

1 **A mechanistic model of an upper bound on oceanic carbon export as a**
2 **function of mixed layer depth and temperature**

3 Zuchuan Li*, Nicolas Cassar

4 Division of Earth and Ocean Sciences, Nicholas School of the Environment, Duke University,
5 Durham, North Carolina, USA

6
7 * Corresponding author: Zuchuan Li (zuchuan.li@duke.edu)

8
9
10 **Key points**

11 1. A mechanistic model of an upper bound on carbon export is developed based on the metabolic
12 balance of photosynthesis and respiration in the oceanic mixed layer

13 2. Using parameters available in the literature, the modeled upper bound envelopes field
14 observations of export production estimated from ²³⁴Th and sediment traps and O₂/Ar-derived net
15 community production

16 3. The model identifies regions of the Southern Ocean where carbon export is likely limited by
17 light during part of the growing season

21 **Abstract**

22 Export production reflects the amount of organic matter transferred from the surface ocean to depth
23 through biological processes. This export is in great part controlled by nutrient and light
24 availability, which are conditioned by mixed layer depth (MLD). In this study, building on
25 Sverdrup's critical depth hypothesis, we derive a mechanistic model of an upper bound on carbon
26 export based on the metabolic balance between photosynthesis and respiration as a function of
27 MLD and temperature. We find that the upper bound is a positively skewed bell-shaped function
28 of MLD. Specifically, the upper bound increases with deepening mixed layers down to a critical
29 depth, beyond which a long tail of decreasing carbon export is associated with increasing
30 heterotrophic activity and decreasing light availability. We also show that in cold regions the upper
31 bound on carbon export decreases with increasing temperature when mixed layers are deep, but
32 increases with temperature when mixed layers are shallow. A meta-analysis shows that our model
33 envelopes field estimates of carbon export from the mixed layer. When compared to satellite export
34 production estimates, our model indicates that export production in some regions of the Southern
35 Ocean, most particularly the Subantarctic Zone, is likely limited by light for a significant portion
36 of the growing season.

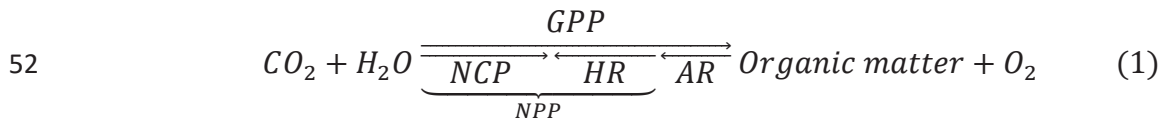
37 **Key words:** Export production, net community production, upper bound, mixed layer depth,
38 temperature

39

40 **1. Introduction**

41 Photosynthesis in excess of respiration at the ocean surface leads to the production of organic
42 matter, part of which is transported to the deep ocean through sinking and mixing (Volk and
43 Hoffert, 1985). This biological process, known as export production (aka soft tissue biological
44 carbon pump) lowers carbon dioxide (CO₂) concentrations at the ocean surface and facilitates the
45 flux of CO₂ from the atmosphere into the ocean (Falkowski et al., 1998; Ito and Follows, 2005;
46 Sigman and Boyle, 2000).

47 Export production is frequently assumed to be a function of net community production (NCP)
48 which is defined as the balance between net primary production (NPP) and heterotrophic
49 respiration (HR), or the difference between gross primary production (GPP) and community
50 respiration (CR; HR plus autotrophic respiration (AR)) (the acronyms used in this study are
51 presented in Table 1) (Li and Cassar, 2016):



53
$$Export\ production = NCP - MLD \times \frac{d(POC + DOC)}{dt} \quad (2)$$

54 where POC, DOC and MLD represent particulate organic carbon, dissolved organic carbon and
55 mixed layer depth, respectively. If the organic carbon inventory (POC+DOC) in the mixed layer
56 is at steady state, NCP is equal to export production (equation (2)). Without allochthonous sources
57 of organic matter, if the organic matter inventory in the mixed layer decreases, NCP will be
58 predicted to be transiently smaller than export production. Conversely, export may lag NPP
59 (Henson et al., 2015; Stange et al., 2017), in which case NCP is expected to be greater than export
60 production.

61 Net community production is in great part regulated by the availability of nutrients and light.
62 Light availability exponentially decays with depth due to absorption by water and its constituents.
63 The mixing of phytoplankton to depth therefore impacts phytoplankton physiology and
64 productivity (Cullen and Lewis, 1988; Lewis et al., 1984), with the depth-integrated NPP expected
65 to increase down to the euphotic depth. Respiration, however, is often modeled to be some function
66 of organic matter concentration, which is expected to be constant with depth if homogeneously
67 mixed within the mixed layer. Temperature is also believed to be an important control on carbon
68 export because respiration is more temperature-sensitive than photosynthesis (Laws et al., 2000;
69 López-Urrutia et al., 2006; Rivkin and Legendre, 2001). Field observations confirm that NCP is
70 generally lower at high temperatures and consistently low when mixed layers are deep. These
71 patterns have been attributed to the balance between depth-integrated photosynthesis (controlled
72 by the availability of nutrients and light) and respiration as a function of MLD and temperature
73 (Cassar et al., 2011; Eveleth et al., 2016; Huang et al., 2012; Shadwick et al., 2015; Tortell et al.,
74 2015). However, descriptions of the underlying mechanisms heretofore remain qualitative.
75 Likewise, the effects of light and nutrient on carbon fluxes are difficult to disentangle. For example,
76 high-nutrient, low-chlorophyll regimes in the Southern Ocean have been attributed to iron
77 limitation (Boyd et al., 2000), deep mixed layers and light limitation (Nelson and Smith, 1991;
78 Mitchell and Holm-Hanse, 1991; Mitchell et al., 1991), or both (Sunda and Huntsman, 1997). To
79 decompose the influence of light and nutrient availability on NCP, we define the upper bound on
80 carbon export from the mixed layer (NCP^*) as the maximum export achievable should all limiting
81 factors other than light (taking into account self-shading) be alleviated.

82 In his seminal paper, Sverdrup presented an elegant model to demonstrate that vernal
83 phytoplankton blooms (i.e., organic matter accumulation at the ocean surface) may be driven by

84 increased light availability when the MLD shoals above a critical depth (Z_c) (Sverdrup, 1953). In
85 our study, we build upon Sverdrup (1953) and derive a mechanistic model of an upper bound on
86 carbon export based on the metabolic balance of photosynthesis and respiration in the oceanic
87 mixed layer, where the metabolic balance is derived from MLD, temperature, photosynthetically
88 active radiation (PAR), phytoplankton maximum growth rate (μ_{max}), and heterotrophic activity.
89 Our approach is analogous to other efforts where mechanistic models were derived to predict
90 proxies of carbon export (e.g., Dunne et al. (2005) and Cael and Follows (2016)). We compare our
91 *NCP** model to observations, and use this model in conjunction with satellite export production
92 estimates to identify regions in the world's oceans where light may limit export production. Our
93 key findings are that 1) using parameters available in the literature, the modeled upper bound
94 envelopes field observations of export production estimated from ^{234}Th and sediment traps and
95 O_2/Ar -derived NCP, and 2) the model identifies regions of the Southern Ocean where carbon
96 export is likely limited by light during part of the growing season.

97 **2. Model description and comparison to observations**

98 **2.1. Net community production and light availability**

99 A conceptual representation of the metabolic balance between volumetric NCP, NPP, and HR
100 profiles is presented in Figure 1(A). According to equation (1), the volumetric NCP flux at a given
101 depth (z) in the mixed layer results from the difference between volumetric NPP and HR:

$$102 \quad NCP(z) = NPP(z) - HR(z) \quad (3)$$

103 where z increases with depth. $NPP(z)$ is a function of the autotroph's intrinsic growth rate
104 (μ) times their biomass concentration (C). Assuming that the effect of nutrients and light on
105 photosynthetic rates abides by Michaelis-Menten kinetics, and neglecting the effect of

106 photoinhibition (Dutkiewicz et al., 2001; Huisman and Weissing, 1994), $NPP(z)$ may be
 107 expressed as follows:

$$108 \quad NPP(z) = \mu(z) \times C = \frac{N}{N + k_m^N} \times \frac{I(z)}{I(z) + k_m^I} \times \mu_{max} \times C \quad (4)$$

109 where μ_{max} is the maximum intrinsic growth rate of the autotrophic community; N and k_m^N
 110 represent the nutrient concentration and half-saturation constant, respectively; and I and k_m^I
 111 represent the irradiance level and half-saturation constant, respectively. μ_{max} , N , k_m^N , k_m^I and C
 112 are assumed to be constant/uniform within the mixed layer. The first two terms on the right-hand
 113 side of equation (4) account for the effect of nutrient and light availability on autotrophic growth
 114 rates, and they are hereafter defined as follows for simplicity:

$$115 \quad N_m = \frac{N}{N + k_m^N} \quad (5a)$$

$$116 \quad I_m(z) = \frac{I(z)}{I(z) + k_m^I} \quad (5b)$$

117 $I(z)$ is modeled as an exponential decay of PAR just beneath the water surface (I_0):

$$118 \quad I(z) = I_0 \times e^{-K_I \times z} \quad (6)$$

119 where K_I is light attenuation coefficient which is assumed to be independent of depth in the mixed
 120 layer.

121 As a first approximation, we assume that $HR(z)$ is proportional to C as in previous studies
 122 (Dutkiewicz et al., 2001; Huisman and Weissing, 1994; Rivkin and Legendre, 2001; Sverdrup,
 123 1953; White et al., 1991):

$$124 \quad HR(z) = r_{HR} \times C \quad (7)$$

125 where r_{HR} represents the intrinsic heterotrophic respiration rate which is assumed to be dependent
 126 on temperature (see below), and independent of depth. In reality, $HR(z)$ is likely best modeled as

127 a function of the concentration of labile organic matter — an additional term could be included to
 128 account for the relationship of total labile organic matter to C .

