). The deep water below 140 m was completely ventilated until at least May 2015. A moderate MBI triggered in November 2015 reached the EGB in February 2016 and lasted until May 2016 (Mohrholz et al., 2016).

Here, we report on benthic fluxes of nutrients and O_2 in the EGB before and after the strong MBI in December 2014 and the moderate MBI in November 2015. Recent studies of nutrient release measured *in situ* (Viktorsson et al., 2013a; Noffke et al., 2016) and modeled (Almroth-Rosell et al., 2015) from the seabed in the EGB identified the deep sediments as an exceptionally high source of PO_4^{3-} as well as NH_4^+ under euxinic bottom water conditions. However, sediments lying within the HTZ between about 80 and 120 m water depth were also identified as a particularly important zone for the recycling of biogenic material (Noffke et al., 2016). Sediments here were observed to be densely covered with mats of filamentous sulfur bacteria of the family *Beggiatoaceae*. Basin wide release of PO_4^{3-} was extrapolated to 152 kt year^{-1} (Viktorsson et al., 2013a) and 109 kt year^{-1} , from which as much as 70% (76 kt) was released from the HTZ (Noffke et al., 2016). Similar patterns were observed for NH_4^+ (Noffke et al., 2016). These P fluxes are several-fold

TABLE 1 | Locations of the sites of benthic lander (BIGO) deployments in the EGB along with water depth and redox characteristics of the deep water during cruise AL422 (pre-inflow, euxinic bottom water in deep basin, late summer), cruise POS487 (post-inflow, late summer), and cruise AL473 (post-inflow, subsequent winter).

Station	Instrument	Position (°N°E)	Depth (m)	Redox	Date
RV ALKOR CRUISE AL422 (STAGNANT, EUXINIC CONDITIONS IN THE DEEP BASIN)					
651	BIGO-II-6	57°26.26', 20°43.53'	65	Oxycline	08. Sep.2013
584	BIGO-I-2	57°21.80', 20°35.87'	80	HTZ	23. Aug.2013
561	BIGO-II-1	57°20.76', 20°35.32'	95	HTZ	19. Aug.2013
600	BIGO-II-3	57°20.58', 20°34.32'	110	HTZ	26. Aug.2013
658	BIGO-I-6	57°20.59', 20°34.30'	110	HTZ	09. Sep.2013
568	BIGO-I-1	57°18.51', 20°32.99'	123	Anoxic basin	20. Aug.2013
626	BIGO-I-4	57°18.50', 20°33.01'	123	Anoxic basin	05. Sep.2013
642	BIGO-I-5	57°18.50', 20°33.04'	123	Anoxic basin	07. Sep.2013
635	BIGO-II-5	57°14.99', 20°27.13'	140	Anoxic basin	06. Sep.2013
603	BIGO-I-3	57°20.98', 20°28.99'	151	Anoxic basin	27. Aug.2013
618	BIGO-II-4	57°21.05', 20°27.97'	173	Anoxic basin	04. Sep.2013
POSEIDON CRUISE POS487 (DEEP BASIN VENTILATED)					
457	BIGO-I-6	57°26.56', 20°43.34'	63	Oxycline	08. Aug.2015
453	BIGO-II-6	57°21.81', 20°35.85'	79	Oxycline	07. Aug.2015
318	BIGO-I-1	57°21.06', 20°35.88'	80	HTZ	18. Jul.2015
325	BIGO-II-1	57°20.99', 20°35.12'	94	HTZ	19. Jul.2015
410	BIGO-II-4	57°20.86', 20°35.39'	94	HTZ	01. Aug.2015
446	BIGO-II-5	57°20.60', 20°34.35'	108	HTZ	05. Aug.2015
345	BIGO-I-2	57°20.58', 20°34.32'	111	HTZ	22. Jul.2015
354	BIGO-II-2	57°18.42', 20°33.12'	124	Oxygenated	23. Jul.2015
450	BIGO-I-5	57°20.49', 20°29.12'	142	Oxygenated	06. Aug.2015
373	BIGO-I-3	57°20.98', 20°29.03'	151	Oxygenated	25. Jul.2015
420	BIGO-I-4	57°21.05', 20°28.56'	161	Oxygenated	02. Aug.2015
401	BIGO-II-3	57°21.07', 20°27.94'	178	Oxygenated	27. Jul.2015
ALKOR CRUISE AL473 (DEEP BASIN VENTILATED)					
138	BIGO-II-3	57°21.70', 20°35.83'	81	HTZ	22. Mar.2016
92	BIGO-II-1	57°20.81', 20°35.26'	95	HTZ	12. Mar.2016
87	BIGO-I-1	57°20.59', 20°34.29'	109	HTZ	11. Mar.2016
105	BIGO-I-2	57°18.47', 20°33.00'	123	Oxygenated	14. Mar.2016
123	BIGO-I-3	57°20.96', 20°29.00'	151	Oxygenated	20. Mar.2016
115	BIGO-II-2	57°21.04', 20°27.96'	174	Oxygenated	19. Mar.2016

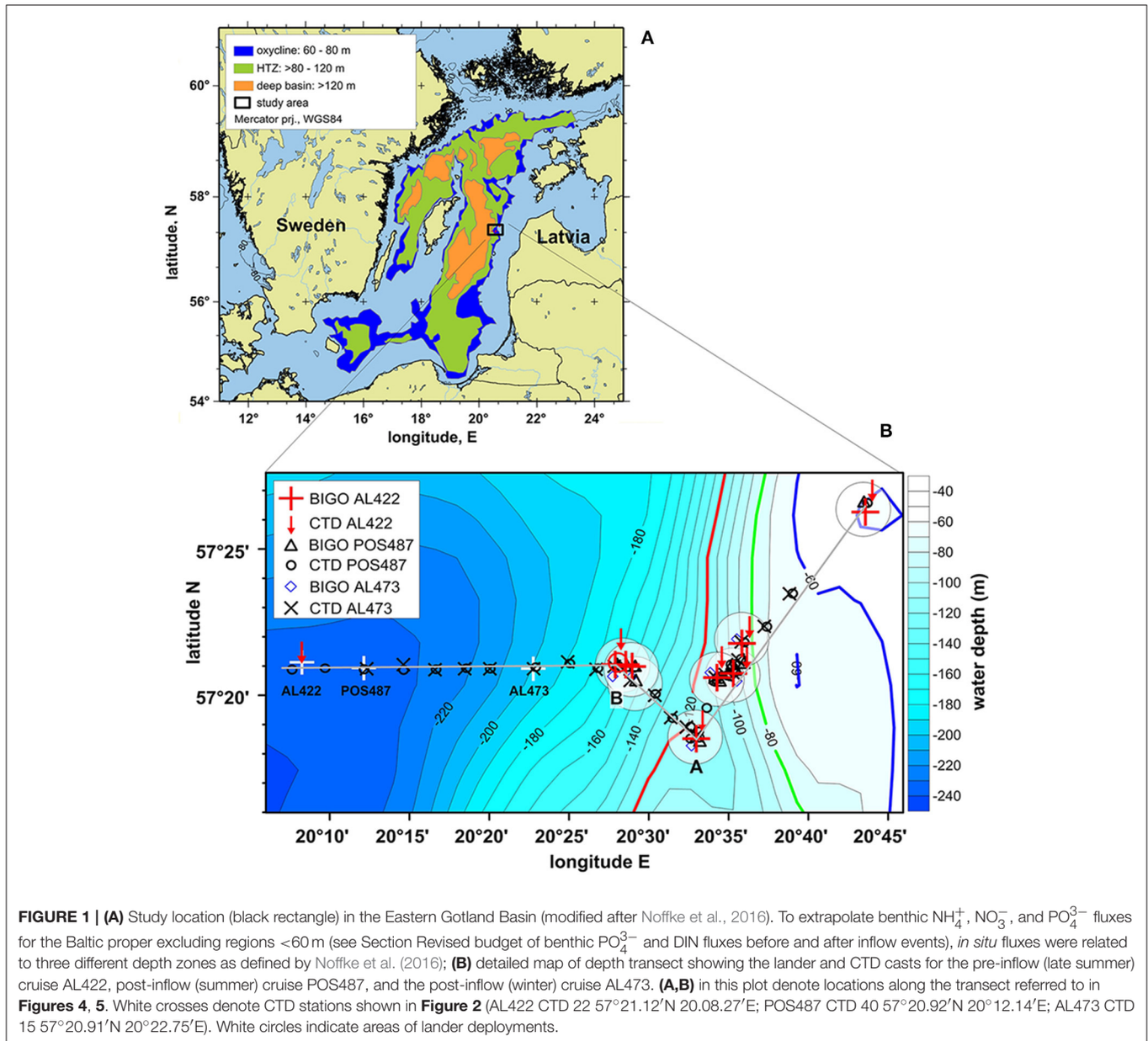
higher than the external P load of 14 kt year⁻¹ reported for 2006 (HELCOM, 2009b). Hence, one of the main motivations for this study was to determine whether rapid P and N cycling at the seafloor is strongly altered during MBIs. Comparison of in situ benthic flux measurements in the pre- and post-inflow phases both made in late summer revealed markedly different dynamics in the deep basin, but little change in the HTZ where most P and N regeneration takes place.

METHODS

Field Campaigns

Sampling campaigns in the EGB were conducted on the RV Alkor cruise AL422 in August/September 2013, RV Poseidon cruise POS487 in July/August 2015 and RV Alkor cruise AL473 in March 2016 (Table 1; Figure 1). A previous cruise on Alkor

AL355 taking place under euxinic conditions of the deep basin in May/June 2010 has been described by Noffke et al. (2016); results from that study are not re-tabulated here. In subsequent sections this cruise will be referred to as pre-inflow (early summer) cruise. Data from AL422 are representative of late summer stagnant, euxinic conditions in the deep basin at a time of cyanobacterial bloom development referred to here as pre-inflow (late summer) cruise. Data from POS487 is referred to as post-inflow (summer) cruise and were taken at the same time of year under ventilated deep basin conditions. Hence, comparison of data from these cruises should allow the effect of the MBI on sediment fluxes to be addressed directly without being too confounded by seasonality, although some degree of inter-annual variation in primary production and benthic respiration is to be expected. Data from winter 2016 [cruise AL473, referred to as post-inflow (winter) cruise] are presented here to show the



evolution of benthic fluxes under prolonged ventilation, but are not the focus of the present study. This cruise took place in March 2016 and captured the moderate MBI that was triggered in November 2015.

