Prolific nitrite re-oxidation across the Eastern Tropical North Pacific Ocean

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Abstract

Marine Oxygen Deficient Zones serve as hotspots for the loss of fixed nitrogen for the world's oceans, and fixed nitrogen limits primary productivity in large expanses of the ocean. This fixed nitrogen loss occurs primarily through denitrification, where the stepwise reduction of nitrate to nitrite and ultimately to dinitrogen gas is coupled to organic matter oxidation. Nitrite, the first intermediate in denitrification, can also be re-oxidized back to nitrate in a reaction by chemoautotrophic microbes. Nitrite's partitioning between reduction and oxidation determines if marine fixed nitrogen is lost or recycled. Nitrite oxidation in anoxic waters has been previously studied through stable and tracer isotope experiments, but the difficulty of these measurements has limited their geographical distribution and therefore requires extrapolation to understand their impact on the nitrogen cycling. Using basin-scale data, we analyze the progression of nutrients within the three water masses that feed the Eastern Tropical North Pacific Oxygen Deficient Zone. Significant deviations from the expected stoichiometry for denitrification demonstrate that 79% of the nitrite produced in the upper region of the Oxygen Deficient Zone is re-oxidized, whereas only 54% of the nitrite produced in the lower region of the Oxygen Deficient Zone is re-oxidized. These large estimates for nitrite re-oxidation reveal significant fixed nitrogen recycling across the Eastern Tropical North Pacific. Prolific nitrite re-oxidation across the Eastern Tropical North Pacific Ocean
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10 Abstract

Marine Oxygen Deficient Zones serve as hotspots for the loss of fixed nitrogen for the 11 12 world's oceans, and fixed nitrogen limits primary productivity in large expanses of the ocean. This fixed nitrogen loss occurs primarily through denitrification, where the stepwise reduction of nitrate 13 14 to nitrite and ultimately to dinitrogen gas is coupled to organic matter oxidation. Nitrite, the first 15 intermediate in denitrification, can also be re-oxidized back to nitrate in a reaction by 16 chemoautotrophic microbes. Nitrite's partitioning between reduction and oxidation determines if marine fixed nitrogen is lost or recycled. Nitrite oxidation in anoxic waters has been previously 17 18 studied through stable and tracer isotope experiments, but the difficulty of these measurements has 19 limited their geographical distribution and therefore requires extrapolation to understand their 20 impact on the nitrogen cycling. Using basin-scale data, we analyze the progression of nutrients 21 within the three water masses that feed the Eastern Tropical North Pacific Oxygen Deficient Zone. 22 Significant deviations from the expected stoichiometry for denitrification demonstrate that 79% of 23 the nitrite produced in the upper region of the Oxygen Deficient Zone is re-oxidized, whereas only 54% of the nitrite produced in the lower region of the Oxygen Deficient Zone is re-oxidized. These 24 large estimates for nitrite re-oxidation reveal significant fixed nitrogen recycling across the Eastern 25 Tropical North Pacific. 26

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28 Introduction

Denitrification consists of a series of reduction reactions that convert fixed nitrogen, which is generally available for biological reactions, to dinitrogen gas, which is unavailable for these reactions. These reactions occur within the oxygen deficient layers of the water column within marine Oxygen Deficient Zones (ODZs) as well as within marine sediments. Denitrification is the

primary process removing fixed nitrogen in the marine fixed nitrogen budget (Brandes and Devol 33 34 2002) and therefore an important regulatory control on marine productivity. During denitrification, organic matter is oxidized to carbon dioxide and inorganic phosphate is produced from organic 35 phosphorus compounds, causing a stoichiometric decrease in nitrate and increases in total carbon 36 as well as phosphate. While the inorganic C:N:P stoichiometry for aerobic respiration of organic 37 matter is 106:16:1 and commonly referred to as Redfield stoichiometry, similar Redfield 38 relationships exist for other metabolic pathways. The inorganic C:N:P stoichiometry during 39 denitrification from nitrate to dinitrogen gas is 106:-94.4:1 (Froelich et al. 1979), and estimates of 40 fixed nitrogen loss commonly rely on comparing nitrate and phosphate concentrations, such as N* 41 (Gruber and Sarmiento 1997). 42

