Regionally variable contribution of dissolved organic phosphorus to marine annual net community production

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Abstract

Marine dissolved organic phosphorus (DOP) can serve as an organic nutrient to marine autotrophs, helping to sustain a portion of annual net community production (ANCP). Numerical models of ocean circulation and biogeochemistry have diagnosed the magnitude of this process at regional to global scales but have thus far been validated against DOP observations concentrated within the Atlantic basin. Here we assimilate a new marine DOP dataset with global coverage to optimize an inverse model of the ocean phosphorus cycle to investigate the regionally variable role of marine DOP utilization by autotrophs contributing to ANCP. We find $^{25\%}$ of ANCP accumulates as DOP with a regionally variable pattern ranging from 8 – 50% across nine biomes investigated. Estimated mean surface ocean DOP lifetimes of $^{0.5}$ – 2 years allow for transport of DOP from regions of net production to net consumption in subtropical gyres. Globally, DOP utilization by autotrophs sustains $^{14\%}$ (0.9 Pg C yr⁻¹) of ANCP with regional contributions as large as $^{75\%}$ within the oligotrophic North Atlantic and North Pacific. Shallow export and remineralization of DOP within the ocean subtropics contributes 30 – 80% of phosphate regeneration within the upper thermocline (< 300 m). These shallow isopycnals beneath the subtropical gyres harboring the preponderance of remineralized DOP outcrop near the poleward edge of each gyre, which when combined with subsequent lateral transport equatorward by Ekman convergence, provide a shallow overturning loop retaining phosphorus within the subtropical biome, likely helping to sustain gyre ANCP over multi-annual to decadal timescales.

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14	Key Points
15	• 25% of marine annual net community production (ANCP) accumulates as DOP with a
16	mean surface ocean lifetime of $0.5 - 2$ years
17	• Autotrophic uptake of DOP supplies 14% of global ANCP with regional contributions as
18	large as 80%
19	• Shallow subsurface DOP consumption and surface Ekman convergence create a shallow
20	overturning loop retaining P within the ocean subtropics
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24 Abstract

25 Marine dissolved organic phosphorus (DOP) can serve as an organic nutrient to marine 26 autotrophs, helping to sustain a portion of annual net community production (ANCP). Numerical 27 models of ocean circulation and biogeochemistry have diagnosed the magnitude of this process 28 at regional to global scales but have thus far been validated against DOP observations 29 concentrated within the Atlantic basin. Here we assimilate a new marine DOP dataset with 30 global coverage to optimize an inverse model of the ocean phosphorus cycle to investigate the 31 regionally variable role of marine DOP utilization by autotrophs contributing to ANCP. We find 32 ~25% of ANCP accumulates as DOP with a regionally variable pattern ranging from 8-50%33 across nine biomes investigated. Estimated mean surface ocean DOP lifetimes of $\sim 0.5 - 2$ years 34 allow for transport of DOP from regions of net production to net consumption in subtropical 35 gyres. Globally, DOP utilization by autotrophs sustains $\sim 14\%$ (0.9 Pg C yr⁻¹) of ANCP with 36 regional contributions as large as ~75% within the oligotrophic North Atlantic and North Pacific. 37 Shallow export and remineralization of DOP within the ocean subtropics contributes $\sim 30 - 80\%$ 38 of phosphate regeneration within the upper thermocline (< 300 m). These shallow isopycnals 39 beneath the subtropical gyres harboring the preponderance of remineralized DOP outcrop near 40 the poleward edge of each gyre, which when combined with subsequent lateral transport 41 equatorward by Ekman convergence, provide a shallow overturning loop retaining phosphorus 42 within the subtropical biome, likely helping to sustain gyre ANCP over multi-annual to decadal 43 timescales.

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45 **1. Introduction**

46 Dissolved phosphorus (P) has long been recognized as a globally important 'macro'-47 nutrient that can limit marine biological productivity across various spatial and temporal scales. 48 Decades of research have focused on the biogeochemical cycling and availability of dissolved 49 inorganic phosphorus in seawater, dominated by the phosphate ion (PO₄³⁻) and its associated 50 protonated forms, to both infer and predict spatiotemporal patterns of marine biomass production 51 and its subsequent turnover by heterotrophic microbes. However, in recent years other pools of P 52 present in seawater have increasingly gained appreciation as potential sources of P supporting 53 marine biological productivity including inorganic polyphosphate (Orchard et al., 2010b; Martin 54 et al., 2014; Diaz et al., 2018) and organic phosphorus-containing esters (e.g., Mather et al., 55 2008; Orchard et al., 2010a; Reynolds et al., 2014), and phosphonates (Dyhrman et al., 2006), 56 collectively termed dissolved organic phosphorus (DOP). Many major lineages of marine 57 phytoplankton are known to harbor enzymes capable of accessing these organic forms of P 58 including cyanobacteria (Dyhrman et al., 2006; Su et al., 2007), diatoms (Lin et al., 2013), and 59 other eukaryotic algae (Shaked et al., 2006; Li et al., 2018), with numerous field and laboratory 60 studies having documented the ability of these algal taxa to use DOP to supplement and/or satisfy their cellular P requirements (e.g. Björkman & Karl, 2003; Dyhrman & Ruttenberg, 2006; 61 62 Orchard et al., 2010a).

Near surface waters (< 150 m) of subtropical oceans ($\sim 10 - 40^{\circ}$) perennially exhibit low concentrations of PO₄³⁻, typically on the order of $\sim 10 - 300$ nM (Martiny et al., 2019), with similarly depleted concentrations of nitrate (Eppley et al., 1990). This macro-nutrient limitation of net primary (NPP) productivity results in the global minima of vertically integrated annual NPP rates found in the subtropical ocean (Behrenfeld & Falkowski, 1997; Behrenfeld et al., 2005). The related biogeochemical flux, net community production (NCP), represents the

69 amount of organic matter fixed in excess of total community energy requirements, typically 70 evaluated over annual timescales, and is supported by exogenous inputs of nutrients into the 71 productive euphotic zone (Dugdale & Goering 1967; Eppley & Peterson, 1979). The estimated 72 subtropical to extratropical NCP gradient is less variable than for NPP (Emerson, 2014), 73 however there is a longstanding conundrum in ocean biogeochemistry to explain subtropical 74 ocean NCP rates given estimates of the exogenous inorganic nutrient inputs at these latitudes 75 (Michaels et al., 1994; Gruber et al., 1998; Johnson et al., 2010). 76 Lateral supply of organic nutrients including DOP and dissolved organic nitrogen (DON) 77 to gyres from the more productive equatorial, eastern, and poleward boundaries have been 78 suggested to supplement subtropical ocean nutrient budgets (Roussenov et al., 2006; Charria et 79 al., 2008; Torres-Valdes et al., 2009; Reynolds et al., 2014; Letscher et al., 2016), with additional 80 heterotrophic and/or autotrophic processing of the organic macromolecules involved to liberate 81 bioavailable P and N. This process is best characterized for the North Atlantic subtropical gyre 82 where surface PO_4^{3-} concentrations are at or near the global minima (~1-10 nM; Wu et al., 2000; 83 Martiny et al., 2019), leading to P limitation of NPP and NCP (Ammerman et al., 2003; Lomas et 84 al., 2010), for which lateral supply of DOP may provide a nutritional subsidy to marine 85 phytoplankton (Reynolds et al., 2014; Duhamel et al., 2021). Mather et al. (2008) reported on the 86 latitudinal and longitudinal gradients in DOP, PO4³⁻, and alkaline phosphatase activity (APA), a 87 common enzyme used by marine plankton to access P in DOP across the Atlantic basin, finding 88 strong observational support for the microbial utilization of surface ocean DOP when surface 89 PO_4^{3-} dropped below ~100 – 200 nM. Many other field studies have documented the ability of 90 individual phytoplankton taxa and the community to utilize DOP as a P source sustaining growth

across the North Atlantic and North Pacific subtropical gyres (Bjorkman & Karl, 2003; Casey et
al., 2009; Lomas et al., 2010; Orchard et al., 2010a).

