

# 1 Simulations with the Marine Biogeochemistry Library 2 (MARBL)

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## 13 Key Points:

- 14 • MARBL is the ocean biogeochemistry component of the Community Earth Sys-  
15 tem Model (CESM)
- 16 • MARBL is a flexible, plankton functional type model with a modular architec-  
17 ture supporting portability across ocean circulation models
- 18 • CESM2 CMIP6 solutions contain significant biogeochemical biases linked to un-  
19 derlying physics

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## Abstract

The Marine Biogeochemistry Library (MARBL) is a prognostic ocean biogeochemistry model that simulates marine ecosystem dynamics and the coupled cycles of carbon, nitrogen, phosphorus, iron, silicon, and oxygen. MARBL is a component of the Community Earth System Model (CESM); it supports flexible ecosystem configuration of multiple phytoplankton and zooplankton functional types; it is also portable, designed to interface with multiple ocean circulation models. Here, we present scientific documentation of MARBL, describe its configuration in CESM2 experiments included in the Coupled Model Intercomparison Project version 6 (CMIP6), and evaluate its performance against a number of observational datasets. The model simulates an air-sea CO<sub>2</sub> flux and many aspects of the carbon cycle in good agreement with observations. However, the simulated integrated uptake of anthropogenic CO<sub>2</sub> is weak, which we link to poor thermocline ventilation, a feature evident in simulated chlorofluorocarbon distributions. This also contributes to larger-than-observed oxygen minimum zones. Moreover, radiocarbon distributions show that the simulated circulation in the deep North Pacific is extremely sluggish, yielding extensive oxygen depletion and nutrient trapping at depth. Surface macronutrient biases are generally positive at low latitudes and negative at high latitudes. CESM2 simulates globally-integrated net primary production (NPP) of 48 Pg C yr<sup>-1</sup> and particulate export flux at 100 m of 7.1 Pg C yr<sup>-1</sup>. The impacts of climate change include an increase in globally-integrated NPP, but substantial declines in the North Atlantic. Particulate export is projected to decline globally, attributable to decreasing export efficiency associated with changes in phytoplankton community composition.

## Plain Language Summary

Numerical models of the ocean carbon cycle and biogeochemistry play a key role in understanding the fate of human carbon dioxide emissions and the magnitude of expected climate change over the next several decades to a century. Models are needed to quantify changes in the carbon reservoirs of the ocean and atmosphere and to explore interactions between climate change and carbon reservoirs that could amplify or damp future warming. This paper presents the Marine Biogeochemistry Library (MARBL), which is an ocean biogeochemistry model coupled to the Community Earth System Model (CESM). MARBL was designed to be compatible with multiple ocean models, a design motivated by an interest in building a diverse community of researchers around the development of MARBL. This paper presents a technical description of MARBL and an evaluation of the ocean biogeochemical simulation in CESM version 2. Overall, the model captures large-scale biogeochemical distributions, though several important biases are highlighted, including those dependent on the representation of circulation. MARBL provides a robust platform for researchers to address critical questions related to the impacts of climate variability and change on marine ecosystems.

## 1 Introduction

The ocean comprises the largest active carbon reservoir on Earth, storing approximately 38,000 Pg C of natural CO<sub>2</sub>, nearly all of it as dissolved inorganic carbon (DIC). The ocean has also absorbed about 30% of anthropogenic CO<sub>2</sub> emissions (152 Pg through 2007) since the beginning of the industrial revolution (Sabine et al., 2004; Gruber et al., 2019)—and this sink will remain an important control on the airborne fraction of CO<sub>2</sub> emissions (Jones et al., 2013). Given this fundamental importance, Earth system models (ESMs) include ocean biogeochemistry models (OBMs) that seek to represent the ocean carbon cycle mechanistically, enabling future projections inclusive of carbon-climate feedbacks (Friedlingstein et al., 2006). As these models have matured, there has been increasing recognition of their relevance to questions beyond biogeochemistry, and in particular related to ocean ecosystems in the context of climate variability and change (Bopp

70 et al., 2013; Stock et al., 2011; Tommasi et al., 2017). This paper describes the Marine  
71 Biogeochemistry Library (MARBL), which is the ocean biogeochemistry component for  
72 the Community Earth System Model, version 2 (CESM2) (Danabasoglu et al., 2020).  
73 We document MARBL and evaluate the ocean biogeochemistry simulations in the fully-  
74 coupled CESM2 integrations submitted to the Coupled Model Intercomparison Project  
75 phase 6 (CMIP6) (Eyring et al., 2016).