129 NCP integrated over the mixed layer ($NCP(0, MLD)$) can be derived from equations (3-7):

$$\begin{aligned}
 130 \quad NCP(0, MLD) &= NPP(0, MLD) - HR(0, MLD) \\
 131 \quad &= \int_0^{MLD} NPP(z) dz - \int_0^{MLD} HR(z) dz \\
 132 \quad &= N_m \times I_m(0, MLD) \times \mu_{max} \times C - r_{HR} \times MLD \times C \quad (8)
 \end{aligned}$$

133 The first term on the right side of equation (8) represents NPP integrated over the mixed layer
 134 ($NPP(0, MLD)$), which is equivalent to the product of $\int_0^{MLD} \mu(z) dz$ and C , where the former term
 135 is modeled to be a function of μ_{max} conditioned by nutrient and light availability within the mixed
 136 layer. $I_m(0, MLD)$ can be derived as follows:

$$137 \quad I_m(0, MLD) = \int_0^{MLD} I_m(z) dz = -\frac{1}{K_I} \times \ln \left(\frac{I_0 \times e^{-K_I \times MLD} + k_m^I}{I_0 + k_m^I} \right) \quad (9)$$

138 NCP integrated over the mixed layer (equation (8)) is a bell-shaped function of MLD as depicted
 139 in the schematic diagram of Figure 1(B).

140 2.2. Net community production and phytoplankton biomass concentration

141 As can be seen from equation (8), $NCP(0, MLD)$ is a direct function of C because
 142 $NPP(0, MLD)$ and $HR(0, MLD)$ are proportional to C . $NCP(0, MLD)$ is also an indirect function
 143 of C due its effect on light attenuation (i.e., K_I). The attenuation coefficient K_I can be divided into
 144 water and non-water components ($K_I = K_I^w + K_I^{nw}$) (Baker and Smith, 1982; Smith and Baker,
 145 1978a; Smith and Baker, 1978b), where K_I^{nw} is controlled by the concentrations of phytoplankton,
 146 colored dissolved organic matter (CDOM), and non-algal particles (NAP). In the open ocean where
 147 CDOM and NAP co-vary with phytoplankton (Morel and Prieur, 1977), K_I can be related to C as
 148 follows:

149
$$K_I = K_I^w + k_c \times C \quad (10)$$

150 where k_c is a function of the solar zenith angle, the specific absorption and backscattering
 151 coefficients of phytoplankton, and the relationship between phytoplankton, CDOM, and NAP.
 152 Because pure water and phytoplankton attenuate light, K_I^w and k_c should be greater than zero.

153 To calculate how $NCP(0, MLD)$ varies as a function of C , we examine its first ($\frac{dNCP(0, MLD)}{dC}$)
 154 and second ($\frac{d^2NCP(0, MLD)}{dC^2}$) derivatives with respect to C based on equations (8) and (10):

155
$$\frac{dNCP(0, MLD)}{dC}$$

 156
$$= N_m \times \mu_{max} \times \frac{K_I^w \times I_m(0, MLD) + k_c \times C \times MLD \times I_m(MLD)}{K_I^w + k_c \times C} - r_{HR} \times MLD \quad (11)$$

157
$$\frac{d^2NCP(0, MLD)}{dC^2} = N_m \times k_c \times \frac{\mu_{max}}{K_I}$$

 158
$$\times \left\{ \frac{2 \times K_I^w}{K_I} \times (MLD \times I_m(MLD) - I_m(0, MLD)) - \frac{k_c \times C \times I_m(MLD)^2 \times MLD^2 \times k_m^I}{I_0 \times e^{-K_I \times MLD}} \right\} \quad (12)$$

159 when $MLD > 0$, $I_m(0, MLD) > MLD \times I_m(MLD)$:

160
$$I_m(0, MLD) = \int_0^{MLD} \frac{I_0 \times e^{-K_I \times z}}{I_0 \times e^{-K_I \times z} + k_m^I} dz$$

 161
$$> \int_0^{MLD} \frac{I_0 \times e^{-K_I \times MLD}}{I_0 \times e^{-K_I \times MLD} + k_m^I} dz = MLD \times I_m(MLD) \quad (13)$$

162 The detailed derivation of equations (11-12) can be found in the supplementary material.

163 Substituting the inequality (13) into equation (12) gives $\frac{d^2NCP(0, MLD)}{dC^2} < 0$, which suggests that

164 $\frac{dNCP(0, MLD)}{dC}$ decreases with increasing C . Because increasing C decreases light availability due to

165 shelf-shading, $NPP(0, MLD)$ saturates with increasing C . Thus, $NCP(0, MLD)$ will reach an

166 asymptote of $\lim_{C \rightarrow \infty} \left(\frac{dNCP(0, MLD)}{dC} \right) = -r_{HR} \times MLD < 0$, because $HR(0, MLD)$ linearly increases

167 with increasing C while $NPP(0, MLD)$ plateaus (Figure 2). Additionally, because $NCP(0, MLD)$
 168 must be nil when there is no autotrophic biomass ($NCP(0, MLD)|_{C=0} = 0$), $\lim_{C \rightarrow 0} \left(\frac{dNCP(0, MLD)}{dC} \right)$ must
 169 be greater than zero, otherwise the ecosystem would be net heterotrophic which is unachievable
 170 without an allochthonous source of organic matter. $\lim_{C \rightarrow 0} \left(\frac{dNCP(0, MLD)}{dC} \right) > 0$ and $\lim_{C \rightarrow \infty} \left(\frac{dNCP(0, MLD)}{dC} \right) =$
 171 $-r_{HR} \times MLD < 0$ suggest the existence of $\left. \frac{dNCP(0, MLD)}{dC} \right|_{C=C^*} = 0$ where C^* corresponds to an
 172 autotrophic biomass concentration which maximizes $NCP(0, MLD)$ (i.e., NCP^*).

173 The dependence of $NCP(0, MLD)$ on C can be conceptually understood in the following way.
 174 Given a water column with sufficient nutrients, the critical depth Z_c and compensation depth Z_p
 175 are expected to shoal as C increases. When C is low, $NCP(0, MLD)$ increases with C because of
 176 its greater impact on $NPP(0, MLD)$ than on $HR(0, MLD)$. As C further increases, the increase in
 177 $NPP(0, MLD)$ with C slows because of light attenuation (i.e., K_I). There is therefore a C^* which
 178 maximizes the difference between $NPP(0, MLD)$ and $HR(0, MLD)$ leading to NCP^* (Figure 2).
 179 Beyond this point (C^*), further increasing C will cause self-shading and limit photosynthesis in the
 180 deep part of the mixed layer, as a result decreasing $NCP(0, MLD)$. Beyond a critical biomass (C_c),
 181 the ecosystem becomes net heterotrophic. Without an allochthonous source of organic carbon, this
 182 is only transiently sustainable.

183 2.3. Mixed layer depth and compensation depth

184 By definition, if $NCP(MLD)$ is less than zero (i.e., net heterotrophy at the bottom of the mixed
 185 layer), the MLD must be deeper than Z_p ($MLD > Z_p$) (and vice versa). To determine the sign of
 186 $NCP(MLD)$, we substitute inequality (13) into equation (11). According to the inequality
 187 presented in equation (13), $\frac{K_I^W \times I_m(0, MLD) + k_c \times C \times MLD \times I_m(MLD)}{K_I^W + k_c \times C}$ in equation (11) must be greater than

188 $\frac{K_l^w \times MLD \times I_m(MLD) + k_c \times C \times MLD \times I_m(MLD)}{K_l^w + k_c \times C}$ (which is equal to $MLD \times I_m(MLD)$). After simple

189 rearrangements, the substitution of inequality (13) into equation (11) leads to:

$$190 \quad \frac{dNCP(0, MLD)}{dC}$$

$$191 \quad > MLD \times (N_m \times I_m(MLD) \times \mu_{max} - r_{HR}) = \frac{MLD}{C} \times NCP(MLD) \quad (14)$$

192 The inequality in equation (14) in turn suggests that when $NCP(0, MLD)$ is maximized

193 ($\frac{dNCP(0, MLD)}{dC} = 0$), $NCP(MLD)$ is negative (net heterotrophic) and hence the MLD is deeper than

194 Z_p ($MLD > Z_p$). This counterintuitive result is attributable both to the uneven distribution of light

195 availability in the water column (equation (13)) and to water which absorbs light but does not

196 contribute to biomass accumulation. When the mixed layer is at the Z_p , a slight increase in C will

197 leads to negative $NCP(MLD)$ due to decreasing light availability at the base of mixed layer, but

198 will increase NCP higher in the water column because of the increase in biomass. The increase in

199 NCP in the shallow parts of the mixed layer therefore overcompensates for the net heterotrophy at

200 the bottom of the mixed layer, thus maximizing the depth-integrated NCP. If light were uniformly

201 distributed in the water column (i.e., $I_m(0, MLD) = MLD \times I_m(MLD)$) and if water did not

202 attenuate light ($K_l^w = 0$ in equation (11)), $MLD = Z_p$ would maximize $NCP(0, MLD)$, which is

203 consistent with Huisman and Weissing (1994). We note that in equation (14) the NCP profile

204 ($NCP(z)$) varies with increasing C , which is different from what is conceptually presented in

205 Figure 1. The depth-integrated NCP in Figure 1 maximizes at the compensation depth because the

206 NCP profile ($NCP(z)$) is assumed to be invariant.

207 **2.4. An upper bound on carbon export**

208 Equations (11-13) delineate the conditions for an upper bound on carbon export (NCP^*). In

209 order to simplify the relationship of NCP^* to MLD and temperature, we approximate $I_m(0, MLD)$:

210
$$I_m(0, MLD) = -\frac{1}{K_I} \times \ln \left(1 + \frac{I_0}{I_0 + k_m^I} \times (e^{-K_I \times MLD} - 1) \right)$$

211
$$\approx -\frac{1}{K_I} \times \ln(1 - I_m(0)) \quad (15)$$

212 where $I_m(0) = \frac{I_0}{I_0 + k_m^I}$. Based on equation (15), $NCP(0, MLD)$ in equation (8) can be approximated

213 as:

214
$$NCP(0, MLD) = C \times MLD \times \left(\frac{1}{K_I \times MLD} \times \mu^* - r_{HR} \right) \quad (16)$$

215 where $\mu^* = -\ln(1 - I_m(0)) \times N_m \times \mu_{max}$. To evaluate the approximation accuracy of equation

216 (15), we compare the upper bounds estimated from equation (16) and the original model (equations

217 (8-10)). Our comparison suggests that the approximation of equation (15) is accurate for the

218 estimation of NCP^* under most conditions (Figure 3).