93 To date, regional estimates of the role for lateral DOP supply in sustaining subtropical 94 ocean NCP have come from ocean biogeochemistry/ecosystem models coupled with ocean 95 circulation models that have represented marine DOP cycling, including its utilization by 96 autotrophs. These models suggest that on the order of 30 - 70% of NCP in the subtropical North 97 Atlantic is supported by the lateral, exogenous supply of DOP from the gyre margins (Roussenov 98 et al., 2006; Torres-Valdes et al., 2009; Reynolds et al., 2014). A global modeling study extended 99 estimates for the fraction of subtropical NCP supported by lateral DOP utilization to all 100 subtropical gyre regions, corroborating the particular importance of this process within the North 101 Atlantic (~43% of P needed to explain NCP), while finding a reduced but still significant role for 102 DOP, ranging from 19-31% of the P supply for the South Atlantic, Pacific, and Indian ocean 103 subtropical gyres (Letscher et al., 2016). It should be noted that the representation of the marine 104 DOP cycle in all the above modeling studies was validated against an observational DOP 105 concentration database that was largely limited to the Atlantic Ocean, especially the North 106 Atlantic prior to 2015 (Letscher & Moore, 2015). The meridional and zonal gradients in surface 107 ocean DOP concentrations outside of the Atlantic basin are largely unknown outside of two 108 studies (Abell et al., 2000; Moutin et al., 2008). These observed DOP gradients in the subtropical 109 Pacific generally support the proposed mechanism of lateral DOP supply from the productive 110 margins supporting a portion of NCP in the gyres, however a basin- to regional-scale analysis of 111 DOP support of NCP within the Pacific and Indian Oceans, calibrated against a large-scale 112 characterization of DOP concentrations in these basins, is currently lacking.

113 Here we make use of recently compiled DOP concentration data for the open ocean, the 114 DOPv2021 database (Liang et al., 2021), which has doubled the number of DOP observations 115 compared to its predecessor (Letscher & Moore, 2015), with improved spatial coverage of the 116 Pacific basin and some coverage of the Indian basin for the first time. The new DOP database is 117 used to more accurately assess the importance of DOP cycling for sustaining a portion of NCP 118 within a regional context while also providing the first global estimates of this important 119 biogeochemical flux by inclusion of extra-subtropical regions. The DOP observational data are 120 assimilated into an inverse model of the marine P-cycle coupled to a representation of the 121 contemporary ocean circulation to diagnose the biogeochemical fluxes of the marine DOP cycle, 122 e.g., accumulation from biological productivity, heterotrophic remineralization, and autotrophic 123 utilization, that are most consistent with the spatial and vertical gradients observed for DOP. 124 Output from the inverse model is then used to quantify the regionally variable pattern of DOP-125 supported NCP across the global ocean. Model output is also used to identify and quantify the 126 spatial pattern of exported DOP below the productive layer as it is regenerated within the 127 shallow surfaces of the ocean interior.

128

129 **2.** Methods

2.1 Model Design – This study builds on the equilibrium, steady-state P-cycling model of
Wang et al. (2019) that was used to diagnose spatial patterns in marine nitrogen fixation rates.
The model couples a representation of the time-mean ocean circulation in the form of a tracer
transport matrix, the Ocean Circulation Inverse Model v2 (OCIM2; DeVries & Holzer, 2019),
with a parameterization of the marine biogeochemical fluxes of P, which we term the OCIM Pcycling model. The OCIM2 is a data-constrained ocean circulation model that assimilates

136	observational tracers to estimate the climatological mean (steady-state) circulation and has been
137	employed in numerous biogeochemical studies (e.g., DeVries et al., 2013; DeVries & Deutsch,
138	2014; Weber et al., 2016; John et al., 2020). The OCIM P-cycling model used here predicts 3D
139	distributions of dissolved inorganic phosphorus (DIP) (i.e., PO4 ³⁻ but we use DIP to differentiate
140	model predicted phosphate from observed PO4 ³⁻), particulate organic phosphorus (POP), and
141	'semilabile' DOP (lifetime ~months to years) representative of the time-mean conditions for the
142	contemporary ocean. The version of the model employed in this study improves the
143	representation of the marine DOP cycle by adding a term for autotrophic DOP utilization in the
144	upper ocean while also assimilating observations of marine DOP concentration. Assimilation of
145	observed DOP and PO ₄ ³⁻ concentrations is used to objectively constrain a suite of tunable
146	parameters that govern the representation of NCP, the remineralization of its exported products,
147	and specifically the sources and sinks of DOP within the global marine P-cycle.
148	The \sim 200,000 ocean grid boxes within the OCIM2 model grid are divided into nine
149	biogeochemical regions (Fig. 1) to find the spatially variable set of tunable parameters which
150	optimize the representation of marine DOP cycling regionally across the global ocean. The nine
151	biogeochemical regions are defined using the 0.3 μ M surface PO ₄ ³⁻ isoline from the World
152	Ocean Atlas climatology to separate surface ocean-depleted and -replete nutrient regimes within
153	each of the major ocean basins. The choice of nine regions was selected after also considering
154	the distribution of DOP observational data to distribute the available data approximately evenly
155	across regions (n = \sim 375 – 465 per region). Additional regional configurations were tested
156	totaling eight, six, and four regions, with deteriorating goodness of fit and increasing uncertainty
157	in tunable parameter most probable values with decreasing number of regions.

2.2 Datasets – Climatological mean concentrations of observed PO_4^{3-} concentration 158 159 ([PO4]_{obs}) from the World Ocean Atlas 2013 climatology (Garcia et al., 2013) at a resolution of 160 1° lon \times 1° lat \times 33 vertical layers are triangularly interpolated onto the OCIM2 model grid at 2° $lon \times 2^{\circ}$ lat $\times 24$ vertical-layer resolution. A recent global compilation of DOP concentration 161 162 ([DOP]) measurements (the DOPv2021 database; Liang et al., 2021) is similarly interpolated 163 onto the OCIM2 model grid. The [DOP] database contains ~3800 observations spanning the 164 major global ocean basins except for the Arctic, with 97% of observations located in the upper 165 1000 m of the water column. For the OCIM P-cycling model we simulate 'semilabile' DOP 166 (defined in Section 2.3), for which we subtract a value of 0.05 μ M, the asymptotic deep ocean (> 167 1000 m) [DOP] found in DOPv2021, from all DOP data in DOPv2021 to obtain observed, 168 semilabile [DOP].

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2.3 Model Equations & Tunable Parameters – The governing time tendency mass
balance equations for [DIP], [DOP], and [POP] are as follows:

171
$$\frac{d[\text{DIP}]}{dt} + \mathbf{T}[\text{DIP}] = -\gamma[\text{DIP}] + \kappa_d[\text{DOP}] + \kappa_p[\text{POP}] + \kappa_g([\text{DIP}] - \overline{[PO4]}_{obs}) \quad , \tag{1}$$

172
$$\frac{d[\text{DOP}]}{dt} + \mathbf{T}[\text{DOP}] = \sigma \gamma [\text{DIP}] - \kappa_d [\text{DOP}] - \mu [\text{DOP}] \qquad , \qquad (2)$$

173
$$\frac{d[\text{POP}]}{dt} + \mathbf{F}[\text{POP}] = (1 - \sigma)\gamma[\text{DIP}] - \kappa_p[\text{POP}] + \mu[\text{DOP}] \qquad . \tag{3}$$

174 The sources of DIP include from the heterotrophic remineralization of DOP and POP scaled by

175 rate constants k_d and k_p , respectively, and the primary sink is autotrophic uptake scaled by a

- 176 parameter, γ , which is applied to the top two vertical layers of the model (0 73m),
- 177 corresponding to the OCIM model's euphotic zone. The parameter k_p , governing the
- 178 remineralization rate of POP, is not well constrained by the PO₄³⁻ distribution alone and is a
- 179 prescribed value (30 d)⁻¹ (Wang et al., 2019); k_d is a tunable parameter in the model with units =
- 180 yr⁻¹. Gamma (γ) takes the functional form with respect to grid location (**r**):