For this study we will adopt the definitions suggested by Noffke et al. (2016) based on water column O_2 profiles in the Baltic Proper under stagnant conditions. The depth range 60–80 m was identified as the oxycline where O_2 in the bottom water is mostly present although highly variable. The range > 80–120 m was characterized as the HTZ with O_2 levels typically <30 μM but which can be variable. The zone below 120 m and a well-expressed redoxcline defines the deep anoxic and sulfidic basin. During MBIs, this zone becomes oxygenated (**Table 1**).

Conductivity, temperature, depth (CTD) measurements were performed during casts of a Seabird CTD system equipped with a water-sampling rosette. These casts were made at water depths between 50 and 223 m along the redox depth transect shown in **Table 1** and **Figure 1B**. Immediately after retrieval, water samples from Niskin bottles were analyzed for the nitrogen species nitrate (NO_3^-), nitrite (NO_2^-), and NH_4^+ . Phosphate and hydrogen sulfide measurements include all dissolved species. Their ionic forms PO_4^{3-} and HS^- are referred to in this study.

In situ fluxes were determined at eight sites during the pre-inflow (late summer) cruise AL422 (**Table 1**, **Figure 1**) covering the entire oxic to anoxic/sulfidic gradient. Flux measurements were repeated at 110 and 123/124 m, giving a total of 11 deployments. The same sites were investigated on the post-inflow

(summer) cruise POS487, with a further site at 161 m to achieve a better resolution during this ventilation period. During the post-inflow (winter) cruise AL473, *in situ* measurements were conducted at six sites along the transect.

Sea Floor Observation

Sea floor images were obtained using the towed camera system OFOS (Ocean Floor Observation System) equipped with a video and still camera (Nikon D70s), two Xenon lights (Oktopus) and a flashlight (Benthos). The system was towed ~ 1.5 m above the sea floor at ~ 0.3 knots. Seven OFOS deployments were conducted during the pre-inflow late summer cruise along the depth transect where *in situ* fluxes were determined (**Figure 1B**). Additionally, 11 OFOS deployments were conducted to the north and south of the depth transect to confirm presence of microbial mats in the HTZ as described by Noffke et al. (2016). During the post-inflow (summer) cruise, OFOS deployments completely surveyed the depth gradient where *in situ* fluxes were performed covering a distance of about 27 km of continuous video surveillance, spanning water depths from 70 to 240 m. A second depth transect was investigated to the north of the main working area covering a distance of ~ 18 km and a depth range of ~ 63 –195 m (not shown). During the post-inflow (winter) cruise OFOS deployments were conducted at the depths where flux measurements were made.

In situ Flux Measurements and Sediment Sampling

In situ fluxes were determined using benthic chambers mounted in two Biogeochemical Observatories (BIGO) (Sommer et al., 2009; Pfannkuche and Linke, 2003). Each BIGO contains two chambers. The water volume enclosed by the chambers ranged from ca. 9 to 20 L and incubation times ranged from 29 to 56 h. Solute fluxes in each chamber were determined from concentrations measured in discrete water samples removed periodically with eight glass syringes (each ~ 46 mL). Immediately after retrieval of the observatories, the water samples were stored (max. 4 h) in the onboard cold room (about 6°C) before geochemical analyses. Furthermore, chamber water was sampled into 5 slender glass tubes (each ~ 15 mL) for onboard determination of DIC. TOU was determined from *in situ* O₂ concentration measurements in the chambers using optical sensors (Aanderaa Instruments, Norway; Model 3830, Tengberg et al., 2006). Optode performance was cross-checked with O₂ concentrations determined in the water samples by Winkler titration. Solute fluxes were calculated from the linear increase or decrease of concentration versus time, corrected for the surface area to volume ratio of each chamber.

The landers are also equipped to recover the upper incubated sediment layers (~ 10 –15 cm), which serves as a check for sediment disruption during seafloor operations and chamber insertion. The sediment surface for all lander deployments during the pre-inflow (late summer) cruise was intact and undisturbed. On post-inflow cruises the sediments close to the chamber wall of BIGO-II-3 at the 178 m site and of BIGO-I-3 at 151 m were slightly disturbed. It is not known whether this was caused during insertion of the chambers into the sediment or during lander

retrieval. The concentration data did not indicate any artifacts during the flux measurements, and the initial concentrations inside the chamber at the start of the incubation were close to the bottom-water concentrations. Consequently, we have no reason to disregard the fluxes determined from these deployments.

Geochemical Measurements

Concentration measurements of dissolved inorganic nitrogen (NO₃⁻, NO₂⁻, NH₄⁺), PO₄³⁻ and HS⁻ were performed on board. Nutrients (NO₃⁻, NO₂⁻, PO₄³⁻) were determined on a QuAAtro autoanalyzer (Seal Analytical) using standard photometrical methods (Grasshoff et al., 1999). NH₄⁺ and HS⁻ were measured using photometry with a Hitachi U2800 photometer. Analytical details are described by Sommer et al. (2016). NO₂⁻ fluxes were always <10% of NO₃⁻ fluxes, and here we report NO₃⁻ fluxes as $\Sigma\text{NO}_3^- + \text{NO}_2^-$. Dissolved O₂ in Niskin bottles was determined using automated Winkler titration with a detection limit of 3 $\mu\text{mol L}^{-1}$. Concentrations below the detection limit were considered to be anoxic for purposes of defining the prevailing bottom water O₂ regime.

DIC measurements were performed using a quadrupole membrane inlet mass spectrometer (MIMS, GAM200, In Process Instruments). The instrument was equipped with inline sample acidification to shift the carbonate system entirely to the volatile CO₂ species, which then was measured on the MIMS at a mass to charge ratio of 44 (Bell et al., 2011). The general setup is described in detail in Noffke et al. (2016). In short, the water samples taken during the BIGO deployments were pumped using a peristaltic pump (Ismatec, REGLO Digital MS-4/6) at 1.0 mL min⁻¹ through a membrane inlet (Sommer et al., 2015), where extraction of dissolved gases takes place. Gas flow from the inlet to the mass spectrometer was conducted in a steel capillary supported with helium that was supplied through a fused silica capillary. The distance between the inlet and the ion source of the quadrupole was about 80 cm. An in-line cryo-trap (-35°C , ethanol) between the inlet and the mass spectrometer was used to reduce water vapor. The water samples were first analyzed without acidification and recovered in Labco Exetainers capped without a headspace. Subsequently, an acidification inlet was integrated into the sample flow in front of the membrane inlet and 4 M hydrochloric acid added at a rate of 0.15 mL h⁻¹ using a precision syringe pump (KDS-100-CE, kd Scientific) resulting in a sample-acid ratio of 402:1 transforming all DIC into CO₂. The DIC measurements were calibrated with a Na₂CO₃ standard (Merck) that was diluted with a sodium chloride solution (10 g L⁻¹) to standards in the range of 1.50–2.25 mM. The standards were prepared at the respective *in-situ* temperature and were measured before and after each measurement session.

RESULTS

Water Column Pre-inflow Conditions

The O₂ concentrations in the pre-inflow phase (late summer) were similar to those previously described by Noffke et al. (2016) (**Figure 2**). Below the oxycline (70–80 m) that coincided with the pycnocline, O₂ concentrations were <30 μM down to ca.

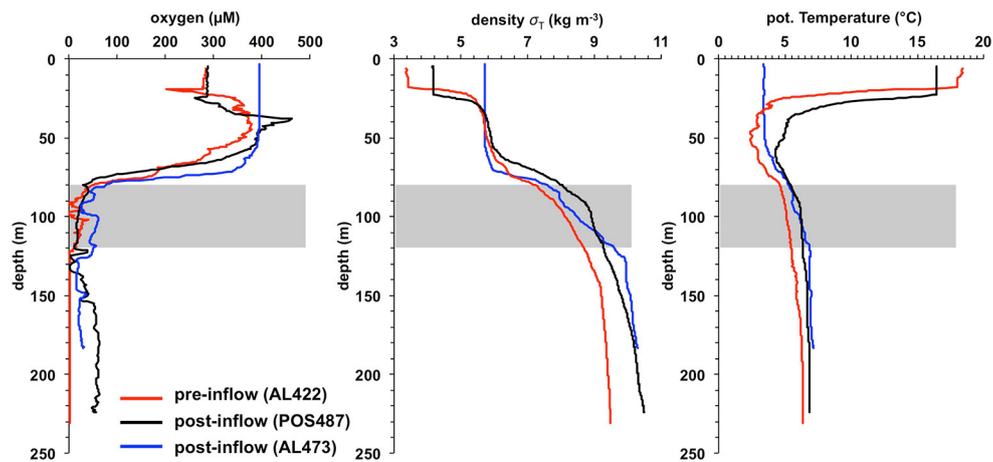


FIGURE 2 | Water-column O_2 concentration, density (σ_T) and potential temperature profiles for the pre- (AL422) and post-inflow conditions in summer (POS487) and winter (AL473). The gray bars indicate the HTZ. Station positions are indicated in Figure 1B.