76 In CESM2, MARBL was configured to invoke an updated version of what has pre-  
77 viously been known as the Biogeochemistry Elemental Cycle (BEC) model (Moore et  
78 al., 2002, 2002a, 2004, 2013). MARBL is a modularized code base, however, consisting  
79 of a self-contained, independent Fortran library that interfaces with an ocean general cir-  
80 culation model (OGCM) through an explicit driver layer. This explicit design decision  
81 enables interoperability between MARBL and different physical models (see section 2.1).  
82 Moreover, MARBL has some flexibility with respect to the configuration of its ecosys-  
83 tem, supporting simulation of an arbitrary number of zooplankton and phytoplankton  
84 functional types (PFTs).

85 The BEC model has been under development for more than two decades, with ini-  
86 tial roots in a one-dimensional configuration applied to represent the seasonal cycle of  
87 upper ocean primary production at the Joint Global Ocean Flux Study (JGOFS), Bermuda  
88 Atlantic Timeseries Study site (Doney et al., 1996). This formulation was extended by  
89 Moore et al. (2002) to include three phytoplankton functional types, multiple nutrient  
90 co-limitation (N, P, Si, Fe), nitrogen fixation, and calcification. At this point, the model  
91 was run in the mixed-layer on a global grid, but there was no lateral exchange and nu-  
92 trient concentrations below the mixed layer were specified from an observationally-based  
93 climatology (Moore et al., 2002). The Moore et al. (2002) model was capable of repro-  
94 ducing the dominant patterns of primary production, nitrogen fixation, and export, in-  
95 cluding the observed high nitrate, low chlorophyll (HNLC) conditions in the Southern  
96 Ocean, Subarctic Northeast Pacific, and equatorial Pacific (Moore et al., 2002a). Moore  
97 et al. (2004) implemented the BEC model in a three-dimensional global OGCM for the  
98 first time, demonstrating skillful biogeochemical solutions under prognostic flow forced  
99 by meteorological fields from atmospheric reanalysis data. This implementation also en-  
100 abled running the model in a “fully-coupled” context, in which atmospheric and ocean  
101 general circulation models exchange information via a flux coupler. A series of studies  
102 examined BEC solutions in the Community Climate System Model, version 3 (CCSM3)  
103 (Collins et al., 2006; Yeager et al., 2006). These include examinations of the effect of at-  
104 mospheric nutrient deposition (Moore & Doney, 2007; Moore et al., 2006; Krishnamurthy  
105 et al., 2007, 2009, 2010; Han et al., 2008; Doney et al., 2007; Mahowald et al., 2011), climate-  
106 change impacts on ecosystems (Marinov et al., 2010), carbon-climate feedbacks (Thornton  
107 et al., 2009), the mechanisms driving carbon fluxes (Lovenduski et al., 2008; Wang et  
108 al., 2012; Doney et al., 2009) and the impacts of volcanic forcing on ocean biogeochem-  
109 istry (Rothenberg et al., 2012). The Community Climate System Model evolved into the  
110 Community Earth System Model, and the BEC model was released to the community  
111 in this context. Moore et al. (2013) documented the marine ecosystem response to cli-  
112 mate change in CESM1; Long et al. (2013) described the simulation of the present-day  
113 ocean carbon sink and Lindsay et al. (2014) examined the fully-coupled carbon cycle in  
114 CESM1. In addition to projections out to 2100, CESM1 was used to conduct Extended  
115 Representative Concentration Pathway (RCP) scenario integrations out to the year 2300  
116 (Moore et al., 2018; Randerson et al., 2015). A version of CESM1 was used to conduct  
117 a large ensemble (CESM-LE) experiment (Kay et al., 2015), which included ocean bio-  
118 geochemistry simulated by BEC. The CESM-LE enabled a series of studies that explic-  
119 itly separated natural variability from anthropogenic forced trends in ocean biogeochem-  
120 istry (Long et al., 2016; Lovenduski et al., 2015, 2016; McKinley et al., 2016; Krumhardt  
121 et al., 2017; Eddebbar et al., 2019). BEC has also been used in Decadal Prediction ex-  
122 periments with CESM1 (Yeager et al., 2012), and a handful of studies have examined

123 predictability of ocean biogeochemical dynamics in this framework (e.g., Krumhardt et  
 124 al., 2020; Lovenduski et al., 2019; Yeager et al., 2018).

125 Our objectives in this paper are to document MARBL and specifically its config-  
 126 uration in the CESM2 integrations submitted to CMIP6. MARBL development has con-  
 127 tinued since the CMIP6 integrations were conducted, and there a handful of MARBL  
 128 features that were not enabled in the CMIP runs. We refer to the CMIP6 configuration  
 129 of MARBL, the default configuration in CESM2.1, as MARBL-CESM2.1 to explicitly  
 130 note the associated model version and its configuration. That is, statements made about  
 131 MARBL hold for all configurations of MARBL, as it was released in CESM2.1, and state-  
 132 ments about MARBL-CESM2.1 hold for the default configuration in that release.

## 133 2 Model description

### 134 2.1 Flexible implementation

135 MARBL is a stand-alone Fortran library designed to be implemented in multiple  
 136 OGCMs. The MARBL framework can be best understood by reference to the prognos-  
 137 tic equation governing the evolution of an arbitrary passive tracer  $\chi$  in an OGCM:

$$\frac{\partial \chi}{\partial t} + \nabla \cdot (\mathbf{u} \chi) - \nabla \cdot (K \nabla \chi) = J_{\chi}(\mathbf{x}), \quad (1)$$

138 where the terms on the left hand side (LHS) are the time-tendency, resolved three-dimensional  
 139 advection by the simulated velocity field,  $\mathbf{u}$ , and diffusion determined by the diffusiv-  
 140 ity  $K$ .  $J_{\chi}(\mathbf{x})$  is the sum of sources-minus-sinks for  $\chi$ , computed as a function of the model  
 141 state vector,  $\mathbf{x}$ ; air-sea, benthic, or riverine fluxes provide boundary conditions. MARBL  
 142 computes the source/sink terms and interfaces with an OGCM through an explicit driver  
 143 layer, such that multiple OGCMs that have implemented the MARBL-driver can call  
 144 the identical MARBL code. The OGCM computes the LHS of Eq. (1) and handles time-  
 145 integration. MARBL has been implemented in the Parallel Ocean Program version 2 (POP2;  
 146 the CESM2 ocean component) (Danabasoglu et al., 2012), the Model for Prediction Across  
 147 Scales, Ocean (MPAS-O) (Ringler et al., 2013; Burrows et al., 2020), and the Modular  
 148 Ocean Model, version 6 (MOM6) (Adcroft et al., 2019), which will be the ocean com-  
 149 ponent in CESM3. MARBL is fully configurable at run-time, and is flexible enough to  
 150 accommodate a variable number of plankton functional types and ecosystem configura-  
 151 tions. MARBL is developed via an open-development process on GitHub, including con-  
 152 tinuous integration and testing, as well as tools supporting appropriate configuration in  
 153 CESM integrations. The repository can be found at [https://github.com/marbl-ecosys/](https://github.com/marbl-ecosys/MARBL)  
 154 MARBL.

### 155 2.2 Ocean biogeochemistry formulations

156 MARBL allows for easy addition of phytoplankton and zooplankton groups to the  
 157 ecosystem. Both are treated as arrays, so to add a phytoplankton or zooplankton group  
 158 simply requires specifying their parameter values in an input file. Key biogeochemical  
 159 processes are built into MARBL, allowing the user to specify the biogeochemical func-  
 160 tions associated with each phytoplankton group (i.e., calcifier, silicifier, capable of N fix-  
 161 ation, etc.). This facilitates supporting multiple ecosystem configurations spanning a range  
 162 in complexity. Here we focus on the implementation in CESM2 CMIP6 integrations, which  
 163 preserved the ecosystem configuration in CESM1: MARBL-CESM2.1 includes one zoo-  
 164 plankton group, three explicit phytoplankton functional groups (diatoms, diazotrophs,  
 165 “small” pico/nano phytoplankton), and one implicit group (calcifiers). MARBL-CESM2.1  
 166 thus simulates 32 tracers, comprising 17 non-living constituents (dissolved inorganic car-  
 167 bon, alkalinity, nutrients, oxygen, and dissolved organic matter) and 15 tracers associ-  
 168 ated with living biomass. Carbonate chemistry is fully-explicit, and there are two par-

169 allel carbonate systems (e.g., with preindustrial and contemporary atmospheric CO<sub>2</sub>),  
 170 which enables cleanly computing anthropogenic CO<sub>2</sub> concentrations.

### 171 **2.2.1 Phytoplankton growth**

172 The source/sink term of the phytoplankton group ( $J_{P_i}$ ) is

$$J_{P_i} = \mu_i P_i - G(P_i') - m_i T_f P_i' - A(P_i') \quad (2)$$

173 where the terms on the right-hand side represent growth, and sinks due to grazing ( $G$ ),  
 174 linear mortality ( $m_i$ ) and aggregation ( $A$ ). The loss terms are dependent on a  $P_i'$ , which  
 175 is the phytoplankton concentration in excess of a temperature- and depth-dependent thresh-  
 176 old. The C-specific growth rate,  $\mu_i$ , is parameterized as the product of the resource-unlimited  
 177 growth rate ( $\mu_{ref}$ ) at a reference temperature (30°C), and temperature ( $T_f$ ), nutrient  
 178 limitation ( $V_i$ ) and light availability ( $L_i$ ) functions:

$$\mu_i = \mu_{ref} T_f V_i L_i. \quad (3)$$

179 The temperature dependence formulation,

$$T_f = 1.7^{\left(\frac{T-30^\circ\text{C}}{10^\circ\text{C}}\right)}, \quad (4)$$

180 is based on results from Sherman et al. (2016).

181 Light-limitation is computed as a function of irradiance,  $I$  (W m<sup>-2</sup>), using a mod-  
 182 ified form of the Geider et al. (1998, 1997) dynamic growth model,

$$L_i = 1 - \exp\left(\frac{-\alpha_i^{Chl} \theta_i^C I}{\mu_{ref} T_f V_i}\right), \quad (5)$$

183 where  $\alpha_i^{Chl}$  (g C m<sup>2</sup> (g Chl W s)<sup>-1</sup>) is the initial slope of the chlorophyll-*a* (*Chl*) spe-  
 184 cific photosynthesis-irradiance (PI) curve and  $\theta_i^C$  is the chlorophyll to carbon ratio (g  
 185 Chl:g C). This equation represents the growth rate as a function of the ratio between  
 186 the supply of energy for photosynthesis in the form of instantaneous light harvesting ca-  
 187 pacity ( $\alpha_i^{Chl} \cdot \theta_i^C \cdot I$ ), and the demand for growth in terms of the maximum photosyn-  
 188 thetic rate constrained by temperature and nutrient limitation ( $\mu_{ref} \cdot T_f \cdot V_i$ ). The chloro-  
 189 phyll to carbon ratio,  $\theta_i^C$ , evolves prognostically in the model, thereby providing a rep-  
 190 resentation of photoadaptation. The source/sink term for chlorophyll is

$$J_{Chl} = \rho_{Chl} \left( \frac{\mu_i}{r_{C:N} \theta^C} \right) \quad (6)$$

191 where  $r_{C:N}$  is the carbon to nitrogen stoichiometry of phytoplankton (see below) and  $\rho_{Chl}$   
 192 is the dimensionless chlorophyll synthesis term (Geider et al., 1998), computed as

$$\rho_{Chl} = \theta_{max,i}^N \frac{\mu_i}{\alpha_i^{Chl} \theta_i^C I}. \quad (7)$$

193 Photosynthetically available radiation ( $I$ ) is assumed to be 45% of incoming short-  
 194 wave radiation (Doney et al., 1996). CESM simulates a subgrid-scale sea-ice thickness  
 195 distribution and computes shortwave penetration independently for each ice thickness  
 196 category, yielding multiple sub-columns with different light levels. MARBL computes  
 197 all light-dependent terms on each sub-column independently, and then computes the grid-  
 198 cell mean terms by taking an area-weighted average across sub-columns (Long et al., 2015).  
 199 Since the photosynthesis equations are non-linear, this order of operations reduces nu-  
 200 merical artifacts that arise where light fields are heterogeneous. Long et al. (2015) showed  
 201 that this approach dramatically reduces biases in the simulation of high-latitude spring  
 202 blooms relative to photosynthesis computed on grid-cell mean light.

**Table 1.** Model parameters.

Parameter	Value	Units	Description
$\mu_{ref}$	5.0	d <sup>-1</sup>	Maximum C-spec growth rate at Tref for small phytoplankton and diatoms
$\mu_{ref, diat}$	2.5	d <sup>-1</sup>	Maximum C-spec growth rate at Tref for diatoms
$Q_{10}$	1.7	unitless	Q10 temperature coefficient
$T_{ref}$	30	°C	Reference temperature
$\theta_N^{max, sp}$	2.5	mg Chl / mmol	Maximum $\theta^N$ (Chl / N) for small phytoplankton
$\theta_N^{max, diat}$	4.0	mg Chl / mmol	Maximum $\theta^N$ (Chl / N) for diatoms
$\theta_N^{max, diat}$	2.5	mg Chl / mmol	Maximum $\theta^N$ (Chl / N) for diatoms
$\alpha_{chl, sp}$	0.39	mmol m <sup>2</sup> / (mg Chl W day)	Chl-spec initial slope of P-I curve (GD98) for small phytoplankton
$\alpha_{chl, diat}$	0.28	mmol m <sup>2</sup> / (mg Chl W day)	Chl-spec initial slope of P-I curve (GD98) for diatoms
$\alpha_{chl, diat}$	0.39	mmol m <sup>2</sup> / (mg Chl W day)	Chl-spec initial slope of P-I curve (GD98) for diatoms
$K_{Fe}$	3.0e-05	mmol/m <sup>3</sup>	Fe uptake half-sat constant for small phytoplankton
$K_{Fe, sp}$	7.0e-05	mmol/m <sup>3</sup>	Fe uptake half-sat constant for diatoms
$K_{Fe, diat}$	4.5e-05	mmol/m <sup>3</sup>	Fe uptake half-sat constant for diatoms
$K_{NO_3}$	0.25	mmol/m <sup>3</sup>	NO3 uptake half-sat constant for small phytoplankton
$K_{NO_3, sp}$	0.5	mmol/m <sup>3</sup>	NO3 uptake half-sat constant for diatoms
$K_{NO_3, diat}$	2.0	mmol/m <sup>3</sup>	NO3 uptake half-sat constant for diatoms
$K_{NH_4}$	0.01	mmol/m <sup>3</sup>	NH4 uptake half-sat constant for small phytoplankton
$K_{NH_4, sp}$	0.05	mmol/m <sup>3</sup>	NH4 uptake half-sat constant for diatoms
$K_{NH_4, diat}$	0.2	mmol/m <sup>3</sup>	NH4 uptake half-sat constant for diatoms
$K_{PO_4}$	0.01	mmol/m <sup>3</sup>	PO4 uptake half-sat constant for small phytoplankton
$K_{PO_4, sp}$	0.05	mmol/m <sup>3</sup>	PO4 uptake half-sat constant for diatoms
$K_{PO_4, diat}$	0.015	mmol/m <sup>3</sup>	PO4 uptake half-sat constant for diatoms
$K_{DOP}$	0.3	mmol/m <sup>3</sup>	DOP uptake half-sat constant for small phytoplankton
$K_{DOP, sp}$	0.5	mmol/m <sup>3</sup>	DOP uptake half-sat constant for diatoms
$K_{DOP, diat}$	0.075	mmol/m <sup>3</sup>	DOP uptake half-sat constant for diatoms
$K_{SiO_3}$	0.7	mmol/m <sup>3</sup>	SiO3 uptake half-sat constant for diatoms
$a_{min, sp}$	0.01	d <sup>-1</sup>	Minimum agg rate for small phytoplankton
$a_{min, diat}$	0.02	d <sup>-1</sup>	Minimum agg rate for diatoms
$a_{min}$	0.01	d <sup>-1</sup>	Minimum agg rate for diatoms
$a_{max}$	0.5	d <sup>-1</sup>	Maximum agg rate for small phytoplankton
$a_{max, sp}$	0.5	d <sup>-1</sup>	Maximum agg rate for diatoms
$a_{max, diat}$	0.5	d <sup>-1</sup>	Maximum agg rate for diatoms
$a_{max}$	3.3	d <sup>-1</sup>	Maximum agg rate for diatoms
$g_{max}$	3.15	d <sup>-1</sup>	Maximum grazing rate at Tref for small phytoplankton
$g_{diat}$	3.3	d <sup>-1</sup>	Maximum grazing rate at Tref for diatoms
$g_{diat}$	3.3	d <sup>-1</sup>	Maximum grazing rate at Tref for diatoms
$K_i$	1.2	mmol/m <sup>3</sup>	Zooplankton grazing half saturation constant for all phytoplankton
$\gamma_{sp}$	0.3	unitless	Gross growth efficiency coefficient for grazing for diatoms
$\gamma_{diat}$	0.25	unitless	Gross growth efficiency coefficient for grazing for diatoms
$\gamma_{diat}$	0.3	unitless	Gross growth efficiency coefficient for grazing for diatoms
$m_Z$	0.1	d <sup>-1</sup>	Zooplankton linear mortality rate at Tref
$a_Z$	0.4	d <sup>-1</sup> / (mmol/m <sup>3</sup> )	Zooplankton aggregation mortality rate at Tref

203 Phytoplankton nutrient requirements vary by taxa; diatoms (*diat*) require N, P,  
 204 Si, and Fe, where as small phytoplankton (*sp*) and diazotrophs (*diaz*) do not assimilate  
 205 Si and diazotrophs are not limited by N. Nutrient limitation is computed using Leibig's  
 206 law of the minimum, such that

$$\begin{aligned} V_{diat} &= \min(V_{diat}^N, V_{diat}^P, V_{diat}^{Si}, V_{diat}^{Fe}), \\ V_{sp} &= \min(V_{sp}^N, V_{sp}^P, V_{sp}^{Fe}), \text{ and} \\ V_{diaz} &= \min(V_{diaz}^P, V_{diaz}^{Fe}). \end{aligned} \quad (8)$$

207 Nutrient limitation is represented according to Michaelis-Menten kinetics, where

$$V_i^{Fe} = \frac{Fe}{Fe + K_i^{Fe}} \text{ and } V_i^{Si} = \frac{SiO_3}{SiO_3 + K_i^{SiO_3}}. \quad (9)$$

208 However, phytoplankton can alternatively assimilate nitrate ( $NO_3$ ) and ammonium ( $NH_4$ ),  
 209 following the substitutable model of O'Neill et al. (1989), such that

$$\begin{aligned} V_i^{NO_3} &= \frac{NO_3/K_i^{NO_3}}{1 + NO_3/K_i^{NO_3} + NH_4/K_i^{NH_4}}; \\ V_i^{NH_4} &= \frac{NH_4/K_i^{NH_4}}{1 + NO_3/K_i^{NO_3} + NH_4/K_i^{NH_4}}; \text{ and} \\ V_i^N &= V_i^{NO_3} + V_i^{NH_4}. \end{aligned} \quad (10)$$

210 All taxa are capable of assimilating both phosphate ( $PO_4$ ) and semi-labile (see below)  
 211 dissolved organic phosphate (DOP); a similar approach is used to compute limitation  
 212 terms for these constituents.

213 Nutrient assimilation ratios are fixed for C:N according to Anderson and Sarmiento  
 214 (1994) (117:16), but permitted to vary for P, Fe, and Si. P:C uptake for all PFTs is com-  
 215 puted according to a modified version of the linear model of Galbraith and Martiny (2015).  
 216 P:C uptake rates are linked to ambient phosphate concentrations, with P:C increasing  
 217 linearly with increasing available phosphate, until a maximum P:C value is reached (Wang  
 218 et al., 2019). The dynamic Fe:C ratios follow a similar formulation, with Fe:C ranging  
 219 between specified minimum and maximum values as a function of ambient iron concen-  
 220 tration (Moore et al., 2004). The Si:C uptake ratio for the diatoms is a function of both  
 221 ambient iron and silicate concentrations, whereby low iron increases Si:C uptake and low  
 222 silicate decreases the Si:C uptake ratio for new growth (Moore et al., 2004). As ambi-  
 223 ent nutrients concentrations change over time, phytoplankton nutrient assimilation ra-  
 224 tios respond, leading to changes in the stoichiometry of phytoