1 Extreme N₂O accumulation in the coastal oxygen minimum zone off Peru

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9 Abstract

10 Depth profiles of nitrous oxide (N₂O) were measured during six cruises to the upwelling area and 11 oxygen minimum zone (OMZ) off Peru in 2009 and 2012/2013, covering both the coastal shelf region 12 and the adjacent open ocean. N_2O profiles displayed a strong sensitivity towards oxygen 13 concentrations. Open ocean profiles with distances to the shelf break larger than the first baroclinic Rossby radius of deformation showed a transition from a broad maximum close to the equator to a 14 15 double-peak structure south of 5°S where the oxygen minimum was more pronounced. Maximum 16 N_2O concentrations in the open ocean were about 80 nM. A linear relationship between ΔN_2O and 17 apparent oxygen utilization (AOU) could be found for all measurements within the upper oxycline, 18 with a slope similar to studies in other oceanic regions. In contrast, N₂O profiles close to the shelf 19 revealed a much higher variability, and N_2O concentrations higher than 100 nM were often observed. 20 The highest N₂O concentration measured at the shelf was ~850 nM. Due to the extremely sharp 21 oxygen gradients at the shelf, N₂O maxima occurred in very shallow water depths of less than 50 m. 22 In the coastal area, a linear relationship between $\Delta N_2 O$ and AOU could not be observed as extremely 23 high $\Delta N_2 O$ values were scattered over the full range of oxygen concentrations. The data points that 24 showed the strongest deviation from a linear $\Delta N_2 O/AOU$ relationship also showed signals of intense 25 nitrogen loss. These results indicate that the coastal upwelling at the Peruvian coast and the 26 subsequent strong remineralization in the water column causes conditions that lead to extreme N₂O 27 accumulation, most likely due to the interplay of intense mixing and high rates of remineralization which lead to a rapid switching of the OMZ waters between anoxic and oxic conditions. This, in turn, 28 29 could trigger incomplete denitrification or pulses of increased nitrification with extreme N₂O 30 production.

31

33 1 Introduction

34 Nitrous oxide (N_2O) acts as a strong atmospheric greenhouse gas and contributes substantially to the 35 stratospheric ozone depletion (IPCC, 2013; WMO, 2011). The ocean is a major source for N₂O as it is 36 naturally produced in the water column (Ciais et al., 2013; Bange, 2008). While in large parts of the 37 surface ocean N₂O concentrations are close to saturation, high emissions of N₂O have been observed in upwelling areas where subsurface waters enriched in N_2O are transported to the surface (e.g. 38 39 Nevison et al., 2004). The global distribution of N_2O in the ocean is closely linked to the oceanic 40 oxygen distribution, and particularly high supersaturations are found in upwelling areas which overlay pronounced oxygen minimum zones (OMZ), e.g. in the Arabian Sea (Bange et al., 2001) or in 41 42 the eastern South Pacific Ocean (Charpentier et al., 2010). 43 These OMZs are key regions for marine nitrogen (N) cycling where active N loss via canonical 44 denitrification and anaerobic ammonium oxidation (anammox) takes place. Particularly in areas 45 where the OMZ is fuelled by high export production, high rates of denitrification and anammox, but 46 also other N transformation processes, such as nitrification, have been observed (Hu et al., 2015; 47 Kalvelage et al., 2013). Oceanic N₂O is mainly produced by nitrification and denitrification, and the 48 interplay of these processes governs the N₂O distribution in OMZs (Bange, 2008). 49 The relationship between N₂O and oxygen concentrations in the ocean in often described by 50 comparing excess N_2O (ΔN_2O) and the apparent oxygen utilization (AOU). As nitrification is one major 51 process accompanying the remineralization of organic matter, a positive correlation between the 52 excess N_2O (ΔN_2O) and the apparent oxygen utilization (AOU) is often interpreted as an indication for 53 nitrification as the main N_2O production pathway (e.g. Walter et al., 2006; Forster et al., 2009). 54 Nitrification can either be performed by bacteria (Arp and Stein, 2003) or archaea (Walker et al., 55 2010). Recent studies indicate that archaea may dominate marine N₂O production under oxic 56 conditions (Löscher et al., 2012; Santoro et al., 2011). The production mechanisms and

57 environmental controls of archaeal N₂O production are subject to ongoing research, however
58 (Stieglmeier et al., 2014).

59 An increase in the $\Delta N_2O/AOU$ ratio at low oxygen concentrations has been observed in several 60 studies in different oceanic areas with reduced oxygen concentrations (Ryabenko et al., 2012; Upstill-61 Goddard et al., 1999; De Wilde and Helder, 1997). This could be explained by several processes: During nitrification, N₂O can either be produced as a side product from the oxidation of ammonium 62 63 to nitrite, or from the reduction of nitrite to N₂O, a process known as nitrifier-denitrification (Stein, 64 2011). Nitrifier-denitrification has been identified as an important production pathway of N_2O at low 65 oxygen concentrations and may thus be responsible for the increased N₂O production under these 66 conditions (Ni et al., 2014). An increase in the N₂O yield of nitrification has indeed been observed in 67 laboratory experiments with bacterial (Goreau et al., 1980) and archaeal ammonium oxidizers 68 (Löscher et al., 2012). The extent to which ammonium oxidation or the nitrifier-denitrification 69 pathway are responsible for N₂O production is yet not well determined (Ostrom et al., 2000; Ni et al., 70 2014), particularly for archaeal nitrification (Löscher et al., 2012; Santoro et al., 2011; Stieglmeier et 71 al., 2014).