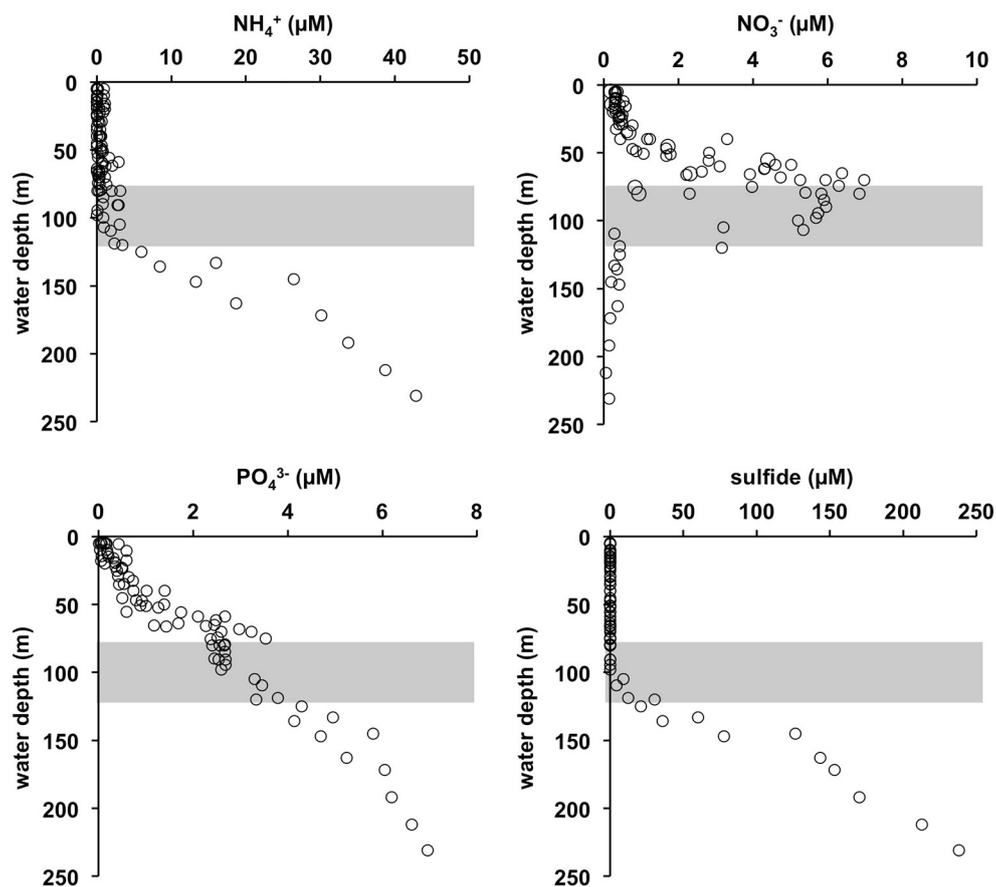
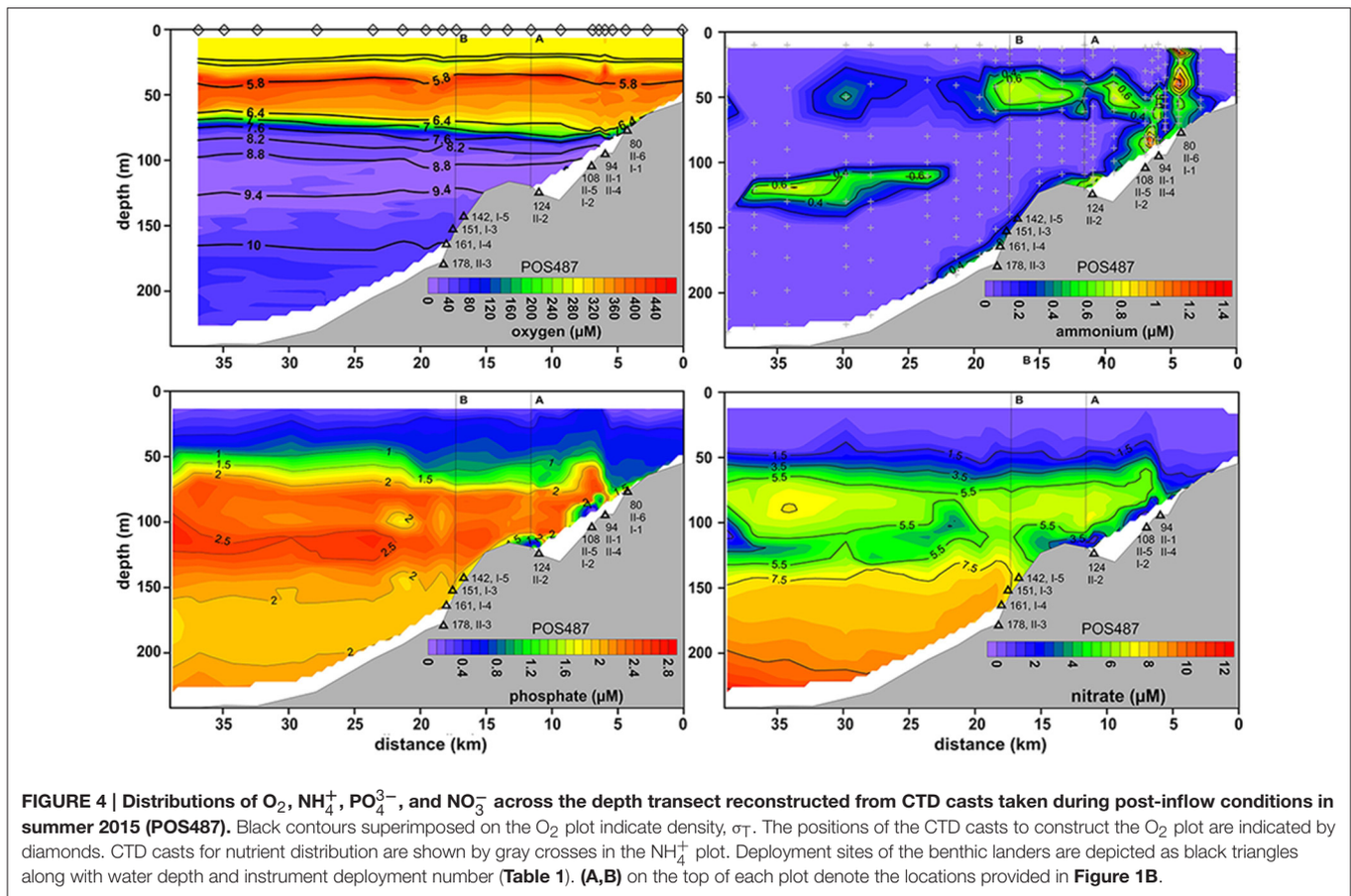


FIGURE 3 | Water column concentration profiles of NH_4^+ , NO_3^- , PO_4^{3-} and sulfide compiled during several CTD casts during pre-inflow cruise AL422 (for positions see Figure 1B). The gray shaded area approximates the extent of the HTZ.

120 m. This depth interval marks the HTZ. In the deep basin below the HTZ, O_2 was undetectable. The deep basin was highly enriched in HS^- (Figure 3), reaching levels of 238 μM ; much

higher than the maximum value of 150 μM reported for euxinic conditions in May/June 2010 (Noffke et al., 2016). HS^- clearly diffuses upwards to the base of the HTZ whereupon it is oxidized.



NH_4^+ accumulated to $43 \mu M$ in the deep basin, again higher than in 2010 where concentrations reached $25 \mu M$ (Noffke et al., 2016). At the base of the HTZ, the NH_4^+ levels were much lower and remained $<3 \mu M$ within the HTZ but were depleted toward the oxycline. NO_3^- showed a peak of $7 \mu mol L^{-1}$ at 70–80 m, but was depleted toward the surface and below ~ 120 m water depth. PO_4^{3-} concentrations were highest in the deep basin ($7 \mu M$) and declined toward the mixed surface layer, illustrating that the deep anoxic basin is a source of PO_4^{3-} to surface waters. As already shown for pre-inflow conditions in early summer 2010 (Noffke et al., 2016), PO_4^{3-} displayed a pronounced gradient change at about 100 m water depth with elevated concentrations ($\sim 2.5 \mu M$) between 60 m and 100 m, indicating an additional source in the HTZ.

Post-inflow Conditions

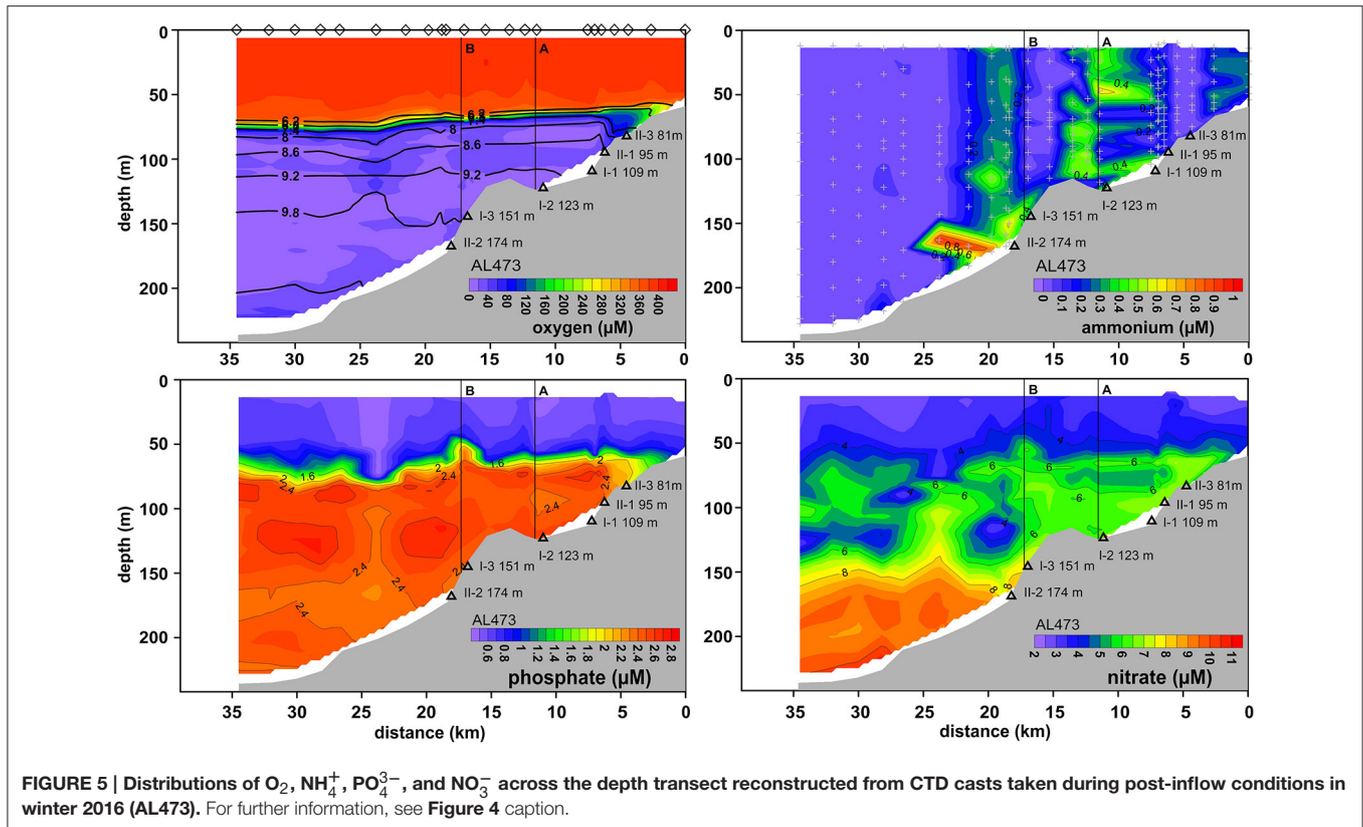
The strong MBI triggered in 2014 oxygenated the deep basin, leading to dissolved O_2 concentrations of up to $\sim 70 \mu M$ at 173 m water depth (Figures 2, 4). In deeper parts of the basin the O_2 levels decreased to about $40 \mu M$. Unexpectedly, the O_2 levels remained below $30 \mu M$ within the HTZ, and the 111 m site was characterized with lowest O_2 concentrations. The oxycline was located at around 60–80 m water depth; similar to the pre-inflow phase. The O_2 level in the surface layer was $90 \mu M$ lower than measured during pre-inflow conditions (early summer)

(Noffke et al., 2016) yet similar to the level measured during pre-inflow conditions (late summer) indicating a seasonal effect of O_2 consumption and temperature effects on O_2 solubility. HS^- was not detectable in the post-inflow water column.

The MBI caused a massive perturbation to nutrient distributions. NO_3^- , which serves as an important electron acceptor during anaerobic respiration, increased with water depth from the base of the HTZ to $12 \mu M$ in the deep basin (Figure 4). As for O_2 , the 110–120 m site appeared to be the station with the lowest availability of electron acceptors with a marked NO_3^- minimum. Above this depth in the HTZ, NO_3^- reached maximum levels of about $7 \mu M$; the same as measured during euxinic conditions (pre-inflow early and late summer; Noffke et al., 2016 and this study).

NH_4^+ did not accumulate in the deep basin post-inflow and remained below $1 \mu M$ (Figure 4). Only at the benthic boundary were NH_4^+ levels slightly increased. However, increased NH_4^+ concentrations were measured in a distinct layer in the water column at about 125 m water depth at the lower edge of the HTZ. Similarly, PO_4^{3-} concentrations in the deep basin were ca. three-fold lower compared to those measured during euxinic conditions. Again, within the HTZ, concentrations were similar to those measured during euxinic conditions ($\sim 2.4 \mu M$).

As a result of the moderate MBI that was triggered in November 2015, oxygenated water masses were detected in the



deep basin although with lower O_2 levels compared to the post-inflow (summer) cruise (Figures 2, 5). The surface layer was well mixed with a steep oxycline between 70 and 80 m resulting in an enhanced O_2 penetration compared to the pre-inflow (summer) and post-inflow (summer) situation. As during all previous cruises, the HTZ was retained yet with slightly elevated O_2 levels. Deep-water renewal maintained elevated levels of NO_3^- , whereas accumulation of NH_4^+ and PO_4^{3-} or HS^- was impeded.

Seafloor Observations

Seafloor imaging during all cruises showed that the sediment surface in the HTZ was densely covered with white filamentous microbial mats, tentatively identified as belonging to the family *Beggiatoaceae*. Also the stations to the south and north of the main working area showed occurrences of microbial mats confirming earlier observations made during cruises R/V Poseidon (POS369 July–Aug. 2008), R/V Alkor (AL346 Sept.–Oct. 2009) as well as R/V Alkor cruise AL355 (Noffke et al., 2016). Notably, under euxinic conditions only sporadic occurrence of microbial mats was observed below 120 m water depth and they became absent with increasing depth. In contrast, during ventilated conditions the microbial mats extended down to water depths of 240 m and extensively covered the sediment surface. Mats were dominated by different size-classes of filaments, with smaller filaments of 4–5 μm in diameter being more abundant at oxic stations and larger filaments with diameters of 22–32 μm

dominating at stations where bottom water O_2 concentrations approached zero.

In situ Fluxes

Pre-inflow Conditions

NH_4^+ fluxes showed a distinct maximum of ca. 1.5 $\text{mmol m}^{-2} \text{d}^{-1}$ in the HTZ between 100 and 120 m (Table 2; Figure 6), with a second maximum of 1.1 $\text{mmol m}^{-2} \text{d}^{-1}$ in the deep basin (173 m). NO_3^- was taken up by the sediment at all sites in the oxycline and HTZ with a maximum of $-0.72 \text{ mmol m}^{-2} \text{d}^{-1}$ at 96 m and 110 m in the HTZ. NO_3^- fluxes were zero at the deep sulfidic stations due to the absence of NO_3^- in the bottom water. Overall, the sediments were a source of DIN with elevated release at the oxycline, inside the HTZ, and at the transition between the HTZ and the deep basin (124 m) (Table 2). As for NH_4^+ , pre-inflow PO_4^{3-} fluxes were directed out of the sediment and elevated inside the HTZ (0.25 $\text{mmol m}^{-2} \text{d}^{-1}$). Fluxes decreased at the lower boundary of the HTZ but increased again in the deep basin to 0.21 $\text{mmol m}^{-2} \text{d}^{-1}$. Below the HTZ, HS^- fluxes increased and reached a maximum value of 10.2 $\text{mmol m}^{-2} \text{d}^{-1}$ at 173 m (Table 2). In general, the above trends are within the uncertainty of the flux measurements made during the pre-inflow (summer) cruise in 2010 by Noffke et al. (2016). The exception is the 80 m site where PO_4^{3-} fluxes were lower in that study and directed into the sediment. DIC fluxes decreased from the oxycline (10.3 $\text{mmol m}^{-2} \text{d}^{-1}$) to a minimum at 140 m (3.4 $\text{mmol m}^{-2} \text{d}^{-1}$) coinciding with a minimum of NH_4^+ and PO_4^{3-} fluxes. TOU of

TABLE 2 | Benthic fluxes in the EGB measured using benthic landers (mmol m⁻² d⁻¹).

Deployment		Depth (m)	Incubation time (h)	NH ₄ ⁺	NO ₃ ⁻	DIN	PO ₄ ³⁻	HS ⁻	DIC	TOU
EUXINIC PRE-INFLOW CONDITIONS (AL422 AUGUST/SEPTEMBER 2013)										
BIGO-II-6	CH1	65	30.0	1.0	-0.22	0.78	0.04	bdl	nd	-12.6
	CH2			1.0	-0.21	0.79	0.06	bdl	nd	-14.6
BIGO-I-2	CH1	80	31.5	0.8	-0.69	0.11	0.22	bdl	nd	-0.37
	CH2			0.3	-0.50	-0.20	0.26	bdl	nd	-0.44
BIGO-II-1	CH1	96	30.0	1.0	-0.52	0.48	0.08	bdl	nd	bdl
	CH2			1.3	-0.72	0.58	0.19	bdl	nd	bdl
BIGO-II-3	CH1	110	31.0	1.1	-0.72	0.38	0.13	bdl	15.3	bdl
	CH2			1.5	-0.72	0.78	0.20	bdl	16.1	bdl
BIGO-I-6	CH1	110	29.0	nd	nd	nd	nd	nd	nd	nd
	CH2			1.3	-0.13	1.17	0.27	bdl	nd	bdl
BIGO-I-5	CH1	123	30.0	nd	nd	nd	nd	nd	nd	nd
	CH2			1.2	-0.35	0.85	0.21	bdl	3.6	bdl
BIGO-I-4	CH1 ^a	123	30.0	nd	nd	nd	nd	nd	nd	nd
	CH2			1.5	-0.84	0.66	0.27	bdl	nd	bdl
BIGO-I-1	CH1	124	30.0	1.4	bdl	1.40	0.24	bdl	nd	bdl
	CH2			1.6	bdl	1.60	0.19	bdl	nd	bdl
BIGO-II-5	CH1	140	29.0	0.7	bdl	0.7	0.09	7.61	nd	bdl
	CH2			0.1	bdl	0.1	0.00	5.27	3.6	bdl
BIGO-I-3	CH1	152	35.0	0.5	bdl	0.5	0.1	3.24	13.2	bdl
	CH2			0.6	bdl	0.6	0.1	4.11	10.9	bdl
BIGO-II-4	CH1	173	31.0	0.9	bdl	0.9	0.19	10.15	7.0	bdl
	CH2			1.3	bdl	1.3	0.22	9.81	2.9	bdl
VENTILATED POST-INFLOW CONDITIONS (POS487 JULY/AUGUST 2015)										
BIGO-I-6	CH1	63	32.0	0.07	0.13	0.20	-0.03	bdl	6.7	8.3
	CH2			0.27	0.24	0.51	0.02	bdl	5.2	8.8
BIGO-II-6	CH1	80	32.0	0.57	-0.57	2.35	0.23	bdl	6.2	2.4
	CH2			0.76	-0.56	2.35	0.30	bdl	5.8	2.1
BIGO-II-1	CH1	94	30.6	1.40	-1.69	-0.29	0.38	bdl	nd	nd
	CH2 ^a			nd	nd	nd	nd	nd	nd	nd
BIGO-II-4	CH1	95	35.0	1.63	-0.64	0.99	0.20	bdl	11.8	-2.8
	CH2			1.13	-0.48	0.65	0.16	bdl	11.5	-3.0
BIGO-I-2	CH1	111	36.1	0.59	-0.62	-0.03	0.11	bdl	6.4	-1.0
	CH2 ^b			0.50	-0.50	0.00	0.05	bdl	nd	nd
BIGO-II-5	CH1	111	34.0	0.52	-0.51	0.01	0.07	bdl	4.2	-0.7
	CH2 ^b			0.72	-0.78	-0.06	0.08	bdl	3.7	-1.0
BIGO-II-2	CH1	124	36.0	1.14	-0.99	0.15	0.21	bdl	15.7	0.8
	CH2 ^a			nd	nd	nd	nd	nd	nd	nd
BIGO-I-5	CH1	142	35.0	0.77	-0.81	-0.04	-0.01	bdl	bdl	2.9
	CH2 ^b			0.80	-0.77	0.03	-0.08	bdl	nd	nd
BIGO-I-3	CH1	151	56.0	0.33	-0.41	-0.08	-0.02	bdl	4.0	5.2
	CH2			0.20	-0.30	-0.10	-0.03	bdl	2.2	3.5
BIGO-I-4	CH1	161	36.0	0.41	-0.43	-0.02	-0.07	bdl	4.1	4.3
	CH2			0.40	-0.38	0.02	-0.04	bdl	2.8	3.2
BIGO-II-3	CH1	178	30.0	0.87	-0.54	0.33	0.19	bdl	9.9	3.8
	CH2			0.78	-0.59	0.19	0.15	bdl	6.8	3.8