181
$$\gamma(\mathbf{r}) \equiv \begin{cases} \alpha \frac{\left[\frac{1}{r_{C:P}} \frac{NPP(\mathbf{r})}{NPP_0}\right]^{\beta}}{\frac{\left[PO4\right]_{obs}(\mathbf{r})}{\left[PO4\right]_{obs}(\mathbf{r})}}, & \text{if } z < z_c, \\ 0, & \text{otherwise}, \end{cases}$$
(4)

where α (units = s⁻¹) and β (unitless) are two tunable parameters to model the autotrophic 182 183 biological uptake rate of DIP after considering information on the spatial distribution of marine 184 NPP and upper ocean [PO4]_{obs} gradients and has been used in previous versions of the OCIM P-185 cycling model (Primeau et al., 2013; Teng et al., 2014; Wang et al., 2019). The input field of 186 vertically-integrated, annual marine net primary productivity, NPP(r), is taken from the output of 187 the Carbon-based Productivity Model using the SeaWiFS ocean color climatology (Behrenfeld et 188 al., 2005) and converted to P units using a variable phosphorus to carbon ratio for marine particle 189 production ($r_{C:P}$) as a linear function of *in situ* [PO4]_{obs}(**r**) concentrations (Galbraith & Martiny, 190 2015) in the World Ocean Atlas climatology. Biological uptake of DIP occurs when $z < z_c$, where 191 $z_c = 73$ m, corresponding to the model's euphotic zone, and set to zero below. NPP₀ and [PO4]_{obs0} are equivalent to 1 mmol C m⁻² s⁻¹ and 1 mmol m⁻³, respectively to ensure that the 192 dimensions of γ are s⁻¹. The total ocean inventory of P is conserved in our equilibrium 193 simulations by inclusion of a small source of DIP with a restoring timescale of $k_g = (10^6 \text{ yr})^{-1}$ 194 195 back to the global mean observed [PO4]_{obs} to account for the small loss of P from the model 196 domain when POP reaches the seafloor. 197 Owing to a general lack of knowledge about the sources and sinks of the refractory

(lifetime ~centuries to millennia) fraction of marine DOP, we have chosen to simulate nonrefractory, 'semilabile', marine DOP which is thought to cycle on monthly to yearly timescales (Karl & Björkman, 2015), while also ignoring 'labile' DOP which cycles on ~hourly timescales. The source of semilabile DOP in the model is as a uniform fraction, σ (a tunable parameter), of autotrophic production of organic matter (γ [DIP]). The two sinks for semilabile DOP include

heterotrophic remineralization with rate constant k_d that occurs throughout the model domain and autotrophic uptake of DOP with rate constant μ , which is applied in the upper two vertical layers of the model. Mu takes the form of the reciprocal of the Monod equation:

206
$$\mu = v \frac{K_s + [PO4]_{obs}}{[PO4]_{obs}}$$
, (5)

207 where v is the minimum semilabile DOP uptake rate (units = yr^{-1}) at elevated *in situ*

208 concentrations of $[PO4]_{obs}$ and K_s (units = μ M) is a tunable parameter describing the half-209 saturation constant for maximal DOP uptake rates at low $[PO4]_{obs}$. Initial simulations with both *v*

and K_s as tunable parameters revealed a large covariance inhibiting the ability to constrain both

211 parameters within acceptable levels of uncertainty. We therefore prescribed v = 0.05 yr⁻¹, the

approximate minimum DOP uptake rates observed in extant field studies (Mather et al., 2008;

The production of POP in the model is from the fraction $(1-\sigma)$ of autotrophic DIP uptake (γ [DIP]) and from the autotrophic uptake of DOP, μ [DOP]. Loss of POP below the surface productive layer (at depths > 73 m) occurs at the timescale of $k_g = (30 \text{ d})^{-1}$. T is the advectiondiffusion operator for tracer ([C]) transport defined as T[C] = $\nabla \cdot (\mathbf{u}[C] - \mathbf{K}\nabla [C])$ taken from the output of the OCIM2 expressed as a tracer transport matrix. F is the vertical flux divergence operator defined as:

220
$$\mathbf{F} = \left[\frac{d}{dz} \left(\frac{z}{z_{73m}}\right)^{-b}\right] \int_{z_{73m}}^{0} (\gamma[\text{DIP}] + \mu[\text{DOP}]) dz \quad , \tag{6}$$

where the vertically integrated flux of POP production from the sum of DIP and DOP uptake
above 73 m is attenuated with depth following a power law function fit with the tunable
parameter, *b*.

In summary, the OCIM P-cycling model assimilates observational PO₄³⁻ and DOP data to constrain a set of tunable parameters; σ , k_d , K_s , α , β , and b; yielding prognostic estimates of the 3D distribution of [DIP], [DOP], and [POP] for the contemporary, steady-state ocean. A unique σ , k_d , and K_s are found for each of the nine biogeochemical regions with a single global value for α , β , and b; thus the total number of tunable parameters is thirty.

229 2.4 Model simulations – Assuming the modern marine P cycle is at steady-state, the time 230 tendency terms in Equations 1-3 disappear and the system of three linear algebraic equations can 231 be solved using direct matrix inversion carried out in Matlab® software. The solution to each 232 equation yields the prognostic 3D distribution of [DIP], [DOP], and [POP] as an implicit 233 function of the model's tunable parameters. The set of tunable parameters, which we define as θ , 234 are optimized using a Bayesian method that minimizes the negative of the log posterior of the 235 likelihood function to estimate the most probable parameter values conditioned on the 236 assimilated PO₄³⁻ and DOP observational data. At the core of the likelihood function is the 237 computed error, e, between model predicted and observed tracer concentrations for both DIP and 238 DOP, which we assume are normally and independently distributed:

239
$$e_{[\text{DIP}],i} = \frac{[\text{DIP}]_{i}(\theta) - [\text{PO4}]_{obs}}{S_{\text{DIP},i}}, \text{ for } i = 1...n_{Pi} \qquad , \qquad (7)$$

240
$$e_{[\text{DOP}],i} = \frac{[\text{DOP}]_i(\theta) - [\text{DOP}]_{obs}}{s_{\text{DOP},i}}, \text{ for } i = 1...n_{Po} \qquad , \qquad (8)$$

where $[PO4]_{obs}$ and $[DOP]_{obs}$ are the observed concentrations of PO_4^{3-} and semilabile DOP (total DOP - 0.05 μ M) from the World Ocean Atlas and DOPv2021 databases, respectively, S_{DIP} and S_{DOP} are the standard deviations of all available observations of each, and n_{Pi} and n_{Po} are the number of independent DIP and DOP observation-model output pairs in the model design. The likelihood function also considers the assigned weighting to the assimilated data, W_{Pi} and W_{Po} , which is computed as the fraction of volume ($V_i/\Sigma V$) at each grid point multiplied by the variance of the DIP and DOP observational data, respectively. The likelihood function is thus:

248
$$L(\theta) = \frac{1}{2} \left(e'_{[\text{DIP}]} W_{Pi} e_{[\text{DIP}]} \right) + \frac{1}{2} \left(e'_{[\text{DOP}]} W_{Po} e_{[\text{DOP}]} \right) \qquad . \tag{9}$$

The negative log posterior of the likelihood function (Eq. 9) is minimized to find the most probable values of the tunable parameters within θ , using the optimization algorithm provided by Matlab®'s fminunc function. To perform efficiently, the algorithm relies on both the gradient, $\nabla_{\theta}L$, and the Hessian, $\nabla_{\theta\theta}L$, of the log posterior with respect to the parameters (Eq. 9), which were coded by hand. Initial values for θ were taken from the global best fit solutions to the previous version of the OCIM P-cycling model (Wang et al., 2019). The optimization generally converged after 30 iterations.

We estimate the uncertainty on each tunable parameter within θ by approximating the posterior probability using a Gaussian centered at θ_p , the most probable values of θ . The posterior covariance matrix for θ_p is given by:

259
$$\Sigma = \langle (\theta - \theta_p)(\theta - \theta_p)' \rangle \approx - \left[\nabla_{\theta \theta} L |_{\theta = \theta_p} \right]^{-1} \qquad (10)$$

All error bars on the parameter values within θ reported in Table 1 represent \pm 1 standard deviation of the posterior after marginalizing out all but the parameter of interest. They are obtained from the square root of the diagonal elements of Σ .

263

264 **3. Results**

265 3.1 Tunable parameters – The most probable values of the thirty tunable parameters 266 within θ_p are presented in Table 1 along with their estimated upper bound (e^+) and lower bound 267 (e^-) uncertainty. Parameter σ , the fraction of P-based net community production that accumulates

268 as semilabile DOP in the productive layer, ranged from 0.08 - 0.50. Accumulating fractions in 269 the three regions covering the Atlantic basin were statistically indistinguishable from each other 270 with a value of ~0.114 ± 0.01. A slightly lower value of $\sigma = 0.080 \pm 0.003$ was found for the 271 Indian Ocean region. Accumulating fractions of DOP were higher in the Pacific Ocean regions, 272 in the narrow range $\sigma = 0.240 - 0.264$. The highest value of σ was found for the Northern region, 273 $\sigma = 0.495 \pm 0.03$. The Southern Ocean exhibited $\sigma = 0.183 \pm 0.004$, intermediate between the 274 Atlantic and Pacific Ocean regions. The relative uncertainty on the nine estimated values of σ is 275 low, ranging from $\sim 2 - 11\%$; mean 5.5%. 276 Parameter k_d represents the heterotrophic remineralization rate of semilabile DOP and has 277 units of yr⁻¹. The inverse of k_d is presented in Table 1, representing the heterotrophic 278 remineralization e-folding timescale expressed in units of years. Timescales for heterotrophic 279 semilabile DOP consumption ranged between 2.0 - 9.7 yr. There is no distinguishable difference 280 in heterotrophic DOP consumption timescales between the Atlantic and Pacific basins which 281 averaged 4.1 ± 0.6 yr across the six relevant regions. Interestingly, the Northern and Southern 282 regions exhibited the minimum and maximum DOP consumption timescales at 2.0 ± 0.1 and 9.8283 \pm 0.4 yr, respectively. Relative uncertainty on the estimated values of k_d is slightly higher, 284 ranging from $\sim 3 - 22\%$; mean = $\sim 7.9\%$. The globally-averaged, mean heterotrophic 285 remineralization timescale of semilabile DOP is 3.8 ± 0.9 yr. Parameter K_s represents the half-286 saturation constant governing the autotrophic uptake rate of semilabile DOP within the 287 productive layer. The most probable values of K_s fell into the narrow range of $0.179 - 0.208 \pm$ 288 0.01 μ M (Table 1). Relative uncertainty on the estimated K_s values ranged from ~4 – 24%; mean 289 11.3%.

290 The most probable values of the global fit constants α , β , and b are presented in Table 1. 291 Parameters α and β , fitting constants for modeling autotrophic DIP uptake while accounting for the spatial pattern of marine NPP, were estimated at values of $\alpha = 1.66 \text{ x } 10^{-4} \pm 9.2 \text{ x } 10^{-7} \text{ s}^{-1}$ and 292 $\beta = 0.0259 \pm 0.0006$, representing a relative uncertainty of 0.55% and 2.4%, respectively. 293 294 Parameter b, the exponent governing POP flux attenuation with depth, was found to have a most 295 probable value of 1.186 ± 0.001 , with a low relative uncertainty of 0.12%. The larger estimated 296 value of b found here as compared to the canonical Martin's b exponent (b = 0.858; Martin et al., 297 1987) expressed in terms of POC flux attenuation represents a more rapid attenuation of POP 298 with depth diagnosed by the OCIM P-cycling model. 299 3.2 [DOP] – The steady-state surface ocean concentration of total [DOP] (simulated 300 semilabile [DOP] + uniform 0.05 µM refractory [DOP]) averaged over the productive layer (0 – 301 73 m) from the OCIM P-cycling model (color mapping) and the [DOP] observations from the 302 DOPv2021 database (colored dots) are presented in Figure 2. Modeled surface ocean [DOP] 303 ranges from $0.03 - 0.6 \mu$ M, with a global mean value of $0.16 \pm 0.08 \mu$ M (± 1 S.D.). The model 304 output and observations both identify the global open ocean maximum in [DOP] = -0.5 - 0.6305 µM within the eastern subtropical to tropical Pacific, largely corresponding to the known regions 306 of eastern boundary upwelling. The model predicts intermediate [DOP] ($\sim 0.25 - 0.4 \mu$ M) within 307 the subarctic North Pacific that are overestimated relative to the observations (Fig. 2). 308 Intermediate modeled [DOP] are also found between $\sim 20^{\circ}N - 20^{\circ}S$ across the Pacific Ocean 309 $(-0.25 - 0.4 \,\mu\text{M})$ and within a similar latitude range in the South Atlantic basin, albeit at lower 310 concentrations ($\sim 0.15 - 0.2 \mu$ M). Both the model and observations identify the global open 311 ocean minimum in [DOP] = $\sim 0.03 - 0.05 \,\mu$ M to lie within the subtropical North Atlantic, 312 especially the western half (Sargasso Sea), which has been well documented (Mather et al.,

2008; Torres-Valdes et al., 2009; Lomas et al., 2010; Reynolds et al., 2014). Modeled surface [DOP] is ~ $0.1 - 0.25 \mu$ M within the polar seas of the Arctic and Southern Oceans. A global ocean and full-depth, pointwise comparison of modeled vs. observed [DOP] yielded a correlation coefficient, r = 0.54, a root mean square error of 0.058 μ M, and an overall negative mean bias of ~-4%. The latter metric implies that the model output underpredicts [DOP] by 4% compared against observations globally, however the variance between comparisons was normally distributed with no systemic regional biases observed.

320 3.3 DOP Source/Sink – The annual, vertically-integrated semilabile DOP production flux [mol P m⁻² yr⁻¹] within the upper 73 m is presented in Figure 3a. Semilabile DOP is produced at 321 322 elevated rates, $\sim 0.015 - 0.03$ mol P m⁻² yr⁻¹, within the Pacific basin eastern boundary upwelling 323 systems and the mid-latitude to subarctic North Pacific and Atlantic basins. The semilabile DOP 324 production flux is similarly elevated within the eastern tropical South Pacific in the region between $\sim 10^{\circ} - 20^{\circ}$ S and extending west to $\sim 140^{\circ}$ W. Within the Atlantic, DOP production 325 326 fluxes are similarly elevated on the eastern side of the basins, approaching rates of 0.01 mol P m⁻ 327 ² yr¹. DOP production fluxes are at a minimum within the subtropical gyre regions of each basin, exhibiting the range $\sim 0.0005 - 0.003$ mol P m⁻² yr⁻¹. The annual, vertically-integrated 328 329 autotrophic uptake rate of semilabile DOP [mmol P m⁻² yr⁻¹] within the upper 73 m is presented 330 in Figure 3b. Autotrophic semilabile DOP utilization ranges from $\sim 0.5 - 8 \text{ mmol P m}^{-2} \text{ yr}^{-1}$ with 331 lower values typically found in the Southern Ocean and within the western half of the subtropical 332 gyres of each basin. Elevated autotrophic DOP utilization rates are found within, and 333 immediately 'downstream' of, the productive ecosystems that border the subtropical gyres: the 334 eastern boundary and tropical upwelling regions as well as the subtropical to mid-latitude 335 transition zone. The Pacific basins exhibit overall higher rates than the Atlantic basin.

336	Autotrophic DOP uptake is similarly elevated in the Sea of Japan in the western Pacific that may
337	be an artefact of not well-represented ocean circulation in this semi-enclosed basin at the 2°
338	spatial resolution of the OCIM model affecting nutrient supply to the Sea. The combined result
339	of the semilabile DOP source minus sink fluxes within the productive layer (DOP production
340	minus combined autotrophic + heterotrophic DOP consumption) coupled with the DOP upper
341	ocean stock are used to calculate the mean lifetime of semilabile DOP within the model euphotic
342	zone (upper 73m; Figure 3c). Semilabile DOP lifetimes are shortest within the subtropical North
343	Atlantic (< 1 yr) and longest within the Southern Ocean (~1.8 yr). Within each major ocean
344	basin, semilabile DOP lifetimes are shortest within the subtropical gyres, on the order of $\sim 0.75 - 0.75$
345	1.25 yr. The fraction of combined autotrophic and heterotrophic DOP consumption within the
346	euphotic zone attributable to autotrophic DOP uptake is presented in Figure 3d. Autotrophs
347	contribute the preponderance (>0.5) of euphotic zone DOP consumption globally, with maximal
348	contributions ($\sim 0.75 - 0.90$) found for the subtropical North Atlantic (especially the Sargasso
349	Sea) and South Atlantic, subtropical North and South Pacific, and the Southern Ocean.
350	3.4 New Production – Vertically-integrated, annual net community production (ANCP) in
351	phosphorus units [mol P m ⁻² yr ⁻¹] computed from the sum of autotrophic DIP (γ [DIP]) and DOP
352	(μ [DOP]) uptake minus POP (k_p [POP]) and DOP (k_d [DOP]) regeneration occurring within the
353	euphotic zone is presented in Figure 4a. ANCP rates are elevated, $\sim 0.02 - 0.04$ mol P m ⁻² yr ⁻¹ ,
354	within the subarctic North Pacific and Atlantic, Southern Ocean, eastern boundary and Arabian
355	Sea upwelling regions, and across the tropical Pacific. Minima are found within the subtropical
356	gyres and Arctic in the range $\sim 0.005 - 0.015$ mol P m ⁻² yr ⁻¹ . ANCP converted to carbon units
357	[mol C m ⁻² yr ⁻¹] using the regionally variable $r_{C:P}$ based on <i>in situ</i> [PO4] _{obs} (Galbraith & Martiny,
358	2015) is presented in Figure 4b. Carbon-based ANCP rates are maximal within eastern boundary

359 upwelling systems and their immediately adjacent 'downstream' regions in the range $3-5 \mod C$ 360 m⁻² yr⁻¹. Minima in carbon-based ANCP are found within the subtropical gyres, the Arctic, and Southern Oceans in the range $0.04 - 1.5 \text{ mol C} \text{ m}^{-2} \text{ yr}^{-1}$. Intermediate ANCP (1.5 - 3 mol C m⁻² 361 362 yr⁻¹) are found within the mid-latitude to subpolar regions of both hemispheres as well as across 363 the equatorial regions. The fraction of DOP-supported ANCP [i.e., μ [DOP] / ANCP] is presented 364 in Figure 4c. Semilabile DOP contributes up to 80% of ANCP within the subtropical North 365 Atlantic with elevated contributions ($\sim 0.25 - 0.70$) also found for the subtropical North and 366 South Pacific and the Arctic Oceans. The minima in DOP-supported ANCP ($\sim 0.02 - 0.10$) are 367 found within the Southern Ocean, mid-latitude North Pacific and North Atlantic, Indian, South 368 Atlantic, and within upwelling systems.

369 3.5 DOP regeneration – The semilabile DOP remineralization flux (accumulating as 370 DIP) from the net heterotrophic remineralization of DOP immediately below the productive layer 371 is presented in Figure 5. Regenerated semilabile DOP vertically-integrated within the 73 - 279 m 372 depth interval (model z layers 3-6) is maximal within the subarctic North Pacific and Atlantic 373 as well as the Arctic, reaching values of $\sim 5 - 10 \text{ mmol P m}^{-2} \text{ yr}^{-1}$ (Fig 5a). Intermediate values $(\sim 2.5 - 4 \text{ mmol P m}^{-2} \text{ yr}^{-1})$ within this depth interval are found below the five subtropical gyre 374 375 regions. Minima in regenerated semilabile DOP within the 73 - 279 m depth interval (< 2 mmol 376 P m⁻² yr⁻¹) are found beneath the equatorial and eastern boundary upwelling systems as well as 377 the Southern Ocean. The fraction of regenerated DIP within the 73 - 279 m depth interval 378 attributable to semilabile DOP remineralization varies from $\sim 0.1 - 0.8$ (Fig. 5b), with the 379 remaining fraction arising from POP remineralization. Semilabile DOP remineralization makes 380 its largest contribution to regenerated DIP within the 73 - 279 m layer beneath the five 381 subtropical gyre regions as well as the subpolar to polar latitudes of the northern hemisphere

with fractions in the range $\sim 0.3 - 0.8$. Of the five subtropical gyre regions, DOP regeneration makes its largest contribution in the North Atlantic followed by the North Pacific, the South Pacific and Indian, followed by the South Atlantic. The fractional contribution of semilabile DOP remineralization to DIP regeneration is at a minimum within the equatorial and eastern boundary upwelling regions as well as the Southern Ocean.

387

388 4. Discussion

389 4.1 Comparison of [DOP] distributions to prior work – The OCIM P-cycling model 390 [DOP] output presented here is the first global data product of DOP concentrations in which the 391 biogeochemical cycle of DOP was the primary tracer optimized via data assimilation of marine 392 DOP observations. The small mean bias of the OCIM P-cycling model [DOP] output, $\sim -4\%$, is 393 an improvement over a prior effort involving a more complex biogeochemical model that 394 included the coupled C, N, P, O₂, Fe, and Si cycles (Letscher et al., 2015b), that achieved a mean 395 [DOP] bias of $\sim +7\%$. The correlation coefficient has also nearly doubled for model-observation 396 pairs between studies, i.e., from r = 0.30, n = 2026 (Letscher et al., 2015b) to r = 0.54, n = 3871397 in the current study. While the prior study did assimilate DOP concentration data to optimize 398 marine DOM cycling, the representation of any particular tracer within the high complexity 399 marine ecosystem-biogeochemical model employed in Letscher et al., [2015b] is subject to trade-400 offs in the necessity to jointly optimize dozens of biogeochemical tracers simultaneously. Likely 401 also contributing to the observed improvement in representation of bulk [DOP] is the significant 402 increase in the available DOP observations assimilated in the current study with the availability 403 of the DOPv2021 database (Liang et al., 2021) which contains double the open ocean [DOP] 404 measurements. The largest difference between the surface ocean [DOP] distribution between the

405 prior and new model data products is an overall near doubling of simulated [DOP] within more 406 productive biomes of the world ocean (e.g., upwelling regions), $\sim 0.4 - 0.5 \mu$ M (this study) vs. 407 $\sim 0.2 - 0.3 \mu$ M, and a near halving of simulated [DOP] within subtropical gyres, $\sim 0.05 - 0.1 \mu$ M 408 (this study) vs. $\sim 0.1 - 0.2 \mu$ M (Fig. 2 (this study); Letscher et al. [2015b]). The simulated Indian 409 Ocean surface [DOP] is $\sim 50\%$ lower in the current study as compared to Letscher et al. [2015b], 410 a region of the world ocean that continues to be under sampled for DOP.

411 The OCIM P-cycling output provides the first global scale picture of the spatial 412 distribution of surface ocean [DOP] that is validated against DOP observations from each of the 413 major ocean basins afforded by the DOPv2021 observational database. Surface [DOP] is at a 414 global minimum within the subtropical North Atlantic as has been previously documented 415 (Mather et al., 2008; Torres-Valdes et al., 2009), however similar surface [DOP] depletions are 416 observed for the other four subtropical gyre regions (Fig. 2). Surface [DOP] is elevated along the 417 eastern flanks of the low latitude ocean basins and the tropical ocean as well as the subarctic 418 North Pacific. These spatial patterns for surface [DOP] share a resemblance to that observed for 419 surface ocean dissolved organic nitrogen concentrations [DON] (Letscher et al., 2013), which 420 could be called that of an 'organic nutrient' (Torres-Valdes et al., 2009; Letscher et al., 2016). 421 This spatial pattern for DON and DOP contrasts with that for dissolved organic carbon 422 concentrations [DOC] in the surface ocean, which are known to exhibit regional maxima within 423 the subtropical gyres resulting from the Ekman convergence of more recalcitrant DOC moieties 424 in surface waters (Hansell et al., 2009; Hansell, 2013). Thus, for the organic nutrients, 425 subtropical Ekman convergence supplies DON and DOP towards the oligotrophic gyres where a 426 combination of heterotrophic and autotrophic consumption releases N and P from their organic 427 macromolecules (e.g., Fig. 3d), resulting in the observed sink of DON and DOP from surface

waters across this gradient. For DOC, this same Ekman convergence contributes to the *accumulation* of surface [DOC] within the oligotrophic ocean, the difference perhaps reflecting
the importance of organic nutrients in otherwise nutrient-impoverished gyres, and/or the more
recalcitrant nature of DOC with respect to heterotrophic consumption in surface waters (Carlson
et al., 2004; Letscher et al., 2015a) as compared to DON and DOP (Fig. 3; Hopkinson & Vallino,
2005; Letscher & Moore, 2015).

434 4.2 Spatial patterns of surface ocean DOP sources and sinks – The spatial patterns of 435 semilabile DOP production, autotrophic uptake, the fraction of euphotic zone DOP consumption 436 attributed to autotrophs, and the resulting estimates of semilabile DOP lifetime within the upper 437 ocean predicted by the OCIM P-cycling model are consistent with the role of DOP as an organic 438 nutrient sustaining a portion of oligotrophic ocean productivity (Fig. 3). The eastern tropical to 439 subtropical Pacific, the mid-latitude and subarctic North Pacific, and to a lesser extent the eastern 440 boundary upwelling systems and mid-latitudes of the Atlantic are identified as the biomes 441 leading to the highest DOP production fluxes in the upper ocean, approaching 15 - 30 mmol P m^{-2} yr⁻¹ (Fig. 3a), which translate to elevated surface [DOP] (> 0.3 μ M) in the same regions (Fig. 442 443 2). Conversely, the subtropical gyre regions of each basin exhibit the lowest DOP production 444 fluxes (Fig. 3a) and lowered surface [DOP] (Fig. 2) relative to neighboring biomes. This pattern, 445 elevated DOP production along the equatorial, eastern, and mid-latitude flanks of the subtropical 446 gyres with depressed DOP production within the subtropical gyres, sets up a lateral gradient in 447 surface ocean [DOP] which, when supplied towards the gyre centers by Ekman convergence, 448 helps support a significant fraction of subtropical gyre net community production (this study; 449 Letscher et al., 2016).

450 Autotrophic uptake of semilabile DOP as a P-source sustaining new production was modeled following a reciprocal Monod function with observed PO_4^{3-} as the controlling factor. 451 The minimum autotrophic uptake rate of semilabile DOP at elevated PO₄³⁻ concentrations was 452 453 prescribed a value of 0.05 yr⁻¹ (mean lifetime of 20 years) estimated from the existing literature 454 of field-based alkaline phosphatase activities, the main enzyme used by phytoplankton to access 455 P from DOP substrates (Mather et al., 2008; Sato et al., 2013; Reynolds et al., 2014). The 456 regionally-resolved, most probable values of the half saturation constant, K_s (µM), governing the 457 concentration of *in situ* PO₄³⁻ below which DOP uptake is stimulated, were found to cluster in 458 the narrow range of 179 – 208 nM (Table 1). The resulting reciprocal Monod function (Eq. 5), 459 using parameters of $v = (20 \text{ yr})^{-1}$ and $K_s = \sim 200 \text{ nM}$, yields an uptake curve in which autotrophic semilabile DOP uptake is $v = (10 \text{ yr})^{-1}$ when DIP = 200 nM, $v = (4 \text{ yr})^{-1}$ when DIP = 50 nM, and 460 $v = (1 \text{ yr})^{-1}$ when DIP = 10 nM. However, depleted surface ocean [PO4]_{obs} is not the only factor 461 462 determining autotrophic DOP uptake patterns, with the pattern of DOP production and its lateral 463 transport away from source regions an important additional control. Autotrophic DOP uptake is 464 greatest in the eastern Pacific, reaching $6 - 8 \text{ mmol P m}^{-2} \text{ yr}^{-1}$ in the eastern tropical North Pacific and $4 - 5 \text{ mmol P m}^{-2} \text{ yr}^{-1}$ in the eastern tropical South Pacific (Fig. 3b) coincident with, 465 466 and immediately 'downstream' of, the regions of greatest DOP production (Fig. 3a). In general, 467 autotrophic DOP uptake fluxes are moderately elevated ($\sim 2.5 - 4 \text{ mmol P m}^{-2} \text{ yr}^{-1}$) within the 468 regions immediately 'downstream' in a surface current-flow sense, from equatorial and eastern 469 boundary upwelling regions, as well as the neighboring mid-latitudes along the equatorward, 470 poleward, and eastern flanks of the subtropical gyres (Fig. 3b). Enhanced autotrophic DOP 471 uptake along gyre margins contributes to reduced estimates of the euphotic zone DOP lifetime 472 within the subtropical ocean which are at their regional minima there with lifetimes of $\leq \sim 1.25$ yr 473 (Fig. 3c). Autotrophic uptake of DOP outweighs heterotrophic DOP consumption nearly 474 everywhere within the euphotic zone (Fig. 3d), and is thus the primary control on DOP lifetime 475 within the upper ocean (< 73 m), especially within the subtropical gyres. Our model also predicts 476 a strong contribution of autotrophic DOP uptake to DOP loss within surface waters of the 477 Southern Ocean (Fig. 3d), which is counterintuitive given this region's elevated PO_4^{3-} . Our estimates of autotrophic DOP uptake fluxes are low there, $< 2 \text{ mmol P m}^{-2} \text{ yr}^{-1}$, as is the 478 479 estimated heterotrophic DOP remineralization rate, lifetime = 9.7 years (Table 1). Thus, for 480 comparisons of two fluxes of comparable magnitude, small changes in the estimates of either 481 would have a large impact on their computed ratio, likely explaining our Southern Ocean result. 482 DOP biogeochemistry is comparatively under studied within the Southern Ocean and could be 483 targeted for future investigation.

484 4.3 Role for DOP in support of ANCP – The OCIM P-cycling model presented here 485 improves on earlier versions (e.g., Primeau et al., 2013; Teng et al., 2014; Wang et al., 2019) by 486 assimilating DOP observational data to objectively constrain the regionally-variable parameters 487 governing its biogeochemical cycling. Thus, the resulting spatial pattern (Fig. 4a) and globally-488 integrated magnitude of the model estimated ANCP flux is the most consistent with the observed 489 spatial patterns of PO₄³⁻ and DOP concentrations, satellite-estimated marine NPP rates, and 490 consideration of variable C:P stoichiometry in marine organic matter cycling. The globally-491 integrated, P-based ANCP rate is estimated to be 5.4 Tmol P yr⁻¹ (167.4 Tg P yr⁻¹). Carbon-based 492 annual net community production obtained from multiplication of P-based ANCP by the 493 spatially variable $r_{C:P}$ stoichiometry of marine production based on *in situ* [PO4]_{obs} (Galbraith & 494 Martiny, 2015) yields an ANCP rate of 6.7 Pg C yr⁻¹. This places the OCIM P-cycling ANCP 495 estimate within the lower end of the literature range of this globally important flux (5 - 14 Pg C)

496	yr ⁻¹) computed from a range of methods (Emerson, 2014; Siegel et al., 2016). The Southern and
497	Northern Oceans combined contribute approximately a third of this ANCP at a ratio of ~1.6:1,
498	respectively (Table 2). The four subtropical gyres of the Pacific and Atlantic contribute $\sim 40\%$ of
499	global ANCP at a Pacific:Atlantic basin ratio of ~1.6:1. The remaining ~27% of global ANCP
500	occurs in the equatorial Pacific, equatorial Atlantic, and Indian Ocean. A notable bias is in the
501	North Atlantic subtropical gyre where the OCIM P-cycling ANCP averages $\sim 0.5 - 1 \text{ mol C m}^{-2}$
502	yr ⁻¹ , ~74 – 87% lower than the mean of observational estimates, 3.8 ± 1.2 mol C m ⁻² yr ⁻¹
503	(summarized in Emerson, 2014).
504	Because the OCIM P-cycling model considers both DIP- and DOP-supported ANCP (Eq.
505	1–3), it is possible to compute the fractional contribution of autotrophic semilabile DOP uptake
506	to ANCP (Table 2). The resulting spatial pattern (Fig. 4c) is largely consistent with that of an
507	organic nutrient discussed above, with DOP supplying $\sim 0.1 - 0.8$ of the P needed to sustain
508	ANCP across the five subtropical gyres. Similar to previous estimates (Torres-Valdes et al.,
509	2009; Letscher et al., 2016), DOP makes its largest fractional contribution to ANCP $(0.3 - 0.8)$
510	within a region across the subtropical North Atlantic and the subtropical North Pacific between
511	$\sim 15 - 30^{\circ}$ N that is elevated towards the western half of each basin (Fig. 4c). Similar elevated
512	fractional contributions of DOP-supported ANCP are observed in the western subtropical South
513	Pacific and South Atlantic and the Indian Ocean subtropical gyre, albeit at reduced levels, $\sim 0.1 - 0.1 - 0.1$
514	0.3. Regionally-averaged, DOP-supported ANCP ranges from 15 – 198 Tg C yr ⁻¹ , representing
515	\sim 7 – 21% of overall ANCP in each region (Table 2). The largest DOP-supported contributions to
516	regional ANCP, ~21%, occur in the North and South Pacific subtropical gyres. Elsewhere,
517	autotrophic DOP uptake supports $\sim 7 - 17\%$ of regionally averaged ANCP. An interesting
518	observation of our model result is that regions of elevated autotrophic DOP uptake fluxes (Fig.

519	3b) do not exactly map onto the regions where DOP makes its largest fractional contribution to
520	ANCP (Fig. 4c). For example, the global maxima in autotrophic DOP uptake are found along the
521	eastern boundary and across the mid- to subarctic latitudes of the North Pacific Ocean, yet
522	fractional contributions of DOP to ANCP are low to intermediate there, $\sim 0.1 - 0.2$ (Fig. 4c).
523	Instead, fractional contributions of DOP-supported ANCP are elevated, $\sim 0.3 - 0.7$, elsewhere in
524	the North Pacific, within the adjacent subtropical gyre margins and gyre core. A similar pattern
525	holds for the other five major ocean basins, albeit at varying levels of DOP-supported ANCP. Of
526	note, the subtropical North Atlantic exhibits intermediate rates of autotrophic DOP uptake, ~ 1
527	-3 mmol P m ⁻² yr ⁻¹ , yet the Sargasso Sea harbors the global maximum in DOP-supported
528	ANCP, $\sim 0.5 - 0.8$. Our globally integrated, autotrophic uptake of DOP flux supplies $\sim 14\%$ of
529	marine export production, or ~0.92 Pg C yr ⁻¹ of the total 6.67 Pg C yr ⁻¹ ANCP. Comparing the
530	subtropical ocean gyre regions, estimates of the DOP-supported ANCP fraction are reduced $\sim 2 -$
531	30% by the OCIM P-cycling model as compared to an earlier study that used a high complexity,
532	full marine ecosystem-biogeochemistry framework (Letscher et al., 2016). Estimates by the latter
533	also included the fraction of ANCP supported indirectly by externally supplied DOP that is
534	subsequently remineralized to PO_4^{3-} within the gyres, a flux that is not diagnosed in the present
535	study, and which likely contributes to the lower estimates of DOP-supported ANCP.
536	4.4 Biogeochemical impacts of DOP remineralization – Because of its observed non-
537	Redfield stoichiometry (Hopkinson & Vallino, 2005; Letscher & Moore, 2015), the fractional

538 contribution of dissolved organic matter (DOM) to apparent oxygen utilization (AOU) within the

539 ocean interior has long been of interest to estimate the spatiotemporal patterns of marine export

540 production, nutrient cycling, and oxygen consumption more accurately (Abell et al., 2000;

541 Hansell & Carlson, 2001; Hansell et al., 2009; Carlson et al., 2010; Emerson, 2014; Fawcett et

542	al., 2018; Letscher & Villareal, 2018). At the global scale and assuming steady-state, this
543	fraction should match the fractional accumulation of ANCP as DOM, which for DOC has been
544	estimated in the range $\sim 0.15 - 0.25$ (Hansell et al., 2009; Letscher et al., 2015b; Romera-Castillo
545	et al., 2016; Roshan & DeVries, 2017). However, this fraction is known to vary with both depth
546	and biome across the ocean with typically higher contributions, $\sim 0.25 - 0.8$, estimated for the
547	shallower depth horizons (< 500 m) beneath the subtropical gyre regions (Abell et al., 2000;
548	Hansell & Carlson, 2001; Carlson et al., 2010; Roshan & DeVries, 2017; Letscher & Villareal,
549	2018). Here for semilabile DOP, its globally-integrated fractional accumulation is estimated at
550	24.9% of ANCP with a matching fractional contribution to DIP regeneration below 73 m. We
551	investigated the semilabile DOP remineralization flux within the shallow interior ocean (73 -
552	279 m) at depth horizons where DOM remineralization contributes most to AOU. Similar to
553	previously identified spatial patterns for DOC remineralization, DOP remineralization between
554	73 - 279 m makes its largest fractional contribution to DIP regeneration beneath the subtropical
555	gyre regions as well as the subpolar and polar latitudes of the northern hemisphere (Fig. 5b), in
556	the range of $\sim 0.4 - 0.8$. These locations correspond to regions of large-scale downwelling and/or
557	minima in water mass 'ideal age', estimated at a few years (DeVries & Deutsch, 2014).
558	Considering the subtropical gyres, this shallow depth horizon $(73 - 279 \text{ m})$ corresponds to
559	isopycnal layers in the range $\sigma_{\theta} = \sim 24.2 - 25.5$ in the Pacific and $\sigma_{\theta} = \sim 25.8 - 26.5$ in the
560	Atlantic (Abell et al., 2000; Letscher & Villareal, 2018) which seasonally outcrop at the
561	poleward edge of the gyres and/or immediately poleward within the mid-latitudes (Letscher et
562	al., 2013). These latitudes exhibit equatorward Ekman flow in the upper ocean, contributing to
563	the subtropical Ekman convergence in each basin. Thus, regenerated DIP resulting from shallow
564	remineralization of DOP in the subtropics that eventually returns to the surface near the poleward

655 edge of the gyre is likely partially retained within the subtropical gyre system as surface ocean 566 DIP and any DOP resulting from autotrophic production it supports is laterally advected towards 567 the gyre centers. This shallow overturning loop between upper thermocline DOM 568 remineralization near gyre centers and surface ocean lateral nutrient supply occurring near the 569 gyre margins operating within the ocean subtropics has been suggested to be one mechanism by 570 which marine nitrogen fixation (Abell et al., 2000), and net community production more 571 generally, can be sustained over multi-annual to decadal timescales (Letscher et al., 2016).

572

573 5. Final Comments

574 This work builds and improves on previous studies which have quantified the importance 575 of autotrophic DOP uptake to contribute to ANCP regionally within the Atlantic basin (e.g., 576 Torres-Valdes et al., 2009) and for the five subtropical gyre regions (Letscher et al., 2016), by 577 providing the first global estimates of autotrophic DOP uptake (Fig. 3b) and its fractional 578 contribution to ANCP (Fig. 4b). These global estimates are produced from an inverse model of 579 marine P cycling that assimilated the largest dataset of marine DOP observations to date, 580 including global basin coverage. We note a decoupling between regions where DOP supports the 581 highest fraction of ANCP versus where the highest rates of autotrophic DOP consumption occur. 582 Consistent with the earlier studies, the subtropical North Atlantic gyre (specifically, the Sargasso 583 Sea) is identified as the region where autotrophic DOP uptake makes its largest fractional contribution to ANCP, consistent with the perennially low PO₄³⁻ concentrations and its vertical 584 585 flux in this region (Lomas et al., 2010; Martiny et al., 2019). The subtropical North Pacific gyre 586 is another region exhibiting elevated rates of autotrophic DOP utilization, consistent with 587 previous work performed at Station ALOHA (Bjorkman & Karl, 2003). However, the model

588 output predicts patterns of enhanced autotrophic DOP utilization with highest absolute rates 589 immediately "downstream" of regions of net DOP production. This model prediction offers 590 opportunities for validation by future field studies, including within the subtropical South 591 Pacific, tropical Pacific, the Indo-Pacific throughflow region, and the western Arctic Ocean. 592 Furthermore, the model predicts minima in autotrophic DOP utilization within recently upwelled 593 waters, the mid-latitude recirculation regions of the surface western boundary currents (e.g., Gulf 594 Stream, Kuroshio), the Southern Ocean, and the Indian Ocean. The latter two regions as well as 595 the Arctic continue to suffer from poor data coverage for DOP which could be targeted for future 596 research. Lastly, recent work has identified a potentially important role for the availability of 597 dissolved iron (Fe) and/or ecosystem Fe stress in determining the biogeochemical context in 598 which marine DOP accumulates or is utilized as an additional autotrophic P source within 599 surface waters (Liang et al., in revision). The OCIM P-cycling model presented here does not 600 explicitly model the linkages between the marine P and Fe cycles, however Fe limitation, as well 601 as nitrogen and/or light stress, are implicitly included in the formulation of the autotrophic DIP 602 uptake term that is a spatially variable function based on the pattern of marine NPP (γ term, Eq. 603 4). Many marine diazotrophs are known to obtain a disproportionately large fraction of their 604 cellular P demand from DOP (Mulholland et al., 2002; Sohm & Capone, 2006; Orchard et al., 605 2010b) and are often co-limited by Fe (Kustka et al., 2002; Mills et al., 2004). Future modeling 606 work could aim to explicitly couple the marine P and Fe cycles to investigate the nexus of iron 607 and DOP supply with diazotrophy contributing to marine ANCP.

608

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- 613

614 **Open Research**

- 615 The recent global compilation of [DOP] measurements (the DOPv2021 database v1) is available
- 616 at the Biological and Chemical Data Management Office (BCO-DMO), doi:10.26008/1912/bco-
- dmo.855139.1, under a Creative Commons Attribution 4.0 license. The World Ocean Atlas 2013
- 618 v2 PO₄³⁻ data are available the National Centers for Environmental Information (NOAA),
- doi.org/10.7289/V5NZ85MT, under a public domain license. Model code is available at GitHub,
- 620 <u>https://github.com/rletscher/GBC_DOP_Letscher</u>, under a GNU General Public License.
- 621

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Region	ID	Parameter	Value	e^+	e⁻
Northern	1	σ	0.495	0.030	0.030
Indian	2	σ	0.080	0.003	0.003
Southern	3	σ	0.183 0.004		0.004
NPSG	4	σ	0.253	0.005	0.005
EqPac	5	σ	0.264	0.008	0.008
SPSG	6	σ	0.249	0.017	0.017
NASG	7	σ	0.121	0.008	0.008
EqAtl	8	σ	0.100	0.011	0.010
SASG	9	σ	0.122	0.012	0.011
Northern	1	$(k_d)^{-1}$	2.032	0.080	0.080
Indian	2	$(k_d)^{-1}$	3.807	0.640	0.548
Southern	3	$(k_d)^{-1}$	9.748	0.412	0.396
NPSG	4	$(k_d)^{-1}$	3.048	0.085	0.085
EqPac	5	$(k_d)^{-1}$	4.711	0.227	0.217
SPSG	6	$(k_d)^{-1}$	4.555	0.167	0.161
NASG	7	$(k_d)^{-1}$	4.109	0.392	0.358
EqAtl	8	$(k_d)^{-1}$	3.660	0.791	0.650
SASG	9	$(k_d)^{-1}$	4.308	0.323	0.300
Northern	1	K_s	0.188	0.017	0.016
Indian	2	K_s	0.205	0.009	0.009
Southern	3	K_s	0.208	0.051	0.051
NPSG	4	K_s	0.179	0.019	0.017
EqPac	5	K_s	0.201	0.020	0.020
SPSG	6	K_s	0.187	0.021	0.019
NASG	7	K_s	0.192	0.047	0.038
EqAtl	8	K_s	0.206 0.014		0.014
SASG	9	K_s	0.195	0.013	0.008
		α	1.66 x 10 ⁻⁴	9.25 x 10 ⁻⁷	9.20 x 10 ⁻⁷
		β	0.0259	0.0006	0.0006
		b	1.186	0.0014	0.0014

898 **Table 1.** The optimized parameter values, collectively ' θ ', including the upper bound (e^+) and

899 lower bound (*e*⁻) estimates of the uncertainty. Units: $\sigma =$ unitless; (*k_d*)⁻¹ = yr; *K_s* = μ M; $\alpha = s^{-1}$; β

900 = unitless; b = unitless.

Region	ID	ANCP	% of total	ANCP- DOP	% of total
		Pg C yr ⁻¹		Pg C yr ⁻¹	
Northern	1	0.87	13.1%	0.150	17.2%
Indian	2	0.72	10.8%	0.071	9.9%
Southern	3	1.36	20.4%	0.096	7.1%
NPSG	4	0.95	14.3%	0.198	20.8%
EqPac	5	0.87	13.1%	0.140	16.0%
SPSG	6	0.71	10.6%	0.149	21.0%
NASG	7	0.56	8.3%	0.062	11.2%
EqAtl	8	0.20	3.0%	0.015	7.7%
SASG	9	0.43	6.5%	0.039	9.0%

902	Table 2. Regional	contributions to	ANCP [$Pg C vr^{-1}$].	% of global	ANCP, DOP	-supported
			L		0)	11

903 ANCP [Pg C yr⁻¹], and % of regional ANCP supported by DOP.



Figure 1. Nine unique biogeochemical regions separated by the 0.3 μM surface ocean phosphate

- 914 contour. Region 1 = Northern; 2 = Indian; 3 = Southern; 4 = North Pacific subtropical gyre
- 915 (NPSG); 5 = equatorial Pacific (EqPac); 6 = South Pacific subtropical gyre (SPSG); 7 = North
- 916 Atlantic subtropical gyre (NASG); 8 = equatorial Atlantic (EqAtl); 9 = South Atlantic
- 917 subtropical gyre (SASG).



Figure 2. Surface ocean bulk [DOP] (μM) predicted by the OCIM P-cycling model output (color
mapping), overlain with observations of [DOP] from the DOPv2021 database (Liang et al., 2021)
(colored circles).





942 Figure 3. Autotrophic sources and sinks contributing to the mean lifetime of semilabile DOP 943 within the euphotic zone (Ez = 0 - 73 m). (a) Ez vertically-integrated semilabile DOP production flux in mol P m⁻² yr⁻¹. (b) Ez vertically-integrated autotrophic semilabile DOP uptake flux in 944 mmol P m⁻² yr⁻¹. (c) Mean lifetime of semilabile DOP (yr) within the Ez resulting from the 945 946 combination of autotrophic DOP production, autotrophic and heterotrophic DOP consumption, 947 and ocean circulation. (d) Fraction of semilabile DOP consumption within the Ez contributed by 948 autotrophic DOP uptake; remainder contributed by heterotrophic DOP remineralization. White 949 isoline in (a) and (b) is the surface ocean $[PO4]_{obs} = 0.3 \mu M$ contour from the World Ocean 950 Atlas.



Figure 4. (a) OCIM P-cycling model predicted annual net community production (ANCP) in phosphorus units, mol P m⁻² yr⁻¹. (b) ANCP in carbon units, mol C m⁻² yr⁻¹, applying the regionally variable $r_{C:P.}$ (c) The fraction of ANCP supported by autotrophic uptake of DOP.



956Figure 5. (a) Vertically-integrated interior ocean DOP remineralization flux accumulating as957DIP within the 73 - 279 m layer, mmol P m⁻² yr⁻¹. (b) The fraction of regenerated DIP arising958from DOP remineralization within the 73 - 279 m layer; remaining fraction arises from POP959remineralization.