72 Additional N₂O production from denitrification has also been proposed as a potential mechanism 73 leading to an increased $\Delta N_2 O/AOU$ at low oxygen concentrations (e.g. Farías et al., 2009; Ji et al., 74 2015). During denitrification, the canonical reduction of nitrate to molecular nitrogen, N₂O evolves as an intermediate product. Denitrification is stimulated by the supply of organic carbon or hydrogen 75 76 sulfide (Chang et al., 2014; Dalsgaard et al., 2014; Galan et al., 2014), and active denitrification is 77 restricted to suboxic to anoxic conditions (e.g. Firestone et al., 1980; Dalsgaard et al., 78 2014).Depending on the environmental conditions, N₂O production or consumption due to 79 denitrification can be observed in the environment. There has been evidence that N₂O consumption 80 is more sensitive to trace amounts of oxygen than N_2O production. This could lead to N_2O 81 accumulation when oxygen is present in low concentrations (Tiedje, 1988). Exceptionally high N₂O 82 concentrations off the West Indian Coast were thus associated with an increased N₂O production

from denitrification during transient oxygen concentrations (Naqvi et al., 2000). In a recent study it
was furthermore shown that N₂O production from denitrification could be stimulated by H₂S addition
(Dalsgaard et al., 2014) which could indicate a coupling between N₂O production and sulfur cycling.
At oxygen concentrations below a threshold value of 4 - 10 µM, (Nevison et al., 2003; Ryabenko et

al., 2012; Cornejo and Farias, 2012), consumption of N_2O in the water column is observed, which 87 leads to a breakdown in the previously described positive $\Delta N_2 O/AOU$ relationship. The exact oxygen 88 89 concentration at which N₂O consumption starts is not yet well determined, however (Cornejo and 90 Farias, 2012; Zamora et al., 2012). N₂O consumption has been associated with denitrification as the 91 only known process to consume N₂O in OMZ waters (Cornejo and Farias, 2012). Although rate 92 measurements only rarely detected active denitrification in the water column of the ETSP (Kalvelage 93 et al., 2013; Hamersley et al., 2007; Thamdrup et al., 2006), the widespread N₂O consumption in the 94 OMZ core is an indicator for active denitrification (Farias et al., 2007).

There is a strong indication that at low oxygen concentrations nitrification and denitrification may
take place in close proximity (Kalvelage et al., 2011), and the N₂O production and consumption under
these conditions are strongly influenced by the interaction of both processes (Ji et al., 2015).
Measurements of N₂O consumption rates in the eastern tropical North Pacific Ocean (ETNP)
furthermore provided evidence for a rapid N₂O cycling, although depth profiles of N₂O seemed to be
relatively invariant over time (Babbin et al., 2015). These quasi-stable conditions may be disturbed by
rapid changes in the environmental conditions.

The eastern tropical South Pacific Ocean (ETSP) harbors one of the four major eastern boundary
upwelling systems (EBUS): alongshore trade winds induce offshore Ekman transport of the surface
water masses which leads to strong coastal upwelling off Peru and Chile (Chavez and Messié, 2009).
While year-round upwelling and high primary productivity can be observed along the Peruvian coast
(Messie et al., 2009), the highest upwelling intensity can be observed during austral winter, whereas
primary production seems to be higher during autumn and spring (Pennington et al., 2006), which

108 may be caused by nutrient and light limitation during phases of intense upwelling (Echevin et al.,109 2008).

The region is influenced by strong seasonal and interannual variability caused by the influence of Equatorial Kelvin waves and the El Niño Southern Oscillation (ENSO). ENSO could cause the interruption of the upwelling during El Niño events (Dewitte et al., 2012; Graco et al., 2016). While the OMZ core is largely unaffected by ENSO, a deepening of the upper oxycline and the reoxygenation of the Peruvian shelf due to the propagation of coastal trapped waves can be observed (Gutierrez et al., 2008).

The ETSP is characterized by one of the largest and most intense OMZs in the oceans, extending from 116 117 the Peruvian shelf about 1000 km offshore with a maximum thickness of more than 600 m 118 (Fuenzalida et al., 2009). It is located in the shadow zone of large ocean current systems which leads 119 to a sluggish ventilation and long residence times of waters within the OMZ (Karstensen et al., 2008). 120 Equatorial current bands such as the Equatorial Undercurrent (EUC) and the Southern Subsurface 121 Countercurrents (SSCC) supply waters to the ETSP which leads to slightly elevated oxygen 122 concentrations in the northern part of our study area, with minimum oxygen concentrations of 10 -123 20 μ M (Stramma et al., 2010), whereas oxygen concentrations below 3 μ M are common in the OMZ 124 core south of 5°S (Paulmier et al., 2006). The equatorial current bands also feed the poleward Peru-125 Chile Undercurrent (PCUC) which is the main source for waters upwelled along the coast (Montes et 126 al., 2010; Chaigneau et al., 2013) and which transports Equatorial Subsurface Water (ESSW) 127 southward. During its spreading, the ESSW is subject to oxygen depletion and mixing with 128 surrounding water masses, e.g. the Antarctic Intermediate Water (AAIW) below and the Eastern South Pacific Intermediate Water (ESPIW) which originates from the South (Wyrtki, 1967; Chaigneau 129 130 et al., 2013). Mixing of different water masses in the upwelling zone creates a distinct coastal water 131 mass which is called Cold Coastal Water (CCW) (Pietri et al., 2013).