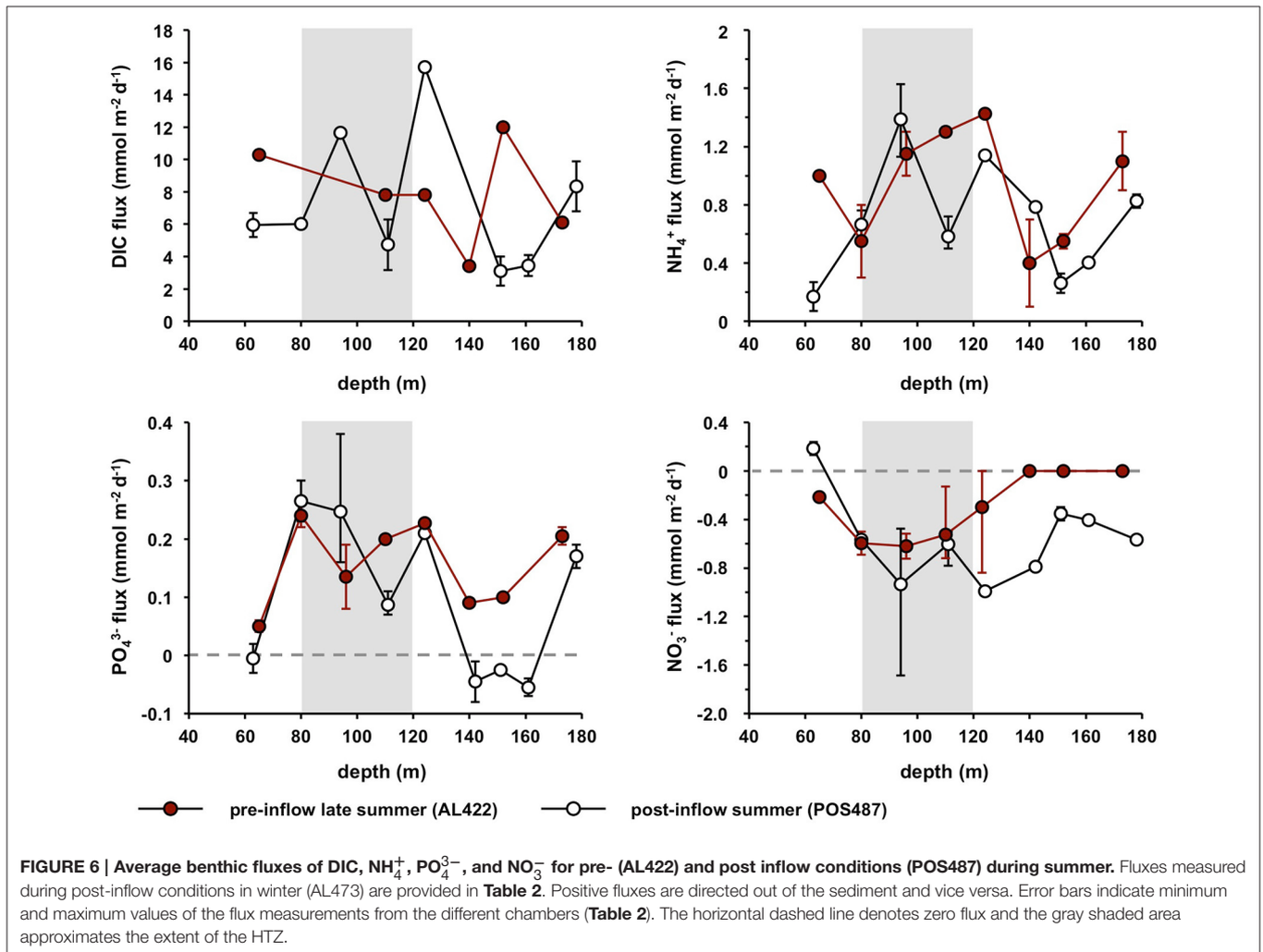
(Continued)

TABLE 2 | Continued

Deployment	Depth (m)	Incubation time (h)	NH_4^+	NO_3^-	DIN	PO_4^{3-}	HS^-	DIC	TOU	
VENTILATED POST-INFLOW CONDITIONS (AL473 MARCH 2016)										
BIGO-II-3	CH1	81	30.7	0.29	-0.03	0.59	-0.07	bdl	nd	-4.5
	CH2			0.36	-0.49	0.72	-0.07	bdl	nd	-4.3
BIGO-II-1	CH1	95	31.7	1.65	-0.82	0.83	0.19	bdl	16.4	-3.5
	CH2			1.29	-0.64	0.65	0.16	bdl	15.1	-2.9
BIGO-I-1	CH1	109	29.7	1.48	-0.74	0.74	0.17	bdl	nd	-1.4
	CH2			1.31	-0.65	0.66	0.16	bdl	nd	-2.1
BIGO-I-2	CH1	123	33.7	1.23	-0.55	0.68	0.14	bdl	15.1	-2.5
	CH2			1.08	-0.73	0.34	0.17	bdl	8.2	-2.2
BIGO-I-3	CH1	151	30.7	0.44	-0.34	0.10	0.04	bdl	5.1	-1.4
	CH2			0.62	-0.42	0.20	-0.001	bdl	5.2	-1.4
BIGO-II-2	CH1	174	30.7	2.14	-0.88	1.26	0.20	bdl	13.9	-2.4
	CH2			2.42	-0.75	1.71	0.26	bdl	13.2	-3.0

^aCH1 failed; ^bChamber volume was assumed to be the same as that of CH1.

Fluxes for both chambers (CH1, CH2) of each lander deployment are provided. Positive fluxes are directed out of the sediment. $\text{DIN} = \text{NH}_4^+ + \text{NO}_3^-$. nd, no data, bdl, below detection limit. O_2 uptake is referred to as total oxygen uptake (TOU). DIC is considered equivalent to organic carbon degradation.



up to $14.6 \text{ mmol m}^{-2} \text{ d}^{-1}$ was only measurable at the oxycline station and at the upper boundary of the HTZ ($0.44 \text{ mmol m}^{-2} \text{ d}^{-1}$) (Table 2).

Post-inflow Conditions

During both post-inflow cruises, ventilation resulted in elevated TOU rates in the deep basin ranging from 0.8 to $5.2 \text{ mmol m}^{-2} \text{ d}^{-1}$ (post-inflow, summer) and from 1.4 to $3.0 \text{ mmol m}^{-2} \text{ d}^{-1}$ (post-inflow, winter) (Table 2). Within the HTZ, TOU was now higher at the 80 m site with rates of 2.4 and $4.5 \text{ mmol m}^{-2} \text{ d}^{-1}$ during cruises POS487 (summer) and AL473 (winter), respectively. During both post-inflow cruises, DIC fluxes were elevated at the deepest station with maximum rates of 9.9 and $13.9 \text{ mmol m}^{-2} \text{ d}^{-1}$ as well as at the upper boundary of the deep basin at 123 m water depth (15.7 and $15.1 \text{ mmol m}^{-2} \text{ d}^{-1}$) (Figure 6, Table 2).

Despite oxygenated bottom waters, the depth distribution of NH_4^+ fluxes during the post-inflow (summer) cruise was similar as described above for the pre-inflow situation (Figure 6, Table 2). Maximum average NH_4^+ release was $1.4 \text{ mmol m}^{-2} \text{ d}^{-1}$ at 96 m inside the HTZ with a second pronounced peak of $0.8 \text{ mmol m}^{-2} \text{ d}^{-1}$ at the deepest station. Fluxes at 110 m were lower than measured previously ($0.6 \text{ mmol m}^{-2} \text{ d}^{-1}$) but had increased again to high values ($1.4 \text{ mmol m}^{-2} \text{ d}^{-1}$) in the following winter (cruise AL473, Table 2). In contrast to the late summer pre-inflow scenario, NO_3^- was slightly released at the oxycline with an average rate of $0.2 \text{ mmol m}^{-2} \text{ d}^{-1}$. Due to the presence of NO_3^- in the bottom water, NO_3^- was taken up by sediments between 80 and 173 m at average rates between $-0.4 \text{ mmol m}^{-2} \text{ d}^{-1}$ at 151 m and $-1.0 \text{ mmol m}^{-2} \text{ d}^{-1}$ at 124 m. During winter (AL473), NO_3^- was taken up at all sites at rates similar to the summer post-inflow condition (Table 2). PO_4^{3-} fluxes in the HTZ measured during pre- and post-inflow summer were similar, with elevated release rates in the range of 0.09 – $0.27 \text{ mmol m}^{-2} \text{ d}^{-1}$ (Figure 6). However, during summer post-inflow conditions, PO_4^{3-} was taken up by the sediments at the oxycline (65 m) and at the now oxic deep-water stations (142, 151, and 161 m) with a maximum uptake rate of $0.06 \text{ mmol m}^{-2} \text{ d}^{-1}$. PO_4^{3-} was still being released to bottom waters at the deepest site at $0.17 \text{ mmol m}^{-2} \text{ d}^{-1}$ despite ventilated bottom water conditions. In the following winter, PO_4^{3-} was released from all sites at rates similar to the pre-inflow conditions except for the 80 m site (Table 2). Benthic HS^- release was apparently efficiently diminished to below detection limit under oxic conditions.

DISCUSSION

The objective of this study is to first identify changes in benthic nutrient fluxes in response to two MBIs. These inflows led to increased availability of O_2 and NO_3^- in the deep basin, and lowered NH_4^+ and PO_4^{3-} (this study, Mohrholz et al., 2015; Nausch et al., 2016). Benthic fluxes can then be extrapolated for the Baltic Proper in order to assess MBIs with regard to their potential to mitigate eutrophication in this area.

Differential Response to Ventilation in Deep Basin and HTZ

As known from previous MBIs in 1993 and 2003 (e.g., Nausch and Nehring, 1994), the ventilation in 2015/2016 profoundly changed the redox landscape of the water column below ca. 125 m. Deep-water renewal resulted in similar low NH_4^+ and PO_4^{3-} concentrations to those observed after the inflow event in 1993 (Nausch and Nehring, 1994) as well as elevated NO_3^- . The water column HS^- inventory was completely oxidized and not detectable, in stark contrast to euxinic conditions ($240 \mu\text{M}$, this study). This entails very efficient HS^- oxidation mechanisms in the water column as well as at the sediment surface. Oxidation of reduced compounds likely becomes enhanced in the water column following ventilation due to bottom boundary mixing. Furthermore, during propagation of dense bottom currents, entrainment of less saline water also leads to interleaving of incoming oxygenated water masses with the ambient water mass at the level of neutral buoyancy and concomitant ventilation at different depths (Reissmann et al., 2009, cf. their Figure 2). In the sediments, sulfate reduction below the oxic sediment surface still produces high amounts of HS^- whose oxidation represents a strong sink for O_2 . Mohrholz et al. (2015) estimate that the total amount of O_2 transported into the Baltic Sea during 2015 amounts to $\sim 2.04 \times 10^6 \text{ t}$. Given a benthic HS^- flux of $6.7 \text{ mmol m}^{-2} \text{ d}^{-1}$ during pre-inflow (Table 2), and a deep basin area of $\sim 19,000 \text{ km}^2$ (Table 3), one can estimate that the sediments alone could consume this flux within 8 months. This is a maximum lifetime of O_2 in the basin with respect to benthic respiration in the deep basin since the O_2 inventory in the Gotland Basin will be lower than the total inflow into the Baltic (Schmidt, 2014) and a fraction of O_2 will be consumed by respiration of reduced dissolved and particulate compounds in the water column. Indeed, a fast decrease of O_2 levels was recorded in September and December 2015 during time-series observations conducted at 2 and 20 m above the seafloor, respectively (Mohrholz et al., 2016). This calculation, whilst necessarily first-order, nonetheless emphasizes that HS^- oxidation at the sediment water interface strongly contributes to the rapid depletion in O_2 following deep basin ventilation.

In contrast to elevated P release under euxinic bottom water conditions (this study; Jilbert et al., 2011; Viktorsson et al., 2013a; Noffke et al., 2016; and references therein), PO_4^{3-} was taken up by the sediment at the sites at 142–161 m water depth under ventilated conditions during July/August 2015. A significant removal pathway for phosphorus occurs by way of its adsorption onto iron oxides while $\text{pH} < 9$ (Berner, 1973; Van Cappellen and Ingall, 1996), and reactive iron might become increasingly available in surface sediments under oxic conditions. Remarkably, though, despite oxic conditions during both inflow cruises POS487 (summer) and AL473 (winter) at the deepest sites, average PO_4^{3-} release (0.17 and $0.23 \text{ mmol m}^{-2} \text{ d}^{-1}$ respectively) was similar to respective fluxes during euxinic conditions in July/August ($0.21 \text{ mmol m}^{-2} \text{ d}^{-1}$, this study) or even higher compared to May/June ($0.15 \text{ mmol m}^{-2} \text{ d}^{-1}$, Noffke et al., 2016). This might be related to the time course of the inflow spreading into the Gotland Basin. O_2 -rich intrusions

TABLE 3 | Regionalization of P and N fluxes (ktons year⁻¹) based on average local PO₄³⁻, NH₄⁺, and NO₃⁻ fluxes measured in the different depth zones in the Baltic Proper.

Depth zone	Pre-inflow, euxinic deep basin		Post-inflow, ventilated deep basin	
	AL355	AL422	POS487	AL473
PHOSPHATE				
Oxycline (60–<80 m)	11.5 ± 19.9	14.8 ± 4.2	–1.5 ± 10.4	nd
HTZ (80–120 m)	66.4 ± 78.5	102.4 ± 28.3	106.3 ± 52.5	48.1 ± 74.1
Deep basin (>120 m)	21.4 ± 9.3	32.3 ± 13.8	21.6 ± 13.7	31.4 ± 5.7
Grand total*	87.9 ± 43.9	134.8 ± 21.1	128.0 ± 33.1	79.5 ± 39.9
AMMONIUM				
Oxycline	31.0 ± 10.7	133.3 ± 0	22.6 ± 18.6	nd
HTZ	191.9 ± 83.9	241.3 ± 95.8	211.7 ± 106.9	256.6 ± 154.6
Deep basin	34.9 ± 19.1	81.6 ± 42.4	73.9 ± 17	93.2 ± 21.5
Grand total*	226.7 ± 51.5	322.9 ± 69.1	285.5 ± 62	349.9 ± 88.1
NITRATE				
Oxycline	30.0 ± 13.5	–28.7 ± 0.9	24.7 ± 10.4	nd
HTZ	–178.6 ± 101.6	–139.6 ± 12.6	–169.4 ± 49.3	–135.6 ± 63.2
Deep basin	–12.6 ± 21.8	–5.8 ± 11.5	–32.9 ± 12.9	–31.9 ± 5.1
Grand total*	–191.2 ± 61.7	–145.3 ± 12.0	–202.3 ± 31.1	–167.5 ± 34.2

Area of the oxycline is 26,088 km², the HTZ is 47,230 km² and the deep basin is 18,954 km²; see also **Figure 1**. Fluxes are shown for euxinic, stagnant conditions before (AL422 this study; AL355 Noffke et al., 2016) as well as oxygenated conditions after the strong ventilation event in December 2014 (POS487) and after a moderate ventilation event in November 2015 (AL473).

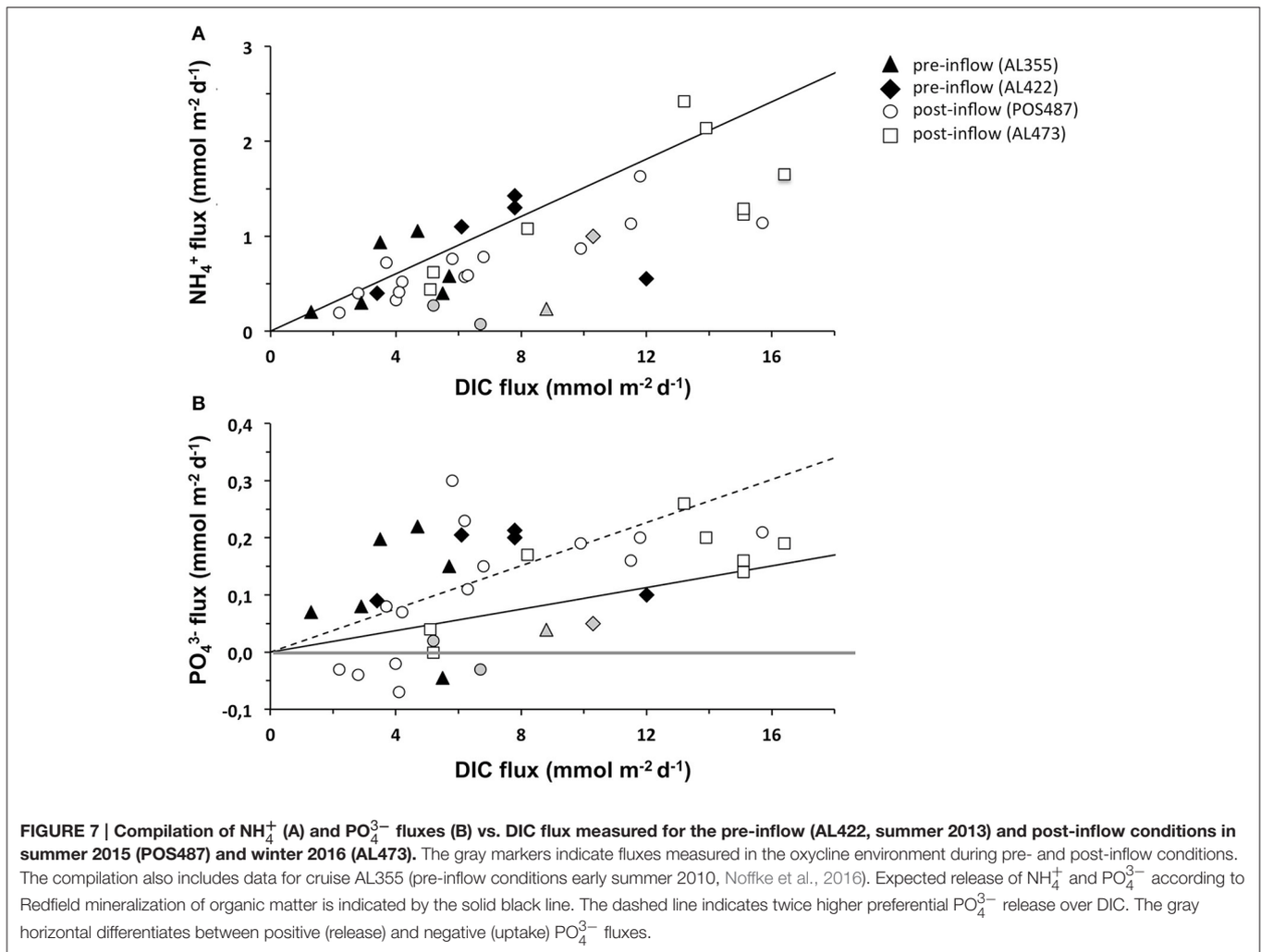
*Excludes the oxycline.

of North Sea waters propagate as dense bottom currents, such that the deepest sites were likely ventilated first followed by ventilation of the shallower sites sometime later as the deep water is progressively replaced. Hence, reactive iron for P sequestration might already have become limited at the deeper sites since organic P degradation rates are relatively elevated here resulting in a low P retention capacity of the sediments (Noffke et al., 2016). During both post-inflow cruises, PO₄³⁻ release from the deepest station resulted in the gradual accumulation of PO₄³⁻ in the water column as observed in the deepest part of the EGB (compare **Figures 4, 5**).

At the upper boundary of the deep basin at 124 m water depth, the PO₄³⁻ release during both post inflow cruises in summer and winter was as high or almost as high as under pre-inflow euxinia. An O₂ time series recorded at this site over the entire cruise POS487 revealed strong bottom water O₂ fluctuations between 0 and 15 μM and extended periods of O₂ levels <5 μM (not shown). The O₂ profiles and NO₃⁻ distributions (**Figures 2, 4**) suggest that the bottom water at this site has the lowest availability of electron acceptors following ventilation. Hence, this site can be considered as strongly hypoxic, thereby providing a partial explanation for the ongoing high PO₄³⁻ release rates. These O₂ fluctuations are likely caused by internal waves and seiches (Reissmann et al., 2009), which drive water mass movement across the 124 m site resulting in the periodic entrainment of colder anoxic water from above (HTZ) as well as warmer North Sea water rich in O₂ and NO₃⁻ from below.

Seafloor imaging revealed that sulfur bacteria belonging to the family *Beggiatoaceae* extensively colonized the sediment surface even down to the deepest part of the EGB. So far these organisms

were only observed in water depths of about 70–120 m where electron acceptors are at least temporarily available (Noffke et al., 2016; Sommer et al., unpublished results). These chemotrophic organisms gain their energy from HS⁻ oxidation using O₂ as terminal electron acceptor (e.g., Teske and Nelson, 2006). However, some members of the *Beggiatoaceae* have the capacity to switch to a NO₃⁻ based HS⁻ oxidation, releasing NH₄⁺ to the environment as a metabolic waste product (reviewed by Jørgensen and Nelson, 2004). In contrast to denitrification and anammox, this process known as dissimilatory nitrate reduction to ammonium (DNRA) retains DIN in the ecosystem and has been argued to be an important benthic process under low O₂ conditions in the HTZ (Noffke et al., 2016). It has also been reported for sediments from Eckernförde Bay (Dale et al., 2011, 2013), the northern Baltic Proper, and the coastal Gulf of Finland (Kuparinen and Tuominen, 2001; Hietanen and Lukkari, 2007; Jäntti et al., 2011; Jäntti and Hietanen, 2012). When all the data is considered collectively, NH₄⁺ release rates under ventilated conditions were similar to those measured during euxinic conditions and thus apparently not greatly enhanced by DNRA. The NH₄⁺ release rates measured during post inflow conditions are consistently below the expected based on Redfield organic matter degradation (**Figure 7A**). However, it is possible that a fraction of NH₄⁺ produced by DNRA may be consumed by close coupling with nitrifying bacteria in adjacent sediment layers, thereby explaining the low N:C ratio of the fluxes. Only a few sites, including those from our previous study in 2010 (Noffke et al., 2016), hint toward an excess of NH₄⁺ release from the sediment, and thus an excess of DNRA over nitrification. It is important to note that during the post-inflow cruise in March



(AL473), larger filaments of *Beggiatoaceae* with internal NO_3^- concentrations of 2.5 ± 1.8 mM dominated within the HTZ at 100 and 110 m water depth as well as at the upper boundary of the deep basin (124 m) (Schulz-Vogt, unpublished). This strongly indicates the potential capacity for DNRA in this region. Due to higher concentrations of O_2 in the deep basin, aerobic respiration by HS^- oxidizing bacteria may have been energetically more preferable than anaerobic, NO_3^- respiration.

Filamentous HS^- oxidizing microbes may also play an important role in benthic P cycling. They are able to perform luxury uptake of PO_4^{3-} from the porewater and bottom water under oxic conditions that is subsequently stored as polyphosphates in their vacuoles (Brock and Schulz-Vogt, 2011). During anoxic conditions, these organisms degrade the polyphosphates and release the stored PO_4^{3-} back into the porewater. Evidence for this metabolic mechanism has been provided for the genera *Thiomargarita* spp. and *Beggiatoa* spp. in sediments from the Namibian upwelling system (Schulz and Schulz, 2005; Goldammer et al., 2010; Winkel et al., 2016). Breakdown of polyphosphates may account for the fact that sediments in the deep basin again acted as a PO_4^{3-} source in

March 2016 where O_2 in the bottom water had decreased to <40 μM . Interestingly, though, during March 2016 filaments of *Beggiatoaceae* were void of polyphosphates, whereas smaller granules of polyphosphates were abundantly observed in the surface sediment (Schulz-Vogt, unpublished). The source of these particles remains an open question, and could conceivably be attributed to authigenic formation at the former pelagic oxycline, dead phytoplankton, or polyphosphate accumulated by smaller benthic bacteria (Diaz et al., 2008, 2012). The absence of polyphosphate in the filamentous *Beggiatoaceae* is an additional indication that polyphosphate accumulation is not a trait shared by all *Beggiatoaceae* under all environmental conditions. Members of the genus *Marithioploca*, (former *Thioploca*) prevailing in sediments off the Chilean coast were also reported not to contain visible polyphosphate granules (Høgslund et al., 2009).

A plot of the PO_4^{3-} vs. DIC flux from all cruises demonstrates that the sediments in the EGB behave in a highly non-Redfield manner (Figure 7B). Most sites with a positive PO_4^{3-} flux lie far above the expected Redfield ratio (solid black curve). Even accounting for preferential remineralization of organic P relative

to C by a factor of two (Dale et al., 2016), additional P sources are needed to explain the data distribution (dashed black curve). Off Peru, Lomnitz et al. (2016) observed a clear correlation between PO_4^{3-} flux and cell number of the filamentous bacterium *Marithioploca*. In principal, as described above, transient P storage and release from bacterial mats of *Beggiatoaceae* spp. that almost completely cover the sediment surface in the HTZ could conceivably contribute to the measured P fluxes if they had accumulated polyphosphate during conditions prior to March 2016. As mentioned, smaller less conspicuous sediment bacteria may instead be responsible for polyphosphate mediated P cycling. In addition, iron-bound P dissolution could be significant and deserves greater attention to constrain potential PO_4^{3-} fluxes from this source (e.g., Jilbert et al., 2011). Clearly, future work must now focus on process-based studies to fully address the factors leading to the highly dynamic nature of N and P fluxes in the EGB.

The O_2 depth profiles obtained during post inflow summer conditions indicate that the MBI only affected water masses deeper than ~ 120 m. Above this, the O_2 levels were close to zero and reached $\sim 30 \mu\text{M}$ at 80 m at the base of the oxycline. During winter inflow the O_2 profile shows a similar trend, yet with slightly higher O_2 levels in the HTZ. An identical O_2 distribution has been previously described during euxinia and, along with the distribution of microbial mats, was used as a major criterion to define the HTZ (Noffke et al., 2016). Further similar observations (e.g., Schmale et al., 2016), coupled with the observation that the sediments were also densely covered with mats of sulfur bacteria, imply that the HTZ can be considered as a typical and persistent feature of this water depth range. Pre- and post-inflow solute fluxes in the HTZ were variable but generally indistinguishable. These findings have important implications because the HTZ has been recognized to be a major region strongly contributing to the internal loading of PO_4^{3-} and NH_4^+ (Noffke et al., 2016).

Revised Budget of Benthic PO_4^{3-} and DIN Fluxes before and after Inflow Events

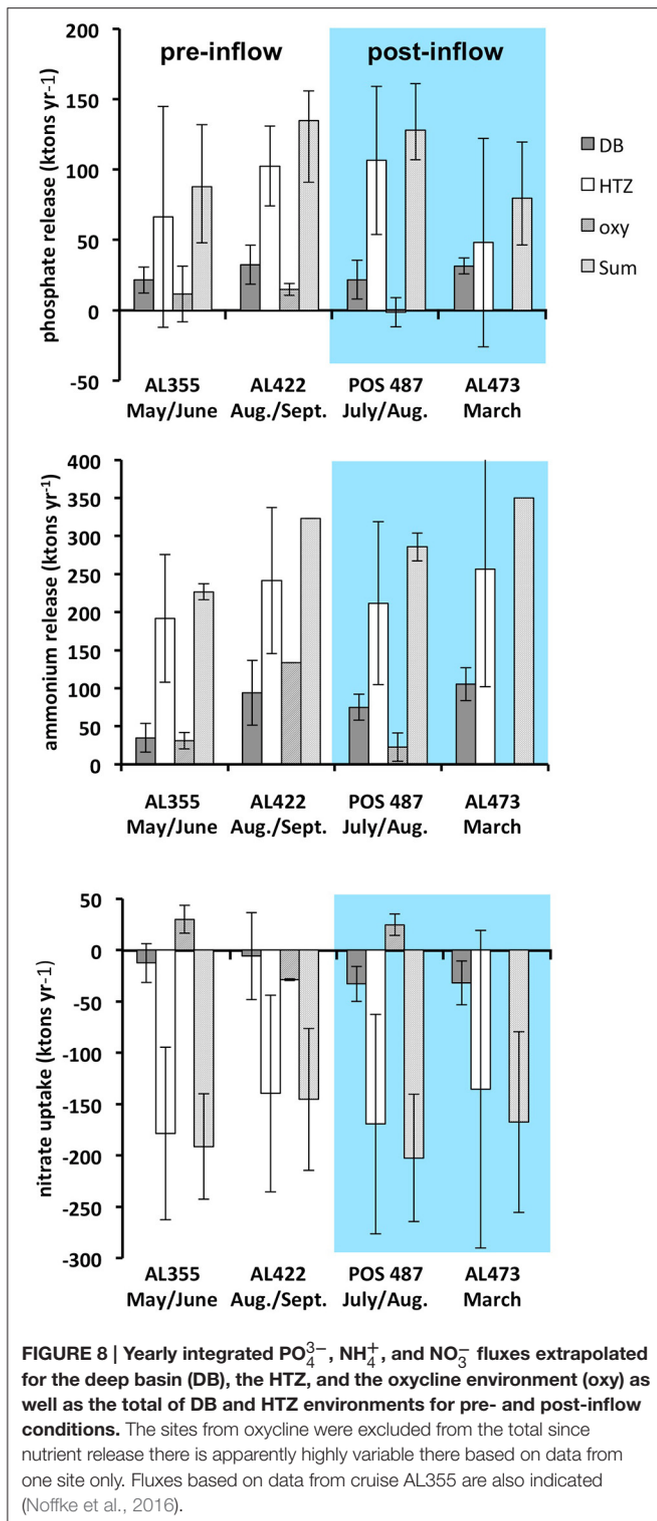
The above discussion has demonstrated that the deep basin and HTZ behaved differently during the most recent MBI events. In order to assess the importance of the expected basin wide reduction of nutrient release during ventilation, we approximated the entire benthic nutrient load for the Baltic Proper before and after ventilation (Table 3) using the approach put forward by Noffke et al. (2016). Here, the Arkona Basin has been excluded from the analysis since this is a seasonally hypoxic setting and the presence of more extensive mats of filamentous sulfur oxidizing bacteria there is in doubt. The Baltic Proper was divided into three depth zones according to the vertical distribution of O_2 in the water column during pre-inflow conditions (Figure 1A, Table 1). To arrive at regional nutrient fluxes, the areas of each depth zone were multiplied with the respective averages of the benthic PO_4^{3-} , NH_4^+ , and NO_3^- ($\text{NO}_3^- + \text{NO}_2^-$) fluxes measured in each depth zone. Our flux measurements were only conducted in water depths below 60 m, which approximately separates

deeper muddy from sandy sediments in the Baltic Proper (www.helcom.fi/baltic-sea-trends/data-maps).

For ease of comparison with other published rates (e.g., Viktorsson et al., 2013a; Noffke et al., 2016) the fluxes are reported on a yearly basis. However, the ventilation events are transient and the strong MBI triggered in December 2014 only lasted from March to September 2015 when O_2 was already consumed 2 m above the seafloor (Mohrholz et al., 2016). The second moderate ventilation event captured during AL473 only lasted for about 3 months (Mohrholz et al., 2016). Hence, we made the assumption that the fluxes we measured in the deep basin in response to the deep-water renewal prevailed only for 6 and 3 months respectively. For the remaining time of the year we assumed fluxes as measured during pre-inflow conditions. This neglects the dynamics of the inflowing water that progressively replaces the anoxic and sulfidic deep water before the shallower sites of the eastern flank of the basin. It also neglects the time needed for the microbiological community to become established in response to oxygenation, although observations of rapid establishment of *Beggiatoaceae* in response to anoxia in Eckernförde Bay implies colonization times of < 1 month (Dale et al., 2013).

Ventilation during the MBI from December 2014 (post-inflow summer cruise POS487) reduces the yearly PO_4^{3-} release from the deep basin sediments by about 33 and about 10% for NH_4^+ (Table 3, Figure 8). In parallel to the flux study during cruise POS487, benthic P fluxes were measured in the ventilated deep basin at 170 and 210 m water depth in July 2015 on the Swedish site of the EGB (Hall et al., 2017). Basin-wide P release extrapolated for the deep basin was $17.4 \text{ ktons year}^{-1}$ compared to $50.3 \text{ ktons year}^{-1}$, which was measured by these authors during euxinic conditions corresponding to a reduction of 65%. During the moderate inflow event from November 2015 (post-inflow winter cruise AL473) there was only a 3% reduction of the PO_4^{3-} release compared to flux measurements made during euxinic conditions (pre-inflow summer cruise AL422). In contrast, NH_4^+ release was even higher compared to all previous cruises.

Importantly, since the fluxes in the HTZ were apparently not greatly affected by the inflows, the HTZ remained a major nutrient release site. With the exception of data from the pre-inflow early summer cruise AL355 (Noffke et al., 2016), differences between pre- and post-inflow conditions are indistinguishable from the range of variability measured in individual chambers. A major source of variability of the fluxes within the HTZ is contributed by the 80 m site located just beneath the oxycline. As already shown (Noffke et al., 2016, cf. their Figure 3), O_2 levels are highly variable there and range between close to zero and $160 \mu\text{M}$, which affects mobilization of PO_4^{3-} as well as N cycling. Despite the reduction of the seabed nutrient release from deep basin sediments, the overall, basin-wide, reduction in nutrient release for the strong MBI (post-inflow summer cruise) only amounts to 5 and 12% for PO_4^{3-} and NH_4^+ respectively. For the moderate MBI (post-inflow winter cruise), there was no reduction with regard to NH_4^+ , yet there was an overall reduction of the PO_4^{3-} release by 42%, which was caused by a drop of PO_4^{3-} release within the HTZ and thus not



greatly affected by the inflow. This implies that changes in benthic fluxes induced by the inflow may simply lie within the natural spatial and temporal variability exhibited by the basin sediments. We excluded the sites from oxycline and shallower waters from

our analysis since nutrient release there is highly variable based only on one site.

The analysis shows that seabed nutrient release in the basin is controlled by processes in the HTZ rather than the deep basin as previously assumed. Even the largest MBI recorded from December 1951 (225 km^3), which compares to 198 km^3 of the MBI in 2014, would be very likely insufficient to ventilate the HTZ and suppress seafloor nutrient release. Despite the uncertainties involved, this simple extrapolation highlights that the effect of deep basin ventilation on the reduction of benthic nutrient release can be considered as minor in the context of the entire nutrient budget. It should be noted that the above extrapolation was conducted under the assumption that the HTZ is present throughout the Baltic Proper (Figure 1). During the MBI triggered in December 2014, an O_2 deficient zone was also retained at the Swedish side of EGB with a still-elevated average P release rate of $0.34 \text{ mmol m}^{-2} \text{ d}^{-1}$ (Hall et al., 2017). Up-scaling of this P flux yields a P-load of $94 \text{ ktons year}^{-1}$, which is very similar to that estimated for the HTZ during euxinia as well as during post inflow conditions presented in this study.

CONCLUSIONS

The intrusion of O_2 and NO_3^- rich North Sea water into the EGB during a major Baltic inflow event in 2014 led to an approximate 33 and 10% reduction of the seabed PO_4^{3-} and NH_4^+ release from deep basin sediments ($>120 \text{ m}$ water depth) compared to euxinic, stagnant conditions that prevailed for the previous decade. Post-inflow, the deep basin, was colonized by vacuolated HS^- oxidizing bacteria tentatively assigned to the family *Beggiatoaceae*. HS^- oxidation was highly efficient and seabed HS^- release was completely suppressed. O_2 consumption during HS^- oxidation in the water column and the sediment-water interface was rapid and must have contributed to the short duration of ventilated conditions that only lasted a few months.

The presence of the HTZ, which has been identified recently as a second major zone for rapid nutrient recycling and nutrient release (Noffke et al., 2016), was confirmed for euxinic deep water conditions as well as for post inflow conditions. Detection of deep-water renewal and associated bottom water ventilation at this particular depth zone where seabed release of PO_4^{3-} and NH_4^+ was ongoing at high rates was not overly apparent. Extrapolation of the *in situ* nutrient fluxes of the deep basin and the HTZ for the Baltic Proper (excluding the Arkona basin and sediments in water depth $<60 \text{ m}$) indicated that the reduction in PO_4^{3-} and NH_4^+ release in response to strong deep-water renewal can be considered as minor, reducing the internal PO_4^{3-} and NH_4^+ load by 5 and 12%, respectively. Despite that the presence of the HTZ has not been confirmed throughout the Baltic Proper, an O_2 deficient zone similar to the HTZ was retained at the Swedish side of the EGB after the strong MBI with similar consequences for the nutrient release.

Ventilation events suppress HS^- toxification and nutrient release only for short time periods of several months. If the inflow events occur as infrequently in the future as during the past decade they have only limited impact to sustainably

reduce internal nutrient loading in the EGB. In the long-term, eutrophication will not be diminished by these events because recycling of P (and N) between the water column and surface sediments is relatively rapid compared to slower sequestration of P by burial in the sediments.

AUTHOR CONTRIBUTIONS

SS, OP, and AD designed the study, coordinated ship operations, lander deployments, sediment sampling, and the data selection process; SS, DC, MY, HS, and AD took and processed samples; All authors contributed ideas and wrote the manuscript.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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