The first reaction in the denitrification process is dissimilatory reduction of nitrate to nitrite. 43 44 As denitrification continues, further reduction of this nitrogen converts nitrite to a series of intermediates consisting of nitric oxide and nitrous oxide before finally converting it to dinitrogen 45 46 gas. Due to their more widespread biological availability, nitrate and nitrite are often considered "fixed nitrogen", unlike nitric/nitrous oxide and dinitrogen gas (Chang et al. 2014). In the 47 48 canonical denitrification process, nitrate proceeds sequentially through these reduction reactions, 49 however, studies have found that nitrite is frequently re-oxidized to nitrate within ODZs. The 50 pathways for nitrite oxidation are currently not well-defined and spatially heterogenous throughout ODZs (Sun et al. 2021), with potential oxidizing agents such as oxygen, iodate, Mn⁴⁺ and even 51 52 nitrite itself through a dismutation reaction (Babbin et al. 2017, 2020). An ammonia-oxidizing archaeon was recently discovered that is able to produce oxygen in dark, anoxic ocean waters 53 54 (Kraft et al. 2022), which could provide an oxygen source within the anoxic ODZ. Previous studies into the fate of nitrite in the Pacific ODZs found that the vertical structure of the ODZ influenced 55 56 the relative amount of oxidation as well as the oxidizing agents of nitrite re-oxidation. Elevated 57 nitrite re-oxidation has been found in the edges of ODZs, surrounding the oxygen deficient layer that primarily denitrifies (Casciotti et al. 2013), due to the flux of trace amounts of oxygen into the 58 ETNP ODZ. Nevertheless, nitrite oxidation can occur anaerobically (Sun et al. 2021), as the 59 additional oxygen atom in nitrate versus nitrite is derived from water (DiSpirito and Hooper 1986). 60

Recent isotopic tracer and stable isotope studies have measured the rates of nitrite oxidation
and, in some studies, estimated the relative partitioning of nitrite to reduction versus re-oxidation.
In the Costa Rica Dome, the southern side of the Eastern Tropical North Pacific (ETNP) ODZ,

50% of the nitrite was re-oxidized (Buchwald et al. 2015). Data from near the center of the ETNP 64 ODZ also found consistently high rates of nitrite oxidation throughout the ODZ, including the 65 oxygen deficient layer (Babbin et al. 2020) as well as the upper oxycline (Peng et al. 2015). Due 66 to the laborious nature of these measurements (Garcia-Robledo et al. 2016), they are still scarce, 67 such that estimates of the impact of nitrite oxidation across the ETNP ODZ requires significant 68 extrapolation. Fortunately, nitrite oxidation does not stoichiometrically influence inorganic 69 phosphate (Anderson et al. 1982) because nitrite oxidizers are chemoautotrophs (Lipschultz et al. 70 1990). We analyze deviations from 106:-94.4:1 C:N:P stoichiometry (Froelich et al. 1979) to 71 estimate the magnitude of chemical transformations occurring within the water masses of the 72 Eastern Tropical North Pacific. We focus on positive deviation in the -94.4 N value in the 106:-73 94.4:1 C:N:P stoichiometry (Froelich et al. 1979), which has three potential explanations. First, 74 75 nitrite is being re-oxidized back to nitrate, which impacts both the C:N and N:P stoichiometries. Second, the proportion of carbon relative to nitrogen in organic matter being oxidized is lower 76 77 (Van Mooy et al. 2002), decreasing the C:N and C:P stoichiometries. Third, carbon fixation through both photo- and chemo-autotrophy as well as carbonate dissolution occurs, which lowers 78 79 or elevates the C:N and C:P stoichiometries. Only nitrite re-oxidation impacts the N:P 80 stoichiometry without significantly changing the C:P stoichiometry, because carbon fixation will 81 be accompanied by phosphate uptake for autotrophs. This behavior suggests that the C:P 82 stoichiometry identifies the magnitude of the second and third processes on these water masses. 83 Therefore, positive deviations in the N:P stoichiometries relative to the C:P stoichiometries indicate ratio of nitrite that is re-oxidized rather than further reduced. This ratio can then be applied 84 to determine the degree that calculations of fixed nitrogen loss based on inorganic nitrogen versus 85 phosphate overestimate denitrification. Here, we analyzed the progression of nutrients within the 86 87 three water masses that feed the Eastern Tropical North Pacific Oxygen Deficient Zone to 88 determine the relative importance of nitrite oxidation compared to nitrite reduction.

89

90 Methods

We analyzed Pacific-wide data products spanning wide spatial and temporal ranges.
Gridded data from World Ocean Atlas 2013 (Garcia et al. 2013) were acquired via the
WOA13_1.00deg_1955-2012_Annual file available from the Ocean Data View web portal
(https://odv.awi.de/data/ocean/world-ocean-atlas-2013/). Available temperature, salinity, oxygen,

phosphate, and nitrate, and dissolved inorganic carbon were extracted from the World Ocean 95 Database 2018 (Garcia al. 2013) 96 et using the WODselect tool (https://www.ncei.noaa.gov/access/world-ocean-database-select/dbsearch.html). the 97 Both WOA13 and the WOD18 data were processed and visualized with Ocean Data View version 5.1.7 98 (Schlitzer 2020) and MATLAB R2018b (The MathWorks, Inc. 2018). TEOS-10 (http://www.teos-99 100 10.org/software.htm) (T. J. McDougall and Barker 2011) was used to convert in situ temperature and salinity to conservative temperature and absolute salinity, as well as calculate potential density 101 anomaly, using IBM ILOG CPLEX Optimization Studio V12.8.0 as an optimizer. Nutrient ratios 102 for each water mass were calculated using Type II linear regressions with pre-existing code 103 (Glover et al. 2011). All data were plotted, but outliers were excluded from these linear regressions. 104 With regards to visualizing data, the viridis colormap was selected for scientific use of color and 105 106 accessibility (Crameri et al. 2020), rather than the "odv" option which is used as default.

We subdivided this data into the three water masses that compose the ETNP ODZ using conservative parameters centered around water mass definitions specified in a water mass analysis of this region (Evans et al. 2020). Table 1 provides these specifications, which are generously wide in an effort to include sufficient data that outliers do not interfere with calculations. With these water mass subsets, we analyzed the C:N:P regeneration ratios resulting from non-aerobic processes. The percentages of nitrite re-oxidized were calculated using Eq. (1) or Eq. (2) for WOD18 or WOA13 data, respectively.

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	13CW		NEPIW		AAIW	
	Min Max		Min	Max	Min	Max
θ/°C	12.5	13.5	9	10	5	6
$S_A/g kg^{-1}$	34.8	35.2	34.7	34.85	34.67	34.72
$\sigma_{\theta}/kg m^{-3}$	26.2	26.4	26.7	26.9	27.2	27.3
*Latitude/°N	0	n/a	0	n/a	0	n/a
*NO3 ⁻ /µmol kg ⁻¹	15	n/a	n/a	n/a	n/a	n/a

¹¹⁵ Table 1) Ranges for filtering WOA13 and WOD18 data into coherent water masses. Starred

variables were only used to filter the data for linear regression of nutrient ratios in Fig. 2, not for

117 geographic distributions in Fig. 1.

119
$$NO_{2}^{-} \text{ oxidized } (\%) = 100 \left(\frac{(Total \ CO_{2}: NO_{3}^{-})_{i} - (-1.12)}{(Total \ CO_{2}: NO_{3}^{-})_{i}} \right)$$
(1)

$$NO_{2}^{-} oxidized (\%) = 100 \left(\frac{-94.4 - (NO_{3}^{-}: PO_{4}^{3-})_{i}}{-94.4} \right)$$
(2)

120

122 Source waters to the Pacific Oxygen Deficient Zones

Three water masses compose both the Pacific ODZs, with slight differences between the 123 ETNP and Eastern Tropical South Pacific (ETSP) (Evans et al. 2020). In the ETNP, these water 124 masses are the 13 °C Water (13CW), Northern Equatorial Pacific Intermediate Water (NEPIW), 125 and Antarctic Intermediate Water (AAIW). Fig. 1 depicts the geographical extent of each water 126 mass using subsets of World Ocean Atlas 2013 gridded data (WOA13) based on the temperature 127 and salinity of each water mass, listed in Table 1. The 13CW subset was selected to be slightly 128 129 deeper than that found in Evans et al. (2020). This decision is based off the results of particle backtracking into the ETNP ODZ, which found the entry of this water mass through the southern 130 boundary of the ODZ slightly deeper (Margolskee et al. 2019). In addition, while other references 131 (Bostock et al. 2010) define the center of the NEPIW and its southern counterpart, SEPIW, closer 132 to $\sigma_{\theta}=27$ kg m⁻³, the Northern Equatorial Undercurrent Jets that inject the NEPIW into the ETNP 133 ODZ are centered at σ_{θ} =26.8 kg m⁻³ (Qiu et al. 2013; Margolskee et al. 2019). To represent the 134 waters that feed the ETNP ODZ, we adjusted our subset to be shallower, even though this process 135 136 selects specifically the upper region of these water masses. 137



Fig. 1) Maps of (a) 13CW, (b) NEPIW/SEPIW, and (c) AAIW, depicting PO₄³⁻/µmol kg⁻¹. The contour line depicts the concentration past which anaerobic respiration appears to begin, as given in Fig. 2. Water mass formation sources are noted.

In Fig. 1, the phosphate concentrations of each water mass are used to depict how the water 143 144 masses age as they are transported from their source regions to the Pacific ODZs. The purple contour represents the approximate location where the signal of anaerobic metabolisms can be 145 detected. This concentration is identified in Fig. 2 and described in surrounding text, however these 146 phosphate concentrations are based on the ETNP ODZ and do not reflect the ETSP as well. 147 Nevertheless, these plots confirm that the Pacific ODZs share coherent source water masses, as 148 149 well as visualize the flow paths for these water masses entering the ETNP ODZ. While these source water masses are coherent between Pacific ODZs, there are slight differences in the water masses 150 between each ODZ. For example, the Equatorial Pacific Intermediate Water that consists of 151 NEPIW and SEPIW is formed by the subsurface mixing of primarily Pacific Deep Water (PDW) 152 and AAIW with some NPIW. Since the AAIW has far higher dissolved oxygen concentrations 153 than the PDW, the SEPIW remains oxygenated and therefore the ETSP ODZ is shallower than the 154 ETNP ODZ (Kwiecinski and Babbin 2021). 155

The 13CW enters from the south via the Northern Subsurface Countercurrent (Fiedler and Talley 2006; Margolskee et al. 2019), and references for these paths and source regions are described further in Evans et al. (2020), except the AAIW. The AAIW has several formation regions across the Southern Hemisphere, two of which are displayed in Fig. 1c; however, the geographical extent of this water mass definition does not intersect directly with either of these locations. Instead, the AAIW observed in both the ETNP and ETSP ODZ is likely a form of AAIW that has been mixed with other water masses, likely PDW, as Tasman Sea AAIW has a northwestern mixing location (Qu and Lindstrom 2004; Bostock et al. 2013).

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165 Basin-scale quantification of nitrite re-oxidation

While the ocean's fixed nitrogen budget depends on the relative partitioning of nitrite lost 166 via further denitrification versus recycled via oxidation, measurements of nitrite reactions are 167 particularly challenging. Investigations into nitrite oxidation typically extrapolate results based on 168 169 a small number of cruises with limited spatiotemporal sampling. To overcome these limitations, we analyzed WOA13 data for nitrate:phosphate ratios within the source water masses for the 170 171 ETNP ODZ. If each step of denitrification occurs irreversibly, the nitrate:phosphate ratio should be -94.4:1 for each water mass. However, the 13CW, NEPIW, and AAIW had slopes of -20±1, -172 173 43±1, and -34±3, appropriately. These data are presented in Fig. 2a and Table 2. Since WOA13 did not contain carbon data, we verified that these deviations in the nitrate:phosphate ratio were 174 175 due to nitrate by comparing regressions of both nutrients against total inorganic carbon data from 176 WOD18, presented in Fig. 2c and 2d, appropriately. We found that carbon:phosphate deviated less 177 than 12% from the expected 106:1 ratio and this deviation was positive for the NEPIW and AAIW but not the 13CW. These positive deviations are likely due to carbonate dissolution, since this is 178 179 one of the few processes that increases the carbon:phosphate ratio and it can occur within these deeper waters (Hernandez-Ayon et al. 2019). In contrast, the negative deviation in the 13CW 180 181 occurs due to carbon fixation near the base of the euphotic zone, which the 13CW can mix into. 182 Despite the fact that carbon:phosphate ratios deviated by less than 12%, the carbon:nitrate ratios were -3 ± 0.4 and -3 ± 0.2 for 13CW and NEPIW, appropriately. These ratios of approximately -3183 184 are significantly lower than the -1.12 predicted from reaction stoichiometry (Froelich et al. 1979), indicating that denitrifying a single nitrate requires almost three times the carbon expected. 185





Fig. 2) Evolution of nutrients within the 13CW, NEPIW, and AAIW in the northern hemisphere using (a) NO_3^{-}/μ mol kg⁻¹ vs PO_4^{3-}/μ mol kg⁻¹ from WOA13, (b) NO_3^{-}/μ mol kg⁻¹ vs PO_4^{3-}/μ mol kg⁻¹ from WOD18 for comparison with WOA13, (c) total CO₂/µmol kg⁻¹ vs NO_3^{-}/μ mol kg⁻¹ from WOD18, and (d) total CO₂/µmol kg⁻¹ vs PO_4^{3-}/μ mol kg⁻¹ from WOD18.

Data Source	Water mass	Nutrients	Respiration	Slope
WOA13	13CW	NO ₃ ⁻ :PO ₄ ³⁻	Anaerobic	-20±1
	NEPIW	NO ₃ ⁻ :PO ₄ ³⁻	Anaerobic	-43±1
	AAIW	NO ₃ ⁻ :PO ₄ ³⁻	Anaerobic	-34±3
WOD18	13CW	NO ₃ ⁻ :PO ₄ ³⁻	Anaerobic	-43±4
	NEPIW	NO ₃ ⁻ :PO ₄ ³⁻	Anaerobic	-59±6
	13CW	Total CO ₂ :NO ₃ ⁻	Anaerobic	-3±0.4
	NEPIW	Total CO ₂ :NO ₃ ⁻	Anaerobic	-3±0.2
	13CW	Total CO ₂ :PO ₄ ^{3–}	Aerobic	99±2
	NEPIW	Total CO ₂ :PO ₄ ^{3–}	Aerobic	118±3
	AAIW	Total CO ₂ :PO ₄ ^{3–}	Aerobic	118±4
Froelich et al. (1979)		NO ₃ ⁻ :PO ₄ ³⁻	Anaerobic	-94.4
		Total CO ₂ :NO ₃ ⁻	Anaerobic	-1.12
		Total CO ₂ :PO ₄ ^{3–}	Anaerobic	106

193	Table 2) Ratios of nutrients	depicted in Fig.	1 using	linear fits a	and reaction	stoichiometry	for
194	comparison.						

The approximate phosphate concentrations where anaerobic processes begin is defined as 196 where the nitrate:phosphate stoichiometry deviates from 16:1 in Fig. 2a. These concentrations are 197 plotted as in Fig. 1 to visualize the approximate range where each water mass contains the signal 198 199 of denitrification. Previous analysis of these water masses in Evans et al. (2020) focused on the northern part of the ETNP ODZ, where the 13CW is truly anoxic and facilitates nitrite 200 201 accumulation within the slightly deeper secondary nitrite maximum, whereas the NEPIW 202 interferes with nitrite accumulation, likely due to the fact that it is barely oxic with ~10 nM oxygen (Revsbech et al. 2009). In this northern region of the ETNP ODZ, they found that the AAIW is 203 hypoxic. Contradictory to these results, WOA13 data reveals that denitrification has occurred in 204 205 the NEPIW and even the AAIW. Since the NEPIW is introduced into the north side of the ETNP 206 ODZ (Margolskee et al. 2019), this water mass is likely driven to anoxia as it migrates southeast through the ETNP ODZ. Nitrite data from the Costa Rica Dome (Buchwald et al. 2015; 207 208 Chronopoulou et al. 2017) supports this geographical differentiation in water mass oxygen

availability, as nitrite accumulates far deeper in the south of the ETNP ODZ than the north (Evans 209 et al. 2020). Elevated phosphate concentrations in Fig. 1c indicate that in the AAIW, this 210 211 denitrification signal intensifies within the southern portion of the ETNP ODZ, where the ETNP 212 ODZ is deeper.

We observe the largest nitrate:phosphate slope for the 13CW using the WOA13 dataset. 213 214 The 13CW is the shallowest water mass and it can shoal into the photic zone, enabling microaerophilic production of oxygen (Tiano et al. 2014). Nitrite oxidation is reported to be 215 216 responsible for 40%-80% of the oxygen consumption in the edge of the ETNP ODZ (Beman et al. 2021), and this fact supports the significant fraction of nitrite oxidation observed in the 13CW 217 using the WOA13 dataset. A similar trend can be seen in the nitrate:phosphate slopes for the 13CW 218 and NEPIW in the WOD18 data, though these values are closer to the stoichiometric estimate of -219 220 94.4. The anaerobic portion of the WOD18 data originates solely from cruises on the CLIVAR P18 line. This repeat transect occupies 110 °W and therefore bisects through the middle of the 221 222 ETNP ODZ (Fig. S1), emphasizing the processes within the oxygen deficient layer. The WOA13 gridded data encompasses the ETNP ODZ causing its boundaries to be well-represented in this 223 224 dataset, leading to the higher impact of microaerobic regions on nitrite oxidation.

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Future and global implications

227 We can estimate the percent of nitrite that is re-oxidized in the ETNP by comparing the 228 WOD18 data against the expected -1.12 total inorganic carbon:nitrate ratio and the WOA13 data against the -94.4 nitrate:phosphate ratio. In the case of Eq. 1, which compares carbon:nitrate ratios, 229 230 the difference between the observed -3 and the expected -1.12 was found, then scaled by the observed value to determine the precent difference. This comparison is more accurate than the 231 232 nitrate:phosphate comparison because the carbon:nitrate ratio is less sensitive to artificial 233 adjustment by other metabolisms. Unfortunately, the carbon data in this region is limited, so we compared nitrate:phosphate as well. These comparisons reveal that within the core of the ETNP 234 ODZ, as sampled by the WOD18 data, $66\pm12\%$ of the nitrite produced is re-oxidized. This estimate 235 is similar to a previous study that found 50% nitrite re-oxidation (Buchwald et al. 2015) as well as 236 237 another study that found nitrite oxidation rates exceeded nitrate reduction rates (Babbin et al. 2020). Extending this data to the entire ETNP, presented with WOA13 data in Table 2, reveals that the 238 13CW contains $79\pm7\%$ nitrite oxidation, which is noticeably higher than previous estimates. 239

Interestingly, the nitrate:phosphate slopes of the 13CW and NEPIW differ significantly in the WOD18 dataset, whereas their carbon:nitrate slopes are statistically equivalent. The differences in the carbon:phosphate slopes cause the nitrate:phosphate slopes to differ from the carbon:nitrate slopes for this dataset. Nevertheless, nitrite oxidation is likely highest in the 13CW in the WOA13 because previous studies have also noted that the upper oxycline contains higher measured nitrite oxidation rates than the oxygen deficient layer (Peng et al. 2015), and the WOA13 dataset contains the entire upper oxycline in the ETNP ODZ.

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Data source	Water mass	Nitrite oxidized (%)		
WOD18	13CW	66±12		
	NEPIW	64±8		
WOA13	13CW	79±7		
	NEPIW	54±2		
	AAIW	64±9		

Table 3) Percent of nitrite re-oxidized based on slopes fit in Table 2.

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250 These significant proportions of nitrite re-oxidation indicate that steady-state nitrate and phosphate concentrations cannot indicate the rate or extent of nitrate reduction without a 54%-251 79% correction, depending on location within the ETNP. Inverse modeling of ODZs requires 252 253 nitrite oxidation closely paired with nitrate reduction to maintain realistic concentration and isotope distributions (Martin et al. 2019) or artificially lower remineralization rates to prevent 254 nitrate depletion (Su et al. 2015). Models that simulate fixed nitrogen loss using N* without 255 including prolific and depth-dependent nitrite re-oxidation (Deutsch et al. 2001; Codispoti et al. 256 257 2001; DeVries et al. 2013) likely overestimate the fixed nitrogen loss within marine ODZs. Fu et al. (2018) predicts that ODZs will expand until approximately 2100 then contract, but these 258 259 calculations rely on a fixed stoichiometry and increasing the oxycline area of the 13CW may alter the amount of nitrite denitrified versus recycled. 260

This study is the first effort to use basin-wide data to determine the partitioning of nitrite between reduction and oxidation rather than extrapolating from a subset of points or matching data with models. This approach suffers from significant scatter in water mass data, causing large uncertainties, though these uncertainties may be derived from the fact that the nitrite re-oxidation

varies widely across the ETNP. Ultimately, sparse measurements of both inorganic nitrogen and 265 carbon parameters within Pacific ODZs limit the ability to derive robust conclusions. This large 266 267 scale data manipulation also cannot determine the reducing agent for nitrite oxidation, though oxygen, iodate, and even nitrite dismutation have been proposed (Babbin et al. 2020; Sun et al. 268 2021). We calculate that between 54%-79% of nitrite produced in the ETNP is re-oxidized, with 269 270 64±8% in the core of the ETNP ODZ. We find the highest nitrite oxidation in the shallow 13CW because this water mass contains microaerobic pockets due to the secondary chlorophyll maximum 271 as well as high respiration rates. This analysis demonstrates the significance of nitrite recycling, 272 which exceeds nitrite loss across the ETNP ODZ, and the impact of this recycling on elevating 273 phosphate comparisons. Most importantly, the disconnect between nitrate reduction rates from 274 concentrations of nitrate and phosphate influences how oceanographers and ocean modelers 275 276 represent fixed nitrogen loss in the ocean.

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287 Author contributions

N.E. designed this study, acquired the data, trained J.T., and wrote this manuscript. J.T. did the primary analysis of these data sets. J.W.M. advised throughout and A.D. framed this research within the current state of the field.

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292 Competing interests

293 The authors declare no competing interests.

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423 Supplemental information

WOA13 and WOD18 data needed to be subset into the water masses supplying the ODZ to analyze these nutrient relationships. Table 1 depicts the ranges used to subset these data. Table S1 depicts all the linear fits from this data with additional significant figures. Water masses were subset using the ranges presented in Table 1. All code and data for this project can be found at https://github.com/NatalyaEvans/ETNP_stoich.

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Data Source	Nutrients	Water mass	Respiration	Slope	Error	Intercept	Error
WOA13	NO3-PO4	13CW	Aerobic	15.38	0.13	-4.07	0.23
WOA13	NO3-PO4	13CW	Anaerobic	-19.93	1.41	107.43	2.53
WOA13	NO3-PO4	NEPIW	Aerobic	16.44	0.28	-6.83	0.44
WOA13	NO3-PO4	NEPIW	Anaerobic	-43.33	1.06	151.71	1.76
WOA13	NO3-PO4	AAIW	Aerobic	14.44	0.12	-2.34	0.17
WOA13	NO3-PO4	AAIW	Anaerobic	-34.23	3.11	110.80	4.84
WOA13	SiO4-PO4	13CW	Aerobic	11.92	0.33	-1.09	0.47
WOA13	SiO4-PO4	NEPIW	Aerobic	18.77	0.28	-9.71	0.44
WOA13	SiO4-PO4	AAIW	Aerobic	48.13	0.73	-64.02	1.25
WOD18	NO3-PO4	13CW	Aerobic	16.06	0.36	-3.29	0.50
WOD18	NO3-PO4	13CW	Anaerobic	-43.18	4.03	133.55	6.31
WOD18	NO3-PO4	NEPIW	Aerobic	14.05	0.41	0.12	0.63
WOD18	NO3-PO4	NEPIW	Anaerobic	-58.54	6.10	191.99	10.08
WOD18	NO3-PO4	AAIW	Aerobic	14.29	0.50	-0.80	0.83
WOD18	C-PO4	13CW	Aerobic	98.53	2.43	2012.04	3.55
WOD18	C-PO4	NEPIW	Aerobic	117.89	2.96	1959.06	4.71
WOD18	C-PO4	AAIW	Aerobic	117.92	3.94	1949.92	6.72
WOD18	C-NO3	13CW	Aerobic	6.70	0.18	2019.44	1.20
WOD18	C-NO3	13CW	Anaerobic	-3.35	0.41	2341.27	2.49
WOD18	C-NO3	NEPIW	Aerobic	7.66	0.23	1983.19	1.39
WOD18	C-NO3	NEPIW	Anaerobic	-3.11	0.25	2379.70	1.57
WOD18	C-NO3	AAIW	Aerobic	9.51	0.44	1909.30	2.88
ESM4	NO3-PO4	13CW	Aerobic	14.58	0.01	1.08	0.02
ESM4	NO3-PO4	13CW	Anaerobic	-51	1.0	142.7	1.5
ESM4	NO3-PO4	NEPIW	Aerobic	13.717	0.006	2.160	0.008
ESM4	NO3-PO4	NEPIW	Anaerobic	-70.90	0.53	224.44	0.88
ESM4	NO3-PO4	AAIW	Aerobic	11.502	0.016	6.567	0.027
ESM4	NO3-PO4	AAIW	Anaerobic	-89.82	0.29	332.35	0.53

Table S1) Ratios of nutrients depicted in Fig. 1 using linear fits and reaction stoichiometry forcomparison.

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Carbon data in the Pacific ODZs, is quite limited, unlike WOA13. The map in Fig. S1
depicts where WOD18 data extracted with the WOD18select tool was sampled, and the data in
Fig. 2 originates from overlap in the geographical extent of each water mass (Fig. 1) and the
WOD18 carbon sampling (Fig. S1).



440 Fig. S1) Map of all WOD18 data extracted with the WOD18select tool.