219 We first need to derive the C^* which maximizes $NCP(0, MLD)$ (i.e., NCP^*) in equation (16).

220 C^* can be solved from the first derivative of $NCP(0, MLD)$ in equation (16) with respect to C :

221
$$\left. \frac{dNCP(0, MLD)}{dC} \right|_{NCP(0, MLD)=NCP^*} = \mu^* \times \frac{K_I^W}{(k_c \times C^* + K_I^W)^2} - MLD \times r_{HR} = 0 \quad (17)$$

222

223 and therefore:

224
$$C^* = \frac{1}{k_c} \times \left[-K_I^W + \sqrt{\frac{\mu^* \times K_I^W}{MLD \times r_{HR}}} \right] \quad (18)$$

225 Equation (18) decreases with MLD. As C^* is positive ($C^* \geq 0$) and cannot go to infinity ($C^* \leq$

226 C_{max}^*), MLD should satisfy $MLD_{C_{max}^*} \leq MLD \leq \frac{\mu^*}{r_{HR} \times K_I^W}$, where $MLD_{C_{max}^*}$ represents the MLD

227 corresponding to the maximum achievable autotroph's biomass concentration (C_{max}^*) in the

228 surface ocean. The NCP^* model for $0 \leq MLD < MLD_{C_{max}^*}$ is not discussed here, because we do

229 not have data with very shallow MLD to constrain and evaluate the model. The derivation of the
 230 model is however presented in the supplementary material. Substituting C^* from equation (18) into
 231 equation (16):

$$232 \quad \sqrt{NCP^*} = a_2 \times \sqrt{-\ln(1 - I_m(0))} + a_1 \times \sqrt{MLD} \quad (19)$$

233 where $a_1 = -\sqrt{\frac{K_I^W \times r_{HR}}{k_c}}$ and $a_2 = \sqrt{\frac{N_m \times \mu_{max}}{k_c}}$. Constants a_1 and a_2 are functions of r_{HR} and μ_{max} ,
 234 respectively, which are generally modeled to increase with temperature (T) (Eppley, 1972; Rivkin
 235 and Legendre, 2001):

$$236 \quad \mu_{max} = \mu_{max}^0 \times e^{P_t \times T} \quad (20a)$$

$$237 \quad r_{HR} = r_{HR}^0 \times e^{B_t \times T} \quad (20b)$$

238 where P_t and B_t are constants; and μ_{max}^0 and r_{HR}^0 are maximum growth rate and heterotrophic
 239 respiration ratio for $T = 0$ °C, respectively. P_t is commonly assumed to equal 0.0663 (Eppley,
 240 1972). Substituting equations (20a) and (20b) into equation (19) yields:

$$241 \quad \sqrt{NCP^*} = a_4 \times \sqrt{e^{P_t \times T}} \times \sqrt{-\ln(1 - I_m(0))} + a_3 \times \sqrt{e^{B_t \times T}} \times \sqrt{MLD} \quad (21)$$

$$242 \quad \text{where } a_3 = -\sqrt{\frac{r_{HR}^0 \times K_I^W}{k_c}} \text{ and } a_4 = \sqrt{\frac{\mu_{max}^0 \times N_m}{k_c}}.$$

243 **2.5. Comparison to observations**

244 **2.5.1 Data products**

245 We assess the performance of our modeled upper bound on carbon export using a global dataset
 246 of MLD, PAR, sea surface temperature (SST), O_2/Ar -derived NCP, and export production derived
 247 from sediment traps and ^{234}Th (see supplementary material). MLD was derived from global Argo
 248 profiles (Global Ocean Data Assimilation Experiment; <http://www.usgodae.org/>) and CTD casts
 249 (National Oceanographic Data Center; <https://www.nodc.noaa.gov/>). PAR was downloaded from

250 the NASA ocean color website (<https://oceancolor.gsfc.nasa.gov/>). The NCP estimates are based
 251 on a compilation of O₂/Ar measurements from Li and Cassar (2016), Li et al. (2016), Shadwick et
 252 al. (2015), and Martin et al. (2013). The POC export production estimates were obtained from the
 253 recently compiled dataset of Mouw et al. (2016). These estimates were adjusted to reflect a flux at
 254 the base of mixed layer using the Martin curve of organic carbon attenuation with depth (Martin
 255 et al., 1987). The constants k_c and K_I^w in equation (10) were derived assuming a carbon to
 256 chlorophyll *a* ratio of 90 (Arrigo et al., 2008) and an empirical linear relationship between K_I and
 257 chlorophyll *a* concentration (see Figure S3), calculated based on the NOMAD dataset (Werdell
 258 and Bailey, 2005). k_m^I was set at 4.1 Einstein m⁻² d⁻¹ following Behrenfeld and Falkowski (1997).
 259 In our estimation of the upper bound on carbon export, we set N_m to 1 in the NCP^* calculations.

260 **2.5.2 Results and discussion**

261 Overall, we find that NCP^* calculated using published parameters (Table 2) does a good job
 262 of enveloping carbon export observations reported in the literature (Figure 4(A)). Samples on the
 263 NCP^* envelope (upper bound) are likely regulated by light availability. Conversely, points below
 264 the upper bound may be nutrient limited. As expected, NCP^* increases with μ_{max} and decreases
 265 with r_{HR} . Model parameters $a_1 = -1.78$ and $a_2 = 14.75$ (equation (19)) provide the best fit to the
 266 upper bound of O₂/Ar-NCP as a function of MLD. When compared to parameters available in the
 267 literature (Table 2), we find that the best fit to our modeled upper bound is using μ_{max} and r_{HR} of
 268 1.2 d⁻¹ and 0.2 d⁻¹, respectively. When accounting for the effect of T on μ_{max} and r_{HR} , model
 269 constants $a_3 = -1.53$ and $a_4 = 13.39$ (equation (21)) best fit the upper bound on O₂/Ar-NCP,
 270 SST and MLD observations.

271 Our results show that NCP^* decreases faster with increasing MLD in warmer waters (Figures
 272 4(B) and 4(C)), because the term $a_3 \times \sqrt{e^{Bt \times T}}$ in equation (21) is negative and negatively

273 correlated to T . This temperature effect contributes to part of the relationship between export
 274 production and MLD in Figure 4(A). Interestingly, NCP^* increases with T in colder waters and
 275 shallow mixed layers (Figure 4(C)). This is because NCP^* reflects the balance between
 276 productivity ($a_4 \times \sqrt{e^{P_t \times T}} \times \sqrt{-\ln(1 - I_m(0))}$) and heterotrophic respiration ($a_3 \times \sqrt{e^{B_t \times T}} \times$
 277 \sqrt{MLD}). In a shallow cold mixed layer, the change in productivity with T
 278 ($\frac{d(a_4 \times \sqrt{e^{P_t \times T}} \times \sqrt{-\ln(1 - I_m(0))})}{dT} = \frac{P_t}{2} \times a_4 \times \sqrt{e^{P_t \times T}} \times \sqrt{-\ln(1 - I_m(0))}$) is greater than that of
 279 heterotrophic respiration ($\frac{d(a_3 \times \sqrt{e^{B_t \times T}} \times \sqrt{MLD})}{dT} = \frac{B_t}{2} \times a_3 \times \sqrt{e^{B_t \times T}} \times \sqrt{MLD}$). These results could
 280 explain part of the variability in the relationship between NCP and SST reported in previous studies
 281 (Li and Cassar, 2016). Our NCP^* model does not perform as well in warmer deep mixed layers,
 282 where high variability in export ratio maxima have also been reported (Cael and Follows, 2016).
 283 This may stem from uncertainties in observations, the differing relationship between T , μ_{max} , and
 284 r_{HR} at high temperature, and/or violations of our assumptions (see caveats and limitations).

285 Several recent studies have explored the relationship of NCP to oceanic parameters based on
 286 various statistical approaches (Cassar et al., 2015; Chang et al., 2014; Huang et al., 2012; Li and
 287 Cassar, 2016; Li et al., 2016). Our model can shed some light into the mechanisms driving some
 288 of these patterns. To that end, we substitute equation (9) into equation (8):

$$289 \quad NCP(0, MLD) = C \times MLD \times \left(-\frac{N_m \times \mu_{max}}{K_I \times MLD} \times \ln \left(\frac{I_0 \times e^{-K_I \times MLD} + k_m^I}{I_0 + k_m^I} \right) - r_{HR} \right) \quad (22)$$

290 Rearranging equation (22):

$$291 \quad NCP_B = \frac{NCP(0, MLD)}{C \times MLD} = -\frac{\ln \left(\frac{I_0 \times e^{-K_I \times MLD} + k_m^I}{I_0 + k_m^I} \right)}{I_0 \times (1 - e^{-K_I \times MLD})} \times N_m \times \mu_{max} \times PAR_{ML} - r_{HR} \quad (23)$$

292 where NCP_B is the biomass-normalized volumetric NCP, PAR_{ML} is the average PAR in the mixed

293 layer ($PAR_{ML} = \frac{1-e^{-K_I \times MLD}}{K_I \times MLD} \times I_0$), and $-\frac{\ln\left(\frac{I_0 \times e^{-K_I \times MLD} + k_m^I}{I_0 + k_m^I}\right)}{I_0 \times (1-e^{-K_I \times MLD})} \times N_m \times \mu_{max}$ and $-r_{HR}$ correspond

294 to the slope and offset, respectively. The scatter in the relationship between chlorophyll-

295 normalized volumetric NCP and PAR_{ML} , as reported in previous studies (Bender et al., 2016), can

296 likely be explained by the effect of temperature and the availability of nutrient and light (among

297 other properties) on the slope and offset of equation (23). Equation (22) can also be reorganized to

298 assess how environmental conditions may impact the export ratio (ef):

$$299 \quad ef = \frac{NCP(0, MLD)}{NPP(0, MLD)} = 1 - \frac{K_I \times MLD}{-\ln\left(\frac{I_0 \times e^{-K_I \times MLD} + k_m^I}{I_0 + k_m^I}\right)} \times \frac{1}{N_m} \times \frac{r_{HR}}{\mu_{max}} \quad (24)$$

300 where $\frac{r_{HR}}{\mu_{max}}$ is proportional to $e^{(B_t - P_t) \times T}$. Equation (24) is consistent with multiple studies which

301 predict decreasing ef with increasing temperature (Cael and Follows, 2016; Dunne et al., 2005;

302 Henson et al., 2011; Laws et al., 2000; Li and Cassar, 2016). In fact, equation (5) of Cael and

303 Follows (2016) can easily be derived from equation (24) (see supplementary material). Equation

304 (24) also highlights that a multitude of factors may confound the dependence of ef on temperature

305 (including varying MLD, light attenuation, and availability of nutrient and light). This again may

306 explain some of the conflicting observations recently reported in the literature (e.g., Maiti et al.

307 (2013)), where the effect of temperature may be masked by changes in community composition

308 (Britten et al., 2017; Henson et al., 2015). One therefore needs to account or correct for the

309 multitude of confounding factors when predicting the effect of a given environmental condition

310 (e.g., temperature, mineral ballast, and NPP) on the export ratio.

311 **3. Spatial distribution of the upper bound on carbon export**

312 We estimate the global distribution of the upper bound of carbon export using equation (19)
313 and climatological monthly MLD and PAR. In general, NCP^* is high in low latitudes and low in
314 the North Atlantic and Antarctic Circumpolar Current (ACC) in the Southern Ocean (Figure 5(A)).
315 As expected, this spatial pattern is controlled by MLD (see Figure S1). Satellite-derived estimates
316 of NCP (Li and Cassar, 2016) are approximately 10% of global NCP^* , reflecting the high degree
317 of nutrient limitation in the oceans. We also derive a global NCP^* map using equation (21), and
318 find that the global NCP^* estimate is very sensitive to the temperature dependence of r_{HR} . For
319 example, decreasing the B_t in $r_{HR} = r_{HR}^0 \times e^{B_t \times T}$ from 0.11 to 0.08 (as used in Rivkin and
320 Legendre (2001) and López-Urrutia et al. (2006)) increases the global NCP^* budget by a factor of
321 2.4. Large differences in NCP^* in low-latitudes in great part explain this change. In light of the
322 large uncertainties in the relationship between r_{HR} and T (Cael and Follows, 2016; López-Urrutia
323 et al., 2006), we hereafter only discuss NCP^* estimates derived from equation (19).

324 To estimate how close export production is to its upper bound, we calculate the ratio of export
325 production to NCP^* (f_{pt}). Low f_{pt} regimes represent ecosystems likely regulated by nutrient
326 availability (i.e., ecosystems that have not reached their full export potential based on MLD and
327 surface PAR). As expected, low latitude and subtropical regions have low f_{pt} (Figure 5(B)). High
328 f_{pt} regimes represent ecosystems which have reached their full light potential, and are therefore
329 less likely to respond to nutrient addition because of light limitation (e.g., North Atlantic and ACC
330 (Figure 5(B))). In these regions, especially the subantarctic region, f_{pt} is high in the spring (Figure
331 5(C)) and decreases in the summer (Figure 5(D)), suggesting that export production is likely co-
332 limited by nutrient and light availability. This may in part explain the lower response to iron
333 fertilization in the subantarctic region where substantial increases in surface chlorophyll were only

334 observed in regions with shallower mixed layers (Boyd et al., 2007; Boyd et al., 2000; de Baar et
335 al., 2005).

336 Also shown in Figure 5 are the biological pump efficiency and export ratio ef (panels 5E and
337 5F, respectively). These various proxies reflect different components of the biological pump.
338 Whereas f_{pt} reflects the export potential based on current MLD and light availability, the
339 biological pump efficiency reflects the potential as derived from nutrient distribution in the oceans,
340 estimated from the extent of nutrient removal from the surface ocean (Sarmiento and Gruber, 2006)
341 or the proportion of regenerated nutrients at depth (Ito and Follows, 2005). A revised estimate of
342 the global biological pump efficiency, estimated based on the proportion of regenerated to total
343 nutrients (preformed + regenerated) at depth is around 30-35% (Duteil et al., 2013). The ef ratio
344 however describes how much of production is exported as opposed to recycled in the surface
345 (Dunne et al., 2005). The ultra-oligotrophic subtropical waters have a low export ratio, a strong
346 biological pump efficiency with exhaustion of nutrients at the ocean surface, and therefore have
347 not reached their full light potential (low f_{pt}) because of the strong stratification and nutrient
348 limitation. The seasonal pattern of f_{pt} in the subantarctic region suggests that the low biological
349 pump efficiency is the result of light limitation in the austral spring and nutrient (likely Fe) and
350 light limitation in the austral summer.

351 **4. Caveats and limitations**

352 There are a multitude of uncertainties, simplifications, and approximations in our model and
353 field observations. Among others:

- 354 • In our study, we used a model which builds on Sverdrup's critical depth hypothesis. There
355 are competing hypotheses to explain phytoplankton bloom phenology (timing and
356 intensity), including the "dilution recoupling hypothesis" or "disturbance recovery

357 hypothesis” (Behrenfeld, 2010; Boss and Behrenfeld, 2010) and “critical turbulence
358 hypothesis” (Brody and Lozier, 2015; Huisman et al., 1999; Taylor and Ferrari, 2011). In
359 the case of top-down control, any respiratory grazing loss not accounted for by our loss
360 term would behave as a system not reaching its full light potential (NCP^*). Conversely,
361 any grazing loss associated with export (e.g., rapidly sinking fecal pellets and other
362 zooplankton-mediated export pathways) would minimize respiratory losses thereby
363 bringing NCP closer to its upper bound based on light-availability. These opposing effects
364 are beyond the scope of this study, but could be modeled, especially as we learn more about
365 their impacts on carbon fluxes through new efforts such as NASA’s EXPORTS program
366 (Siegel et al., 2016). See also the point below on mixing vs. mixed layer depth.

- 367 • Phytoplankton biomass concentration (C) may vary with depth in the mixed layer,
368 especially for water columns experiencing varying degrees of turbulent mixing. In addition,
369 MLD is not always the best proxy of light availability with mixing layer in some cases
370 deviating from the mixed layer (Franks, 2015; Huisman et al., 1999). The factors defining
371 the MLD also vary in different oceanic regions.
- 372 • For simplicity, we model the dependence of photosynthesis on irradiance assuming
373 Michaelis-Menten kinetics, which does not account for photoinhibition. More accurate
374 models can be found in other studies (Platt et al., 1980). Due to optional absorption, K_I
375 also varies with depth in the mixed layer. Additionally, the linear relationship between K_I
376 and C is influenced by CDOM, NAP, and other environmental factors (e.g., solar zenith
377 angle) (Gordon, 1989).
- 378 • μ_{max} and r_{HR} are influenced by environmental factors other than temperature, including
379 community structure (Chen and Laws, 2017), and may vary with depth within the mixed

380 layer (Smetacek and Passow, 1990). For these reasons, the equations relating μ_{max} and r_{HR}
381 (i.e., B_t and P_t) to temperature also carry significant uncertainties (Bissinger et al., 2008;
382 Edwards et al., 2016; Kremer et al., 2017; López-Urrutia and Morán, 2007; Rivkin and
383 Legendre, 2001) which impacts our estimates of the upper bound on carbon export,
384 especially in warmer regions. As in other recent studies (Cael and Follows, 2016; Cael et
385 al., 2017; Dutkiewicz et al., 2001; Gong et al., 2015; Gong et al., 2017; Huisman et al.,
386 2006; Taylor and Ferrari, 2011), we model heterotrophic respiration to vary in proportion
387 to phytoplankton concentration. The model could be further improved by explicitly
388 including the concentration of heterotrophs. See point above on the grazing effect on export
389 with regards to r_{HR} .

- 390 • NCP may underestimate export production when accompanied by a decrease in the
391 inventory of organic matter in the mixed layer (see introduction and equation (2)).
- 392 • Our field observations are limited, mostly focusing on the spring and summer seasons, and
393 harbor significant uncertainties. For example, deep mixed layers can bias the O_2/Ar method
394 low if entrainment of deeper waters brings low O_2 into the mixed layer. Descriptions of
395 these uncertainties are presented in other studies (Bender et al., 2011; Cassar et al., 2014;
396 Jonsson et al., 2013).
- 397 • Finally, our study is only relevant to the mixed layer. It does not account for productivity
398 below the mixed layer, which can be important in some regions such as the subtropical
399 ocean.

400 **5. Conclusions**

401 In this study, we derived a mechanistic model of an upper bound on carbon export (NCP^*) based
402 on the metabolic balance between photosynthesis and respiration of the plankton community. The

403 upper bound is a positively skewed bell-shaped function of mixed layer depth (MLD). At low
404 temperatures, the upper bound decreases with temperature if mixed layers are deep, but increases
405 with temperature if mixed layers are shallow. We used this model to derive a global distribution
406 of an upper bound on carbon export as a function of MLD and surface PAR, which shows high
407 values in low latitudes and low values in high latitudes due to deep MLD. To examine how current
408 export production compares to this upper bound in the world's oceans, we calculated the ratio of
409 satellite export production estimates to the upper bound derived by our model. High ratios of export
410 production to *NCP** in the North Atlantic and ACC indicate that export production in these regions
411 is likely co-limited by nutrient and light availability. Overall, our results may explain differences
412 in carbon export measured during past iron fertilization experiments (e.g., subantarctic and polar
413 regions), inform future iron fertilization experiments, help in the development of remotely-sensed
414 carbon export algorithms, and improve predictions of the response of marine ecosystems to a
415 changing climate.

416 **Acknowledgements**

417 We would like to acknowledge NASA GSFC for processing and distributing PAR and SST
418 products (<http://oceancolor.gsfc.nasa.gov/>). Global Argo temperature-salinity profiling floats were
419 downloaded from <http://www.usgodae.org/>. CTD casts were downloaded from National
420 Oceanographic Data Center (<https://www.nodc.noaa.gov/>). N.C. was supported by NSF OPP-
421 1043339. Z.L. was supported by a NASA Earth and Space Science Fellowship (Grant No.
422 NNX13AN85H). The authors thank three anonymous reviewers for their insightful comments.

423 **References**

424 Arrigo, K. R., van Dijken, G. L., and Bushinsky, S.: Primary production in the Southern Ocean,
425 1997-2006, *J. Geophys. Res.*, 113, doi:10.1029/2007JC004551, 2008.

426 Baker, K. S. and Smith, R. C.: Bio-optical classification and model of natural waters .2, *Limnol.*
427 *and Oceanogr.*, 27, 500-509, doi:10.4319/lo.1982.27.3.0500, 1982.

428 Behrenfeld, M. J.: Abandoning Sverdrup's Critical Depth Hypothesis on phytoplankton blooms,
429 *Ecology*, 91, 977–989, doi:10.1890/09-1207.1, 2010.

430 Behrenfeld, M. J. and Falkowski, P. G.: Photosynthetic rates derived from satellite-based
431 chlorophyll concentration, *Limnol. and Oceanogr.*, 42, 1-20,
432 doi:10.4319/lo.1997.42.1.0001, 1997.

433 Bender, M. L., Tilbrook, B., Cassar, N., Jonsson, B. F., Poisson, A., and Trull, T. W.: Ocean
434 productivity south of Australia during spring and summer, *Deep-Sea Res. Pt. I*, 112, 68-78,
435 doi:10.1016/j.dsr.2016.02.018, 2016.

436 Bender, M. L., Kinter, S., Cassar, N., and Wanninkhof, R.: Evaluating gas transfer velocity
437 parameterizations using upper ocean radon distributions, *J. Geophys. Res.*, 116,
438 doi:10.1029/2009JC005805, 2011.

439 Bissinger, J. E., Montagnes, D. J. S., Sharples, J., and Atkinson, D.: Predicting marine
440 phytoplankton maximum growth rates from temperature: Improving on the Eppley curve
441 using quantile regression, *Limnol. Oceanogr.*, 53, 487–493,
442 doi:10.4319/lo.2008.53.2.0487, 2008.

443 Boss, E. and Behrenfeld, M. J.: In situ evaluation of the initiation of the North Atlantic
444 phytoplankton bloom, *Geophys. Res. Lett.*, 37, doi:10.1029/2010GL044174, 2010.

445 Boyd, P. W., Jickells, T., Law, C. S., Blain, S., Boyle, E. A., Buesseler, K. O., Coale, K. H., Cullen,
446 J. J., de Baar, H. J. W., Follows, M., Harvey, M., Lancelot, C., Levasseur, M., Owens, N.
447 P. J., Pollard, R., Rivkin, R. B., Sarmiento, J., Schoemann, V., Smetacek, V., Takeda, S.,
448 Tsuda, A., Turner, S., and Watson, A. J.: Mesoscale iron enrichment experiments 1993-
449 2005: Synthesis and future directions, *Science*, 315, 612-617,
450 doi:10.1126/science.1131669, 2007.

451 Boyd, P. W., Watson, A. J., Law, C. S., Abraham, E. R., Trull, T., Murdoch, R., Bakker, D. C. E.,
452 Bowie, A. R., Buesseler, K. O., Chang, H., Charette, M., Croot, P., Downing, K., Frew, R.,
453 Gall, M., Hadfield, M., Hall, J., Harvey, M., Jameson, G., LaRoche, J., Liddicoat, M., Ling,
454 R., Maldonado, M. T., McKay, R. M., Nodder, S., Pickmere, S., Pridmore, R., Rintoul, S.,
455 Safi, K., Sutton, P., Strzepek, R., Tanneberger, K., Turner, S., Waite, A., and Zeldis, J.: A

456 mesoscale phytoplankton bloom in the polar Southern Ocean stimulated by iron
457 fertilization, *Nature*, 407, 695-702, doi:10.1038/35037500, 2000.

458 Britten, G. L., Wakamatsu, L., and Primeau, F. W.: The temperature-ballast hypothesis explains
459 carbon export efficiency observations in the Southern Ocean, *Geophys. Res. Lett.*, 44,
460 1831-1838, doi:10.1002/2016GL072378, 2017.

461 Brody, S. R. and Lozier, M. S.: Characterizing upper-ocean mixing and its effect on the spring
462 phytoplankton bloom with in situ data, *ICES J. Mar. Sci.*, 72, 1961–1970.
463 doi:10.1093/icesjms/fsv006, 2015

464 Cael B. B., Bisson, K., and Follows, M. J.: How have recent temperature changes affected the
465 efficiency of ocean biological carbon export? *Limnology and Oceanography Letters*, 2,
466 113-118, doi:10.1002/lol2.10042, 2017.

467 Cael, B. B. and Follows, M. J.: On the temperature dependence of oceanic export efficiency,
468 *Geophys. Res. Lett.*, 43, 5170-5175, doi:10.1002/2016GL068877, 2016.

469 Cassar, N., Nevison, C. D., and Manizza, M.: Correcting oceanic O₂/Ar-net community production
470 estimates for vertical mixing using N₂O observations, *Geophys. Res. Lett.*, 41, 8961-8970,
471 doi:10.1002/2014GL062040, 2014.

472 Cassar, N., DiFiore, P. J., Barnett, B. A., Bender, M. L., Bowie, A. R., Tilbrook, B., Petrou, K.,
473 Westwood, K. J., Wright, S. W., and Lefevre, D.: The influence of iron and light on net
474 community production in the Subantarctic and Polar Frontal Zones, *Biogeosciences*, 8,
475 227-237, doi:10.5194/bg-8-227-2011, 2011.

476 Cassar, N., Wright, S. W., Thomson, P. G., Trull, W. T., Westwood, K. J., de Salas, M., Davidson,
477 A., Pearce, I., Davies, D. M., and Matear, R. J.: The relation of mixed-layer net community
478 production to phytoplankton community composition in the Southern Ocean, *Global*
479 *Biogeochem. Cy.*, 29, 446-462, doi:10.1002/2014GB004936, 2015.

480 Chang, C.-H., Johnson, N. C., and Cassar, N.: Neural network-based estimates of Southern Ocean
481 net community production from in situ O₂/Ar and satellite observation: a methodological
482 study, *Biogeosciences*, 11, 3279-3297, doi:10.5194/bg-11-3279-2014, 2014.

483 Chen, B., Laws, E. A.: Is there a difference of temperature sensitivity between marine
484 phytoplankton and heterotrophs? *Limnol. and Oceanogr.*, 62, 806-817,
485 doi:10.1002/lno.10462, 2017.

486 Cullen, J. J. and Lewis, M. R.: The kinetics of algal photoadaptation in the context of vertical
487 mixing, *J. Plankton Res.*, 10, 1039-1063, doi:10.1093/plankt/10.5.1039, 1988.

488 de Barr, J. W. H., Boyd, P. W., Coale, K. H., Landry M. R., Tsuda, A., Assmy, P., Bakker, D. C.
489 E., Bozec, Y., Barber, R. T., Brzezinski, M. A., Buesseler, K. O., Boyé, M., Croot, P. L.,
490 Gervais, F., Gorbunov, M. Y., Harrison, P. J., Hiscock, W. T., Laan, P., Lancelot, C., Law,
491 C. S., Levasseur, M., Marchetti, A., Millero, F. J., Nishioka, J., Nojiri, Y., van Oijen, T.,
492 Riebesell, U., Rijkenberg, M. J. A., Saito, H., Takeda, S., Timmermans, K. R., Veldhuis,
493 M. J. W., Waite, A. M., and Wong, C. S.: Synthesis of iron fertilization experiments: From
494 the Iron age in the Age of Enlightenment, *J. Geophys. Res.*, 110, C09S16,
495 doi:10.1029/2004JC002601, 2005.

496 Dunne, J. P., Armstrong, R. A., Gnanadesikan, A., and Sarmiento, J. L.: Empirical and mechanistic
497 models for the particle export ratio, *Global Biogeochem. Cy.*, 19,
498 doi:10.1029/2004GB002390, 2005.

499 Duteil, O., Koeve, W., Oschlies, A., Bianchi, D., Galbraith, E., Kriest, I., and Matar, R.: A novel
500 estimate of ocean oxygen utilisation points to a reduced rate of respiration in the ocean
501 interior, *Biogeosciences*, 10, 7723-7738, doi:10.5194/bg-10-7723-2013, 2013.

502 Dutkiewicz, S., Follows, M., Marshall, J., and Gregg, W. W.: Interannual variability of
503 phytoplankton abundances in the North Atlantic, *Deep-Sea Res. Pt. II*, 48, 2323-2344,
504 doi:10.1016/S0967-0645(00)00178-8, 2001.

505 Edwards, K. F., Thomas, M. K., Klausmeier, C. A., and Litchman, E.: Phytoplankton growth and
506 the interaction of light and temperature: A synthesis at the species and community
507 level, *Limnol. and Oceanogr.*, 61, 1232–1244, doi:10.1002/lno.10282, 2016.

508 Eppley, R. W.: Temperature and phytoplankton growth in the sea, *Fishery Bulletin*, 70, 1063-1085,
509 1972.

510 Eveleth, R., Cassar, N., Sherrell, R. M., Ducklow, H., Meredith, M., Venables, H., Lin, Y., and Li,
511 Z.: Ice melt influence on summertime net community production along the Western
512 Antarctic Peninsula, *Deep-Sea Res. Pr. II*, 139, 89-102, doi:10.1016/j.dsr2.2016.07.016,
513 2017.

514 Falkowski, P. G., Barber, R. T., and Smetacek, V.: Biogeochemical controls and feedbacks on
515 ocean primary production, *Science*, 281, 200-206, doi:10.1126/science.281.5374.200,
516 1998.

517 Franks, P. J. S.: Has Sverdrup's critical depth hypothesis been tested? Mixed layers vs. turbulent
518 layers, *ICES J. Mar. Sci.*, 72, 1897-1907, doi:10.1093/icesjms/fsu175, 2015.

519 Gong, X., Shi, J., Gao, H. W., and Yao, X. H.: Steady-state solutions for subsurface chlorophyll
520 maximum in stratified water columns with a bell-shaped vertical profile of chlorophyll,
521 *Biogeosciences*, 12, 905-919, doi:10.5194/bg-12-905-2015, 2015.

522 Gong, X., Jiang, W., Wang, L., Gao, H., Boss, E., Yao, X., Kao, S., and Shi, J.: Analytical solution
523 of the nitracline with the evolution of subsurface chlorophyll maximum in stratified water
524 columns, *Biogeosciences*, 14, 2371-2386, doi:10.5194/bg-14-2371-2017, 2017.

525 Gordon, H. R.: Can the Lambert-Beer law be applied to the diffuse attenuation coefficient of ocean
526 water? *Limnol. and Oceanogr.*, 34, 1389-1409, doi:10.4319/lo.1989.34.8.1389, 1989.

527 Henson, S. A., Yool, A., and Sanders, R.: Variability in efficiency of particulate organic carbon
528 export: A model study, *Global Biogeochem. Cy.*, 29, 33-45, doi:10.1002/2014GB004965,
529 2015.

530 Henson, S. A., Sanders, R., Madsen, E., Morris, P. J., Le Moigne, F., and Quartly, G. D.: A reduced
531 estimate of the strength of the ocean's biological carbon pump, *Geophys. Res. Lett.*, 38,
532 L04606, doi:10.1029/2011GL046735, 2011.

533 Huang, K., Ducklow, H., Vernet, M., Cassar, N., and Bender, M. L.: Export production and its
534 regulating factors in the West Antarctica Peninsula region of the Southern Ocean, *Global*
535 *Biogeochem. Cy.*, 26, GB2005, doi:10.1029/2010GB004028, 2012.

536 Huisman, J. and Weissing, F. J.: Light-limited growth and competition for light in well-mixed
537 aquatic environments: An elementary model, *Ecology*, 75, 507-520, doi:10.2307/1939554,
538 1994.

539 Huisman, J., van Oostveen, P., and Weissing, F. J.: Critical depth and critical turbulence: Two
540 different mechanisms for the development of phytoplankton blooms, *Limnol. and*
541 *Oceanogr.*, 44, 1781-1787, doi:10.4319/lo.1999.44.7.1781, 1999.

542 Huisman, J., Thi, N. N. P., Karl, D. M., and Sommeijer B.: Reduced mixing generates oscillations
543 and chaos in the oceanic deep chlorophyll maximum, *Nature*, 439, 322-325,
544 doi:10.1038/nature04245, 2006.

545 Ito, T. and Follows, M. J.: Preformed phosphate, soft tissue pump and atmospheric CO₂, *J. Mar.*
546 *Res.*, 63, 813-839, doi:10.1357/0022240054663231, 2005.

547 Jonsson, B. F., Doney, S. C., Dunne, J., and Bender, M.: Evaluation of the Southern Ocean O₂/Ar-
548 based NCP estimates in a model framework, *J. Geophys. Res.*, 118, 385-399,
549 doi:10.1002/jgrg.20032, 2013.

550 Kremer, C. T., Thomas, M. K., and Litchman, E.: Temperature- and size-scaling of phytoplankton
551 population growth rates: Reconciling the Eppley curve and the metabolic theory of ecology,
552 *Limnol. and Oceanogr.*, 62, 1658-1670, doi:10.1002/lno.10523, 2017.

553 Laws, E. A., Falkowski, P. G., Smith, W. O., Ducklow, H., and McCarthy, J. J.: Temperature
554 effects on export production in the open ocean, *Global Biogeochem. Cy.*, 14, 1231-1246,
555 doi:10.1029/1999GB001229, 2000.

556 Lewis, M. R., Cullen, J. J., and Platt, T.: Relationships between vertical mixing and
557 photoadaptation of phytoplankton: Similarity criteria, *Mar. Ecol. Prog. Ser.*, 15, 141-149,
558 doi:10.3354/meps015141, 1984.

559 Li, Z. and Cassar, N.: Satellite estimates of net community production based on O₂/Ar observations
560 and comparison to other estimates, *Global Biogeochem. Cy.*, 30, 735-752,
561 doi:10.1002/2015GB005314, 2016.

562 Li, Z., Cassar, N., Huang, K., Ducklow, H., and Schofield, O.: Interannual variability in net
563 community production at the Western Antarctic Peninsula region (1997-2014), *J. Geophys.*
564 *Res.*, 121, 4748-4762, doi:10.1002/2015JC011378, 2016.

565 López-Urrutia, A. and Morán, X. A. G.: Resource limitation of bacterial production distorts the
566 temperature dependence of oceanic carbon cycling, *Ecology*, 88, 817-822,
567 doi:10.1890/06-1641, 2007.

568 López-Urrutia, Á., San Martín, E., Harris, R. P., and Irigoien, X.: Scaling the metabolic balance
569 of the oceans, *Proc. Natl Acad. Sci. USA*, 103, 8739-8744, doi:10.1073/pnas.0601137103,
570 2006.

571 Maiti, K., Charette, M. A., Buesseler, K. O., and Kahru M.: An inverse relationship between
572 production and export efficiency in the Southern Ocean, *Geophys. Res. Lett.*, 40, 1557-
573 1561, doi:10.1002/grl.50219, 2013.

574 Martin, J. H., Knauer, G. A., Karl, D. M., and Broenkow, W. W.: VERTEX: carbon cycling in the
575 northeast Pacific, *Deep-Sea Res. Pr. A*, 34, 267-285, doi:10.1016/0198-0149(87)90086-0,
576 1987.

577 Martin, P., Rutgers van der Loeff, M., Cassar, N., Vandromme, P., d'Ovidio, F., Stemman, L.,
578 Rengarajan, R., Soares, M., Gonzalez, H. E., Ebersbach, F., Lampitt, R., Sanders, R.,
579 Barnett, B., Smetacek, V., and Naqvi, S. W. A.: Iron fertilization enhanced net community
580 production but not downward particle flux during the Southern Ocean iron fertilization
581 experiment LOHAFEX, *Global Biogeochem. Cy.*, 27, 871–881, doi:10.1002/gbc.20077,
582 2013.

583 Mitchell, B. G. and Holm-Hansen, O.: Observations and modeling of the Antarctic phytoplankton
584 crop in relation to mixing depth, *Deep-Sea Res. Pr. A*, 38, 981-1007, doi:10.1016/0198-
585 0149(91)90093-U, 1991.

586 Mitchell, B. G., Brody, E. A., Holm-Hansen, O., McClain, C., and Bishop, J.: Light limitation of
587 phytoplankton biomass and macronutrient utilization in the Southern Ocean, *Limnol. and*
588 *Oceanogr.*, 36, 1662-1677, doi:10.4319/lo.1991.36.8.1662, 1991.

589 Morel, A. and Prieur, L.: Analysis of variations in ocean color, *Limnol. and Oceanogr.*, 22, 709-
590 722, doi:10.4319/lo.1977.22.4.0709, 1977.

591 Mouw, C. B., Barnett, A., McKinley, G., Gloege, L., and Pilcher, D.: Global ocean particulate
592 organic carbon flux merged with satellite parameters, *Earth Syst. Sci. Data*, 8, 531-541,
593 doi:10.5194/essd-8-531-2016, 2016.

594 Nelson, D. M. and Smith, W. O.: Sverdrup revisited: Critical depths, maximum chlorophyll levels,
595 and the control of Southern Ocean productivity by the irradiance-mixing regime, *Limnol.*
596 *and Oceanogr.*, 36, 1650-1661, doi:10.4319/lo.1991.36.8.1650, 1991.

597 Platt, T., Gallegos, C. L., and Harrison, W. G.: Photoinhibition of photosynthesis in natural
598 assemblages of marine phytoplankton, *J. Mar. Res.*, 38, 687-701, 1980.

599 Rivkin, R. B. and Legendre, L.: Biogenic carbon cycling in the upper ocean: Effects of microbial
600 respiration, *Science*, 291, 2398-2400, doi:10.1126/science.291.5512.2398, 2001.

601 Sarmiento, J. L. and Gruber, N.: *Ocean Biogeochemical Dynamics*, Princeton University Press,
602 Princeton, New Jersey, 2006.

603 Shadwick, E. H., Tilbrook, B., Cassar, N., Trull, T. W., and Rintoul, S. R.: Summertime physical
604 and biological controls on O₂ and CO₂ in the Australian Sector of the Southern Ocean, *J.*
605 *Marine Syst.*, 147, 21-28, doi:10.1016/j.jmarsys.2013.12.008, 2015.

606 Siegel, D. A., Buesseler, K. O., Behrenfeld, M. J., Benitez-Nelson, C. R., Boss, E., Brzezinski, M.
607 A., Burd, A., Carlson, C. A., D'Asaro, E. A., Doney, S. C., Perry, M. J., Stanley, R. H. R.,

608 and Steinberg, D. K.: Prediction of the export and fate of global ocean net primary
609 production: The exports science plan, *Front. Mar. Sci.*, 3, doi:10.3389/fmars.2016.00022,
610 2016.

611 Sigman, D. M. and Boyle, E. A.: Glacial/interglacial variations in atmospheric carbon dioxide,
612 *Nature*, 407, 859-869, doi:10.1038/35038000, 2000.

613 Smetacek, V. and Passow, U.: Spring bloom initiation and Sverdrup's critical depth model, *Limnol.*
614 *and Oceanogr.*, 35, 228–234, doi:10.4319/lo.1990.35.1.0228, 1990.

615 Smith, R. C. and Baker, K. S.: Optical classification of natural waters, *Limnol. and Oceanogr.*, 23,
616 260-267, doi:10.4319/lo.1978.23.2.0260, 1978a.

617 Smith, R. C. and Baker, K. S.: The bio-optical state of ocean waters and remote sensing, *Limnol.*
618 *and Oceanogr.*, 23, 247-259, doi:10.4319/lo.1978.23.2.0247, 1978b.

619 Stange, P., Bach, L. T., Le Moigne, F. A. C., Taucher, J., Boxhammer, T., and Riebesell, U.:
620 Quantifying the time lag between organic matter production and export in the surface ocean:
621 Implications for estimates of export efficiency, *Geophys. Res. Lett.*, 44, 268-276,
622 doi:10.1002/2016GL07087, 2017.

623 Sunda, W. G. and Huntsman, S. A.: Interrelated influence of iron, light and cell size on marine
624 phytoplankton growth, *Nature*, 390, 389-392, doi:10.1038/37093, 1997.

625 Sverdrup, H. U., On conditions for the vernal blooming of phytoplankton, *Journal du Conseil*
626 *International pour l'Exploration de la Mer*, 18, 287-295, doi:10.1093/icesjms/18.3.287,
627 1953.

628 Taylor, J. R. and Ferrari, R.: Shutdown of turbulent convection as a new criterion for the onset of
629 spring phytoplankton blooms, *Limnol. And Oceanogr.*, 56, 2293-
630 2307, doi:10.4319/lo.2011.56.6.2293, 2011.

631 Tortell, P. D., Bittig, H. C., Körtzinger, A., Jones, E. M., and Hoppema, M.: Biological and
632 physical controls on N₂, O₂, and CO₂ distributions in contrasting Southern Ocean surface
633 waters, *Global Biogeochem. Cy.*, 29, 994-1013, doi:10.1002/2014GB004975, 2015.

634 Volk, T. and Hoffert, M. I.: Ocean carbon pumps: Analysis of relative strengths and efficiencies
635 in ocean-driven atmospheric CO₂ changes, in: *The Carbon Cycle and Atmospheric CO₂:
636 Natural Variations Archean to Present*, *Geophys. Monogr. Ser.*, Vol. 32, edited by
637 Sundquist, E. T. and Broecker, W. S., AGU, Washington, D. C., 99-110, 1985.

638 Werdell, P. J. and Bailey, S. W.: An improved in-situ bio-optical data set for ocean color algorithm
639 development and satellite data product validation, *Remote Sensing of Environment*, 98,
640 122-140, doi:10.1016/j.rse.2005.07.001, 2005.

641 White, P. A., Kalff, J., Rasmussen, J. B., and Gasol, J. M.: The effect of temperature and algal
642 biomass on bacterial production and specific growth rate in fresh water and marine habitats,
643 *Microb. Ecol.*, 21, 99-118, doi:10.1007/BF02539147, 1991.

644

645

646 **Table 1.** Model symbols, abbreviations, and units.

647
648 **Table 2.** Value or range of values with references for the parameters used in the model.

649
650 **Figure 1.** Schematic diagram of depth-profiles of net community production (NCP), net primary
651 production (NPP), and heterotrophic respiration (HR). Yellow and black dots represent the
652 compensation and critical depths, respectively.

653
654 **Figure 2.** Relationship between net primary production (NPP), heterotrophic respiration (HR),
655 net community production (NCP), and phytoplankton biomass concentration (C) for a given
656 mixed layer depth (MLD). Hatched area in panel A represents NCP. The yellow dot represents
657 the maximal NCP (NCP*) obtainable for a given MLD, with the corresponding phytoplankton
658 biomass concentration (C*) denoted with a cyan dot. NCP on the right of the yellow dot
659 decreases with C due to self-shading. Black dot represents depth-integrated NCP =0 (i.e.,
660 NPP=HR), with the corresponding phytoplankton biomass concentration defined as critical
661 biomass (C_c) and denoted with a blue dot. Ecosystems on the left and right of this threshold are
662 net autotrophic and heterotrophic, respectively. The asymptote (dashed blue line) in panel B
663 represents a system dominated by heterotrophic respiration (i.e., NCP ≈ HR >> NPP).

664
665 **Figure 3.** Upper bounds derived using the original and approximated models. The upper bound
666 for the original model (equations (8-10)) is estimated through a non-linear optimization
667 approach. The upper bound for the approximated model is calculated analytically from equation
668 (19). The models use the constants listed in Table 2 and $I_m(0) = 0.9$. Decreasing $I_m(0)$ and
669 increasing r_{HR} results in greater discrepancies between the original and approximated models in
670 regions with shallow mixed layers.

671
672 **Figure 4.** Modeled upper bound on carbon export production compared to field observations as a
673 function of mixed layer depth (MLD) and sea surface temperature (SST). (A) The thick gray line
674 represents the upper bound fitted to the net community production (NCP) data. Dash-lines
675 represent the upper bounds calculated using parameters available in the literature (Table 2). (B)
676 NCP as a function of SST with isopleths of constant upper bounds color coded for MLD. NCP
677 observations are color coded with MLD. (C) Surface representing the envelope of the modeled
678 upper bound of carbon export production as a function of SST and MLD. Bars represent field
679 observations color coded with the ratio of NCP to the upper bound. Observations are based on
680 ²³⁴Th and sediment traps estimates of carbon export production and O₂/Ar-derived NCP. A
681 stoichiometric ratio of O₂/C=1.4 was used to convert NCP from O₂ to C units (Laws, 1991). To
682 account for the effect of PAR on export production, both MLD and carbon fluxes are normalized
683 to $-\log(1 - I_m(0))$ (see equations (19) and (21)). The temperature dependence of r_{HR} was
684 modeled as $r_{HR} = r_{HR}^0 \times e^{0.08 \times T}$.

685
686 **Figure 5.** (A) Modeled upper bound on carbon export derived from equation (19), (B-D) ratios
687 of satellite export production estimates to the upper bound on carbon export, (E) biological pump
688 efficiency calculated as the difference in nutrient concentrations between surface and depth,
689 normalized to nutrient concentrations at depth (Sarmiento and Gruber, 2006) (nitrate
690 concentration from World Ocean Atlas (<https://www.nodc.noaa.gov/OC5/woa13/>)), and (F)
691 export ratio derived from Dunne et al. (2005). Annual represents annually-integrated value.
692 Spring and summer represent average value in spring and summer, respectively. In the northern
693 hemisphere, spring and summer seasons are defined as March-May and June-August,

694 respectively. In the southern hemisphere, spring and summer seasons are defined as September-
695 November and December-February, respectively.

696

697 **Table 1.** Model symbols, abbreviations, and units.

Symbol	Description	Units
MLD	Mixed layer depth	m
$MLD_{C^*_{max}}$	Maximum MLD corresponds to maximum achievable autotroph's biomass concentration	m
z	Depth	m
Z_c	Critical depth	m
Z_p	Compensation depth	m
$GPP(0,z)$	Gross primary production	$\text{mmol C m}^{-2} \text{d}^{-1}$
$NPP(z)$	Net primary production at depth z	$\text{mmol C m}^{-3} \text{d}^{-1}$
$NPP(0,z)$	Net primary production above depth z	$\text{mmol C m}^{-2} \text{d}^{-1}$
$NCP(z)$	Net community production at depth z	$\text{mmol C m}^{-3} \text{d}^{-1}$
$NCP(0,z)$	Net community production above depth z	$\text{mmol C m}^{-2} \text{d}^{-1}$
$HR(z)$	Heterotrophic respiration at depth z	$\text{mmol C m}^{-3} \text{d}^{-1}$
$HR(0,z)$	Heterotrophic respiration above depth z	$\text{mmol C m}^{-2} \text{d}^{-1}$
NCP^*	The maximum NCP for a given MLD (upper bound on carbon export)	$\text{mmol C m}^{-2} \text{d}^{-1}$
NCP_B	NCP normalized to autotroph's biomass inventory in the mixed layer	d^{-1}
ef	Export ratio	unitless
f_{pt}	Ratio of satellite export production estimates to the upper bound on carbon export	unitless
N	Nutrient concentration	mmol m^{-3}
k_m^N	Half-saturation constant for nutrient concentration	mmol m^{-3}
N_m	Nutrient effect on phytoplankton growth $N_m = \frac{N}{N+k_m^N}$	unitless
PAR	Photosynthetically active radiation	$\text{Einstein m}^{-2} \text{d}^{-1}$
I_0	Photosynthetically active radiation just beneath water surface	$\text{Einstein m}^{-2} \text{d}^{-1}$
$I(z)$	Photosynthetically active radiation at depth z	$\text{Einstein m}^{-2} \text{d}^{-1}$
k_m^I	Half-saturation constant for irradiance	$\text{Einstein m}^{-2} \text{d}^{-1}$
$I_m(z)$	Light effect on phytoplankton growth at depth z , $I_m(z) = \frac{I(z)}{I(z)+k_m^I} = \frac{I_0 \times e^{-K_I \times z}}{I_0 \times e^{-K_I \times z} + k_m^I}$	unitless
$I_m(0, z)$	Integrated light effect on phytoplankton growth above depth z , $I_m(0, z) = -\frac{1}{K_I} \times \ln\left(\frac{I_0 \times e^{-K_I \times z} + k_m^I}{I_0 + k_m^I}\right)$	unitless
PAR_{ML}	Average PAR in the mixed layer ($PAR_{ML} = \frac{1-e^{-K_I \times MLD}}{K_I \times MLD} \times I_0$)	$\text{Einstein m}^{-2} \text{d}^{-1}$
μ	Phytoplankton growth rate	d^{-1}
μ_{max}	Maximum phytoplankton growth rate	d^{-1}

μ_{max}^0	Maximum phytoplankton growth rate for $T = 0$ °C	d^{-1}
r_{HR}	Heterotrophic respiration ratio	d^{-1}
r_{HR}^0	Heterotrophic respiration ratio for $T = 0$ °C	d^{-1}
K_I	Light attenuation coefficient ($K_I = K_I^w + K_I^{nw}$)	m^{-1}
K_I^w	Light attenuation coefficient due to water	m^{-1}
K_I^{nw}	Light attenuation coefficient due to optically active components	m^{-1}
k_c	Specific attenuation coefficient for irradiance	$m^2 \text{ mmol}^{-1}$
C	Phytoplankton biomass concentration	mmol m^{-3}
C^*	Phytoplankton biomass concentration that maximizes NCP	mmol m^{-3}
C_{max}^*	Maximum achievable autotroph's biomass concentration	mmol m^{-3}
POC	Particulate organic carbon	mmol m^{-3}
DOC	Dissolved organic carbon	mmol m^{-3}
CDOM	Colored dissolved organic matter	m^{-1}
NAP	Non-algal particles	mmol m^{-3}
T	Temperature	°C
P_t	Temperature dependence for phytoplankton growth rate	°C ⁻¹
B_t	Temperature dependence for heterotrophic respiration ratio	°C ⁻¹
CO ₂	Carbon dioxide	ppmv

698

699

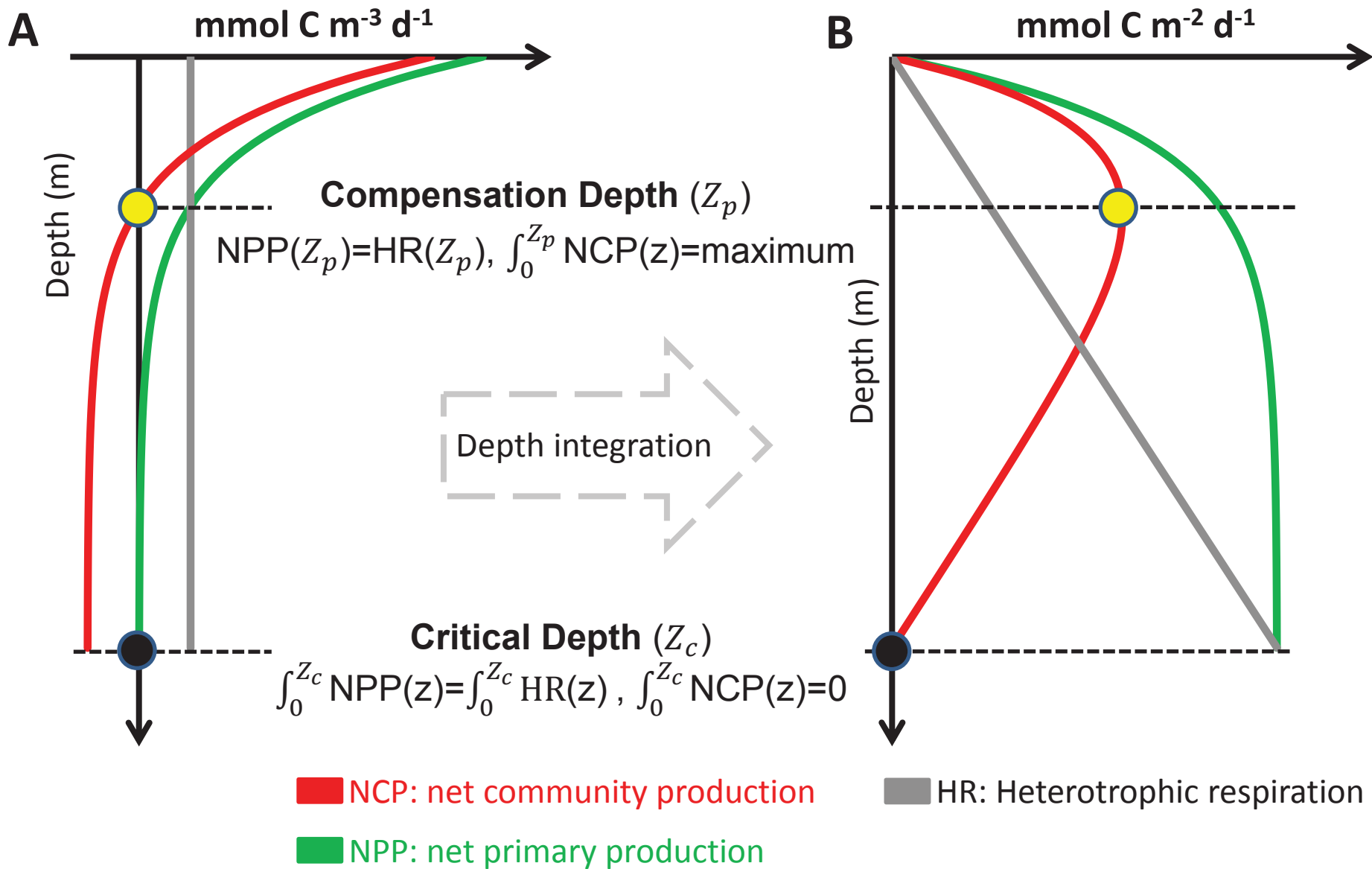
700 **Table 2.** Value or range of values with references for the parameters used in the model.
 701

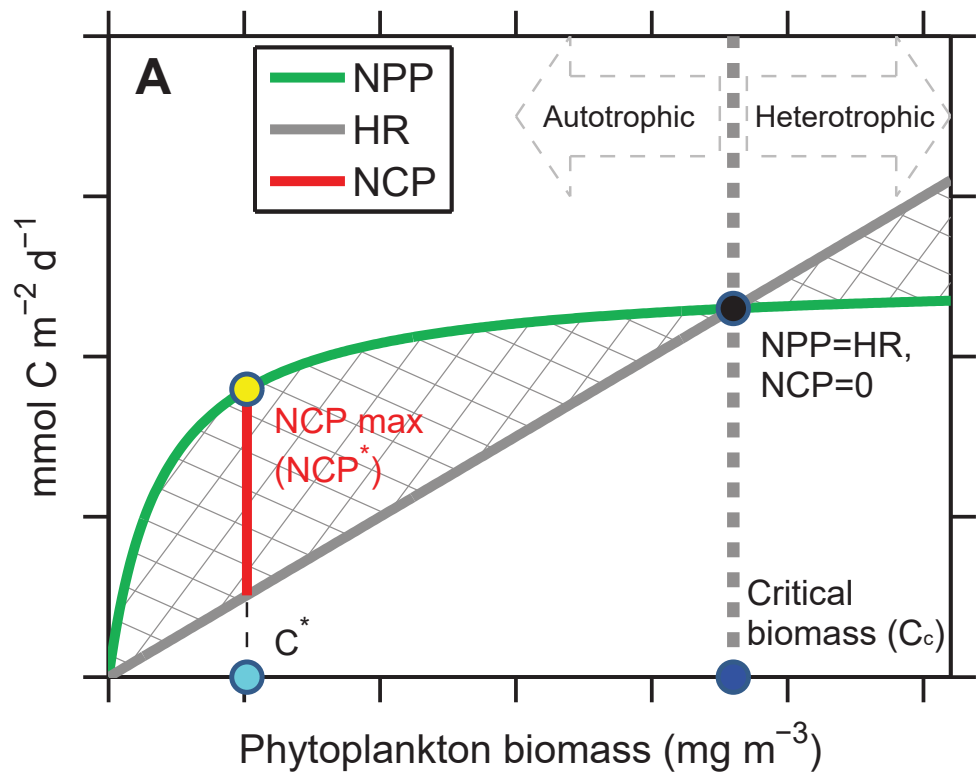
Parameter	Range or value	Reference
K_I^w	0.09	(Werdell and Bailey, 2005)
k_c	0.03	(Werdell and Bailey, 2005)
Carbon to chlorophyll ratio	90	(Arrigo et al., 2008)
k_m^I	4.1 Einstein $m^{-2} d^{-1}$	(Behrenfeld and Falkowski, 1997)
P_t	0.0663	(Eppley, 1972)
B_t	0.08	(Rivkin and Legendre, 2001; López-Urrutia et al., 2006)
μ_{max}	1 d^{-1} , 1.2 d^{-1}	(Laws et al., 2000; Eppley, 1972)
r_{HR}	0.1 d^{-1} , 0.2 d^{-1}	(Laws et al., 2000; Mitchell et al., 1991)

702

703

704





NCP =
NPP - HR