- 132 High primary production and high remineralization rates in the underlying waters lead to a further
- drawdown in subsurface oxygen concentrations to near-depleted conditions (Karstensen et al.,
- 134 2008). Active N loss is observed in large parts of the OMZ which is reflected in a pronounced
- secondary nitrite maximum and a strong nitrogen deficit in the OMZ core (Codispoti et al., 1986). The
- 136 OMZ furthermore frequently extends over large parts of the Peruvian shelf where sulfidic conditions
- 137 within the water column can be observed (Schunck et al., 2013).
- 138 Here we present N₂O measurements in the water column off Peru from six measurement campaigns
- in the ETSP. Previous depth profile measurements in this area showed a pronounced double-peak
- 140 structure off South Peru which merged into a broad maximum north of 5°N (Cornejo and Farias,
- 141 2012; Ryabenko et al., 2012). Surface N₂O measurements off Peru furthermore revealed
- 142 extraordinarily high emissions from the Peruvian shelf area which corresponded to extremely high
- 143 surface and subsurface N₂O concentrations (Arévalo-Martínez et al., 2015).
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147 **2** Methods

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In total, 146 depth profiles (0-~4200 m) of N₂O were measured on two cruises between December
2008 and February 2009 (M77-3 & M77-4) and four cruises between October 2012 and March 2013

- 151 (M90 M93) to the upwelling area and the adjacent open ocean off Peru onboard the German
- 152 research vessel Meteor. The Southern Oscillation Indices
- 153 (http://www.ncdc.noaa.gov/teleconnections/enso/indicators/soi/) from 2008/2009 and 2012/2013
- did not indicate the presence of an El Niño event during our measurement campaigns and similar
- 155 conditions between both measurement campaigns could be expected. The locations of the sampled
- 156 stations are shown in Fig. 1. While the M77-4 and M90 cruises mainly covered the open ocean area,
- the M77-3 and M91-M93 cruises mainly took place in the Peruvian shelf area. The work was part of
- the German DFG collaborative research project (SFB) 754 (<u>https://www.sfb754.de/</u>) and the BMBF
- 159 project SOPRAN (Surface Ocean PRocesses in the ANthropocene, <u>www.sopran.pangaea.de</u>). The N₂O
- 160 data set described here has been archived in MEMENTO, the MarinE MethanE and NiTrous Oxide
- 161 database (<u>https://memento.geomar.de</u>) (Kock and Bange, 2015).
- 162 Triplicate samples were taken from 10 L Niskin bottles mounted on a rosette water sampler or a
- 163 pump-CTD (M77-3) in 25 ± 0.11 mL (M77-3 & M77-4) and 20 ± 0.14 mL (M90 M93) opaque glass
- vials and sealed with butyl rubber stoppers and aluminum caps, thereby avoiding the inclusion of airbubbles.

Samples were treated with 0.2 mL (M77-3 & M77-4) and 0.05 mL (M90 - M93) of a saturated
mercuric chloride solution directly after the sampling to inhibit microbial N₂O production or
consumption. The samples were either analyzed onboard (M77-3 & M77-4, M91, partly M90 & M93)
within a few days or shipped to GEOMAR by air freight for later analysis (M92, partly M90 & M93).
Samples that were shipped to Germany were additionally sealed with paraffin wax and stored upside
down to avoid the formation of air bubbles in the samples due to temperature and pressure changes
during transportation.

173 Samples were analyzed using a static equilibration method: 10 mL helium (99.9999%, AirLiquide, 174 Düsseldorf, Germany) was manually injected into each vial which was equipped with a second 175 syringe to collect the overflowing water. Vials with added headspace were vigorously shaken for 176 about 20 s and allowed to equilibrate at ambient temperature for a minimum of two hours. A 177 subsample of the equilibrated headspace was manually injected into a GC-ECD system (Hewlett-178 Packard 5890 Series II, Agilent Technologies, Santa Clara, CA, USA), equipped with a 6' 1/8" packed 179 column (molsieve, 5Å, W. R. Grace & Co.-Conn., Columbia, MY). The GC was operated at 190 °C, using 180 argon/methane (95%/5%, ECD purity, AirLiquide, Düsseldorf, Germany) as carrier gas at a flow rate of 181 30 mL min^{-1} .

182 The GC was calibrated on a daily basis with a minimum of 2 (M77-3 & M77-4) or 4 (M90 - M93)

183 different standard gas mixtures (N₂O in synthetic air, Deuste-Steininger GmbH, Mühlhausen,

184 Germany and Westfalen AG, Münster, Germany). Standard gases were either injected as pure gas or 185 further diluted with helium (1:3, 1:1 or 3:1) to obtain additional standard gas concentrations. Our 186 standard gases were calibrated against NOAA primary standards at the Max Planck Institute for 187 Biogeochemistry in Jena, Germany, if the standard gas concentrations were within the calibration 188 range of the NOAA gases. Gases with N₂O concentrations outside the NOAA calibration range were 189 internally calibrated using an LGR N₂O/CO analyzer (Los Gatos Research, Mountain View, CA, USA), 190 which was proven to have a linear response and minimal drift within the calibration range (Arévalo-191 Martínez et al., 2013). The N₂O concentration in the samples was calculated according to Walter et 192 al. (2006) using the solubility function of Weiss and Price (1980). The average precision of the 193 measurements, calculated as median standard deviation from triplicate measurements, was 0.7 nM. $\Delta N_2 O$ was calculated as the difference between the in-situ concentration $[N_2 O]_w$ and the equilibrium 194 195 concentration $[N_2O]_{eq}$:

196
$$\Delta N_2 O = [N_2 O]_w - [N_2 O]_{eq}$$
(1)

197 We used the contemporary atmospheric mixing ratio measured at Cape Grim, Tasmania

198 (http://agage.mit.edu/data/agage-data) for the calculation of $[N_2O]_{eq}$. This calculation

199 underestimates the N₂O excess in subsurface waters which have been isolated from the surface for a

200 long time as it does not account for the increase in the atmospheric mixing ratio since the beginning

201 of the industrial revolution (Freing et al., 2009). The use of the contemporary N₂O mixing ratio of

202 2013 would lead to a maximum ~17% overestimate of $[N_2O]_{eq}$, thus leading to only a small error

203 compared to the maximum N₂O concentrations measured in our study, and the use of the

204 contemporary atmospheric mixing ratio still allows a qualitative analysis of the $\Delta N_2 O/AOU$

205 relationship in order to investigate the formation and consumption processes of N₂O.

The potential temperature of the water parcel at a certain depth was calculated using the Gibbs
Seawater Oceanographic Toolbox (McDougall and Barker, 2011).

Oxygen concentrations were measured either with a Seabird (M77-3 & M77-4: SBE-5; M90-M93: SBE
43) oxygen sensor (Sea-Bird Electronics, Bellevue, WA, USA) mounted on the CTD rosette or from
100 mL discrete samples taken from the Niskin bottles and analyzed using the Winkler titration
method (Grasshoff et al., 1999). The oxygen sensor was calibrated against the Winkler
measurements.

213 Recent studies using highly sensitive STOX (Switchable Trace amount Oxygen) sensors for oxygen 214 measurements indicate that measurements with conventional oxygen sensors that are calibrated 215 against Winkler measurements may be biased towards higher concentrations at near-zero oxygen 216 conditions. Thamdrup et al. (2012) therefore argued that anoxic conditions are prevalent in the core 217 of the Peruvian OMZ where oxygen concentrations of several μ M have been found using the 218 conventional Winkler-calibrated measurements. As STOX sensor measurements were not available 219 for all measurement campaigns presented here, the minimum oxygen measurements reported here 220 from the core of the OMZ (3-5 μ M) should be considered as an overestimation.

- The Apparent Oxygen Utilization (AOU) was calculated from the oxygen concentrations $[O_2]_w$ using the CSIRO SeaWater library, version 3.2
- (<u>http://www.cmar.csiro.au/datacentre/ext_docs/seawater.htm</u>) to calculate oxygen saturation
 [O₂]_{eq}:

225
$$AOU = [O_2]_w - [O_2]_{eq}$$
 (2)

Nutrient samples ($[NO_3^{-}]$, $[NO_2^{-}]$, $[PO_4^{3^{-}}]$, $[NH_4^{+}]$) from the CTD rosette were analyzed onboard following the nutrient analysis methods according to Hansen et al. (1999). Samples taken from the pump-CTD during M77-3 were stored at -20°C and shipped to Germany for later analysis. N' was calculated as a measure for the nitrogen deficit from the nitrate ($[NO_3^{-}]$), nitrite ($[NO_2^{-}]$) and phosphate ($[PO_4^{3^{-}}]$) concentrations as follows (Altabet et al., 2012):

231
$$N' = ([NO_3^-] + [NO_2^-]) - 16[PO_4^{3-}]$$
 (3)

To distinguish between coastal and open ocean stations we calculated the distance of each station from the continental slope (2000 m isobath) and used the first baroclinic Rossby radius of deformation as described by Chelton et al. (1998) as threshold distance for stations that were influenced by coastal upwelling.

237 **3 Results**

238 **3.1** Spatial distribution of oxygen, nutrients and N₂O

The distribution of oxygen, nitrite and N₂O along an offshore section between 16°S and 2°N at 86° W from the M77-4 cruise in 2009 and the M90 cruise in 2012 is shown in Figure 2; a coastal cross-shelf section along 12°S with the distribution of oxygen, nutrients, N' and N₂O is shown in Figure 3 and selected depth profiles of oxygen, N₂O and potential density (σ_{θ}) as well as nitrate, nitrite, ammonium and N' are shown in Figure 4.

Along 86° W, a similar distribution of oxygen, nitrite and N₂O was observed during M77-4 and M90.

245 Oxygen and N₂O profiles concentrations were close to saturation in the mixed layer. The mixed layer

246 depth increased from below 20 m in the northern part of the section to more than 100 m south of

247 15 °S. Below the mixed layer, a sharp oxycline with a decrease to oxygen concentrations below

 $10 \,\mu\text{M}$ was observed south of 5°S, whereas in the northern part of the section, below the mixed layer

oxygen concentrations only decreased to \sim 100 μ M in the upper 200 m and further dropped to

250 $\,$ concentrations ~10 μM between 200 and 500 m. Minimum oxygen concentrations in the water

251 column increased towards the north from below 5 μ M south of 5° S to ~10 μ M at the equator.

The nitrite distribution revealed a primary maximum at the base of the mixed layer with maximum nitrite concentrations below 1.5 μ M. This primary maximum is frequently observed in the ocean and is usually associated with nitrification (Codispoti and Christensen, 1985). South of 5° S, a secondary nitrite maximum was observed within the OMZ where oxygen concentrations fell below 5 μ M. Nitrite concentrations in the secondary maximum reached up to ~4 μ M.

Along the cross-shelf section at 12°S, the upper OMZ boundary significantly became shallower
towards the coast as a signal of upwelling, with a well oxygenated mixed layer of ~50 m in the open
ocean and a mixed layer depth of less than 5 m on the shelf. Oxygen was strongly undersaturated in
the surface waters on the shelf as a result of upwelling of waters from the underlying OMZ (Fig. 4).
Elevated phosphate concentrations in the surface waters at the coast also reflected the upwelling on

the shelf, whereas nitrate was depleted in the water column and the surface waters close to the
coast, which was also reflected in very low N' values at the inshore stations (Fig. 3). A primary and
secondary nitrite maximum at the base of the mixed layer and in the OMZ core was observed
throughout the cross-shelf section, but both maxima were much more pronounced on the shelf than
in the deep waters (Fig. 3).

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268 The N₂O distribution along 86°W could be divided into two different regimes: north of 5 °S, a broad 269 N_2O maximum with concentrations of ~60 nM coincided with the depth of the oxygen minimum, 270 whereas in the southern part of the section, the N_2O profiles revealed a double-peak structure with a 271 sharp N_2O maximum in the upper and lower oxycline and N_2O depletion in the OMZ core, also 272 coinciding with the secondary nitrite maximum. This shape of N₂O profiles has been observed in OMZ 273 regions before (e.g. Law and Owens, 1990; Cohen and Gordon, 1978). The transition from profiles 274 with a broad N_2O peak to a double-peak structure coincided with the decrease in the minimum 275 oxygen concentrations towards the South. N₂O depletion was observed in profiles were oxygen 276 concentrations dropped below $\sim 5 \,\mu$ M and nitrite accumulation was observed. 277 Compared to the offshore waters, the N₂O distribution on the shelf and in the adjacent waters 278 showed a much larger variability. N₂O depletion was in fact observed at oxygen concentrations below

5 μ M, too. While several N₂O profiles revealed a shape similar to the offshore profiles, an overall

280 characteristic shape could not be identified, however: profiles with a subsurface N₂O maximum in the

oxycline were observed as well as profiles with multiple maxima or a surface N₂O maximum (Fig. 4).

282 While N₂O concentrations in the offshore waters did not exceed 80 nM, N₂O concentrations above

283 100 nM were frequently observed at the shelf. Several profiles showed an extreme N_2O

accumulation with concentrations above 500 μ M (maximum ~850 nM) (Fig. 4). The location and

shape of the N₂O maxima in the different profiles was highly variable, which resulted in a very patchy

286 distribution of N_2O in the water column (Fig. 3).

287 **3.2** N₂O in different water masses of the ETSP

The water mass distribution in our dataset agrees well with the data presented by Pietri et al. (2014) 288 289 (Fig. 5). Due to the larger area covered by our measurements our data showed a broader scattering, 290 but we could identify the same water masses in our data: below 500 m, both the coastal and the 291 offshore profiles carry relatively fresh (S~34.8) and cool (T_{pot}~5°C) Antarctic Intermediate Water 292 (AAIW) (Pietri et al., 2014) that carried relatively high oxygen and $\Delta N_2 O$ values which corresponded 293 to the secondary N₂O maximum in the lower oxycline. Shallower subthermocline waters are covered 294 by the Equatorial Subsurface Water (ESSW) (S~34.8 – 35.2, T_{pot}~9-14 °C) (Wyrtki, 1967). This water 295 mass carried very low oxygen down to concentrations, while $\Delta N_2 O$ values showed either a maximum 296 or N₂O depletion in this water mass, which reflects the strong sensitivity of net N₂O consumption to 297 variations in the oxygen concentration.

298 Waters with low salinities (~34.7), relatively high oxygen concentrations and potential temperatures 299 between 10°C and 15°C can be traced back to Eastern South Pacific Intermediate Water (ESPIW) 300 (Schneider et al., 2003). $\Delta N_2 O$ values within this water mass were between 20 and 30 nM. Pietri et al. 301 (2014) identified narrow patches of ESPIW below the thermocline about ~100 km offshore. We 302 hardly found this water mass in the coastal data, but it is likely mixed with the ESSW and surface 303 waters on the shelf, where it contributes to the formation of Cold Coastal Water (CCW) (Pietri et al., 304 2014). CCW with S \sim 35.1 and T_{oot} \sim 15.5°C was prevalent over the shelf and could only be identified in 305 the coastal data as it is directly related to the coastal upwelling. Offshore surface data were 306 associated with Subtropical Surface Water (STSW) with salinities above 35.0 and temperatures higher 307 than 17° C (Pietri et al., 2013), while surface waters at the coast showed properties that resulted 308 from the warming of the CCW and the mixing with STSW.

309 Very variable $\Delta N_2 O$ values were associated with the CCW and its related surface waters, and nearly 310 all data points that showed extreme N₂O accumulation fell within these waters. This indicates that 311 the extremely high N₂O concentrations were locally produced in the upwelling area, as none of the 312 source water masses for the upwelling carried similarly high ΔN_2O values.

313 3.3 ΔN₂O/AOU relationship

314 A two-linear $\Delta N_2O/AOU$ relationship has been identified in the upper oxycline for waters with oxygen 315 concentrations higher than 50 μ M and between 50 μ M and 5 μ M during the M77-4 cruise that took 316 place in the offshore waters of the OMZ (Ryabenko et al., 2012). We found a very similar relationship 317 for all offshore data with oxygen concentrations above 50 µM with no systematic difference between 318 the data from the M77-4 (January/February 2009) cruise and the M90 (November 2012) cruise 319 (Figures 2, 6a). This indicates a comparable setting of the open ocean OMZ waters during both 320 cruises. We furthermore found no difference in the $\Delta N_2 O/AOU$ relationship between stations with a 321 broad N₂O maximum and a double-peak structure. These results are similar to previously reported 322 $\Delta N_2O/AOU$ relationships from other oceanic OMZs (Upstill-Goddard et al., 1999; Cohen and Gordon, 323 1978; De Wilde and Helder, 1997).

In contrast to the open ocean waters, a correlation between ΔN₂O and AOU was not observed for the
coastal data (Fig. 6b). Numerous values with much higher ΔN₂O/AOU ratios than in the offshore
waters were observed. These data were highly scattered over the full range of oxygen
concentrations. The ΔN₂O values that showed the strongest deviation from the offshore ΔN₂O/AOU
ratio were associated with highly negative N' values as a signal for a large nitrogen deficit (Fig. 6b).
This indicates that these waters with extreme N₂O accumulation had been subject to extensive N
loss.

332 4 Discussion

To understand the differences between the offshore and the coastal N₂O distribution in the Peruvian
 upwelling, the factors that influence N₂O production or consumption during nitrification and
 denitrification need to be investigated.

336 In the oxycline waters of OMZs, peak N₂O production from nitrification as well as denitrification has 337 been determined under suboxic conditions, whereas N₂O depletion was dominant in the OMZ core (Ji 338 et al., 2015). Rate measurements however provided evidence that N₂O production and consumption 339 co-occur and that interplay between N_2O production and consumption processes regulates net N_2O 340 accumulation or depletion in the water column (Babbin et al., 2015). In open ocean OMZs, however, 341 N₂O profiles reveal a remarkably stable shape which indicates that in these areas N₂O production and 342 consumption processes are well balanced (Babbin et al., 2015). Except for differences between the 343 offshore N₂O profiles with a broad N₂O maximum north of 5 °S and a double-peak structure south of 344 5° S, the offshore N₂O profiles observed in our study indeed showed a relatively invariant N₂O 345 distribution. The differences in the shape of the N₂O profiles can be explained by changing oxygen 346 concentrations in the OMZ core and a threshold oxygen concentration of 5 μ M for net N₂O 347 consumption.

348 In areas where highly oxygen-deficient waters extended over the continental shelf, extreme 349 accumulation of N₂O has been found before in the Arabian Sea (Naqvi et al., 2010; Naqvi et al., 2006) 350 and off Chile (Farías et al., 2015) and has been explained by rapid changes in the environmental 351 conditions: Naqvi et al. (2000) explained the extreme N₂O accumulation over the Indian shelf with 352 the response of denitrifying enzymes to transient oxygen depletion. N₂O thus accumulated when 353 waters reached suboxic conditions. N₂O accumulation coincided with the accumulation of nitrite and 354 consumption of N₂O started when these waters became sulfidic (Naqvi et al., 2010). Farías et al. 355 (2015) measured N₂O accumulation during the transition from oxic to anoxic conditions, too, but at 356 variable oxygen concentrations whereas N₂O depletion became dominant under suboxic conditions.

In contrast to the results from the Indian Ocean, they identified enhanced remineralization due to
 short-term variability in coastal upwelling as the main driver for N₂O accumulation.

359 The large variability we observed in the N₂O distribution at the Peruvian coast could also be 360 explained by an imbalance between N_2O production and consumption processes that may lead to its 361 accumulation. This could have been induced by rapid changes of the oxygen concentrations in the 362 coastal upwelling zone: enhanced mixing of oxygen-rich and oxygen deficient waters and exchange of 363 upwelled waters with the atmosphere supply oxygen to the water column (Schafstall et al., 2010; 364 Thomsen et al., 2016 ; Pietri et al., 2014) while strong remineralization leads to rapid oxygen 365 consumption (Kalvelage et al., 2015). Kalvelage et al. (2011) furthermore showed that these high 366 remineralization rates also induce strong N cycling in the subsurface layer. Turnover rates for 367 different N species are therefore much faster on the shelf than in the open ocean OMZ (Hu et al., 368 2015), which is also reflected in the distribution of different functional gene abundances (Löscher et 369 al., 2014). Hence, it is likely that N_2O production and consumption rates are much higher at the coast 370 than in the offshore waters, and that short periods of increased N_2O production could lead to very 371 high N₂O accumulation.

372 Changes in the oxygen concentrations could influence N₂O production from nitrification as well as 373 from denitrification: enhanced production of N₂O after transition from anoxic to oxic conditions is a 374 known process occurring in soils (e.g. Morley et al., 2008) and may be explained by a different 375 sensitivity of denitrifying enzymes to trace concentrations of oxygen (Tiedje, 1988). In a recent 376 incubation study, Dalsgaard et al. (2014) found no indication of increased N_2O production by 377 denitrification due to changes in the oxygen concentration at nanomolar levels, however. Instead, 378 autotrophic denitrification and N₂O production have been shown to be stimulated by the addition of 379 hydrogen sulfide (H₂S) (Galan et al., 2014; Dalsgaard et al., 2014). We did not find direct evidence for 380 a coupling between N_2O production and the presence of H_2S in our measurements, as high N_2O 381 accumulation was often found in proximity to H₂S plumes but was also detected when H₂S was 382 absent in the water column. We cannot exclude that the high N_2O production we frequently

383 observed at the shelf was stimulated by a coupling of denitrification with sulfur cycling, though:

384 Canfield et al. (2010) found evidence for active sulfur cycling in the ETSP without H₂S accumulation,

385 and a coupling between H₂S oxidation and denitrification has been shown before (Galan et al., 2014;

Jensen et al., 2009). Indeed, active denitrification was found in proximity to H₂S plumes in the water

column during M77-3 (Kalvelage et al., 2013; Schunck et al., 2013).

388 In the ocean, increased N₂O production was also associated with the onset of nitrification after re-389 ventilation of the water column in a seasonal study in the Baltic Sea, but with relatively low resulting 390 N_2O concentrations (Naqvi et al., 2010). Yu et al. (2010) found strongly increased N_2O production by 391 nitrifying bacteria that was stimulated by the availability of ammonium during recovery from anoxic 392 conditions in a chemostat culture experiment. Their results point towards an increased N₂O 393 production via the ammonium-oxidation pathway, while N₂O production by nitrifier-denitrification 394 seemed not to be stimulated by the shift from anoxic to oxic conditions. We frequently measured 395 high ammonium concentrations along the Peruvian shelf, indeed (Fig. 4), which could have 396 stimulated N_2O production from ammonium oxidation. A direct correlation between N_2O and 397 ammonium could not be identified, however.

From our concentration measurements alone we thus cannot distinguish if the observed high production of N₂O is a result of denitrification or nitrification processes. Studies of the isotopic and isotopomeric N₂O composition and N₂O production and consumption rate measurements could reveal more detailed insights whether N₂O is produced via the ammonium oxidation or the nitrite reduction pathway during its extreme accumulation.

In our study, we found strongly elevated N₂O concentrations (>100 nM) over the full range of oxygen
concentrations, coinciding with strong N depletion (Fig. 5), but without nitrite accumulation (Fig. 4).
The high oxygen concentrations found in the majority of our samples with extreme N₂O
accumulation and N depletion excludes in-situ denitrification or anammox (see e.g. Babbin et al.,

407 2014; Dalsgaard et al., 2014).

The extraordinarily high N₂O concentrations as well as the low N' values thus have to be old signals of processes taking place under anoxic to suboxic conditions. There is no known consumption process for N₂O in oxygenated waters (Bange, 2008), and the strong signals of N loss that are produced under anoxic conditions are unlikely to be rapidly compensated by N fixation upon oxygenation. Both signals thus are likely to have remained preserved when oxygen concentrations increased due to mixing with waters of higher oxygen concentration or due to direct contact with the atmosphere as a result of upwelling.

Our observations of high N₂O concentrations in oxygenated waters furthermore indicate that this accumulation could have taken place during re-oxygenation rather than during decreasing oxygen concentrations. An increase in oxygen concentrations would lead to the preservation of the high N₂O signals in the water column whereas further decreasing oxygen concentrations would only lead to a temporal N₂O accumulation and would eventually stimulate N₂O consumption.

420

421 4 Summary and Conclusions

422 We observed extreme N₂O accumulations over the Peruvian shelf and in the adjacent waters with 423 maximum concentrations similar to the observations by Naqvi et al. (2000) over the West Indian shelf 424 and Farías et al. (2015) off Chile, whereas N₂O concentrations in the open ocean OMZ off Peru were 425 comparably moderate. Similar to the findings by Naqvi et al. (2000), we found that N_2O accumulation 426 could be caused by enhanced N_2O production by nitrification or denitrification under transient 427 oxygen concentrations. We found strong evidence that these N₂O accumulations are preserved when 428 oxygen concentrations increased as a result of mixing and exchange with the overlying atmosphere in 429 the upwelling zone. Waters with high N₂O concentrations can thus be directly and frequently 430 transported to the surface ocean. This makes this region one of the most important oceanic regions 431 for N₂O emissions to the atmosphere (Arévalo-Martínez et al., 2015). This direct link between 432 unusually high N₂O production and emissions over the Peruvian shelf makes it necessary to

433 understand the biogeochemical processes involved in N₂O production and consumption to produce 434 reliable predictions of oceanic emissions from this area. Current approaches to model the N₂O distribution rely on parameterizations based on the linear ΔN₂O/AOU relationship (Suntharalingam 435 436 and Sarmiento, 2000; Nevison et al., 2003; Freing et al., 2012). These approaches could in fact 437 reproduce the oxygen distribution in the open ocean OMZ off Peru reasonably well, but they fail to 438 account for the extreme N₂O accumulation and its high spatial and temporal variability over the shelf 439 area. They thus significantly underestimate the emissions from the Peruvian upwelling and 440 potentially other upwelling areas with similar conditions, too.

441

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717 Figures:

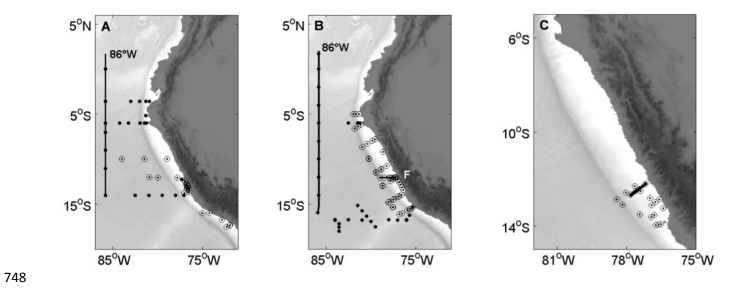
- 718 Figure 1: Station maps of the sampled N₂O stations from cruises A) M77-3, December 2008 January
- 719 2009 (•) and M77-4, January February 2009 (☉), B) M90, November 2012 (•) and M91,
- 720 December 2012 (\odot), C) M92, January 2013 (\bullet) and M93, February March 2013 (\odot). Section
- annotations in A) and B) correspond to the vertical sections shown in Fig. 2 and 3.
- Figure 2: Spatial distributions of oxygen (A, B), nitrite (C, D) and N₂O (E, F) along 86°W during M77-4
 (2009, A, C, E) and M90 (2012, B, D, F). Small dots indicate location and depth of the discrete
- samples. Data gridding: ODV/DIVA.

Figure 3: Cross-shelf distribution of A) Oxygen, B) Phosphate, C) Nitrate, D) N', e) Nitrite and f) N₂O
 during M91 (Section F).

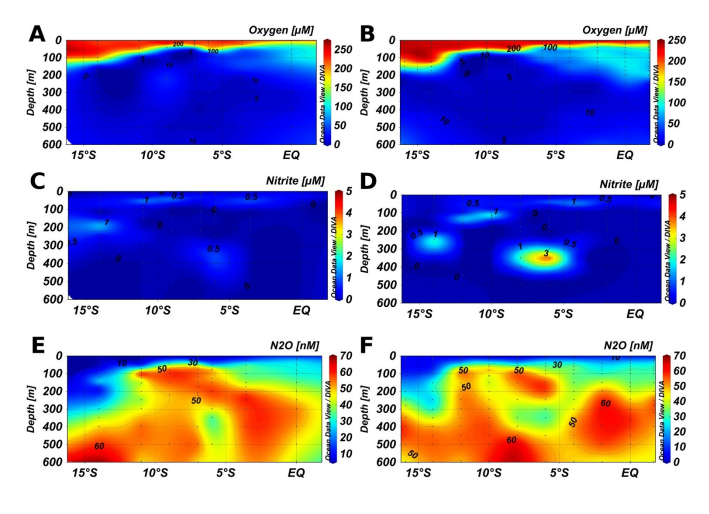
727 Figure 4: Selected depth profiles of oxygen (black dots, dotted line), potential density (σ_{θ} , grey line) 728 and N₂O (red line, open circles) (left panel) and nitrate (grey line), nitrite (black circles, dotted 729 line), ammonium (blue diamonds, straight line) and N' (red line, small dots) (right panel) from 730 selected open ocean and shelf stations during M90-93. Depth profiles of oxygen and σ_{θ} were 731 taken from the CTD sensors, whereas the other parameters were taken from discrete samples. 732 The locations of the respective stations are shown in the map. Red signals denote stations 733 classified as "coastal" stations whereas blue signals denote "offshore" stations. Please note the 734 changes in the scales for N₂O, σ_{θ} , nitrite and ammonium.

Figure 5: Temperature-Salinity diagrams with ΔN₂O color coded for a) the offshore stations and b)
the onshore stations. Gray symbols denote the T-S properties of a) the onshore and b) the
offshore data. The approximate location of the different water masses annotated in the figure is
given by black dots or lines. Different symbols denote different cruises: □ M77-3; ◊ M77-4; ○
M92; ▷M90; < M91; *M93.

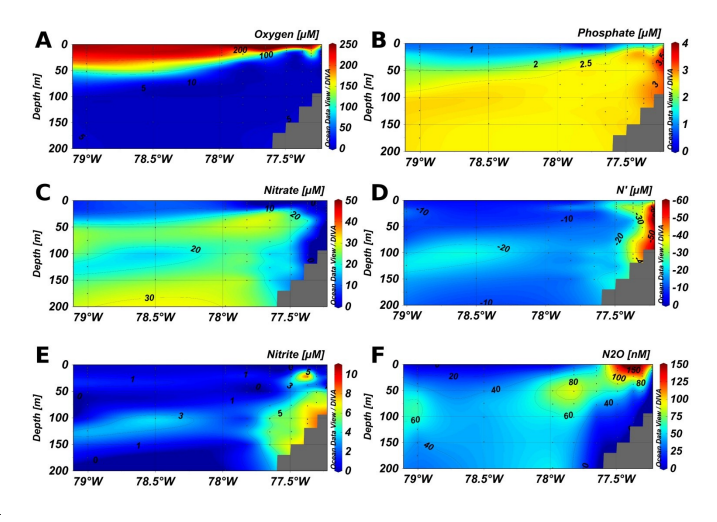
- Figure 6: ΔN₂O/AOU relationship from a) offshore stations and b) coastal stations. Samples from the
- vpper OMZ and oxycline (sample depth < 350 m) are color coded with N', whereas samples from
- below 350 m are shown in gray. Different symbols for different cruises are denoted the same as in
- Figure 5. The black line denotes the $\Delta N_2 O/AOU$ relationship from the offshore data for samples
- with $O_2 > 50 \mu$ M and depth < 350 m (y=0.13x+3.73; r²=0.83). Please note the change in the scaling
- for $\Delta N_2 O$ values of 0 100 nM and 100 1000 nM (dotted line).

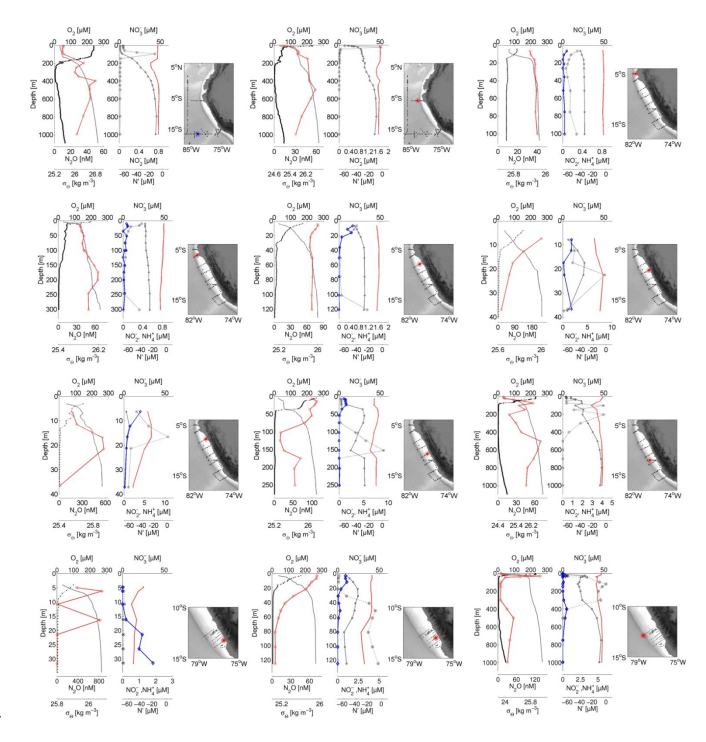


750 Figure 2:

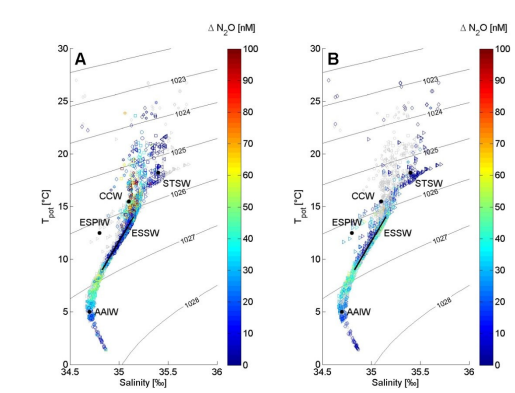


753 Figure 3:









763 Figure 6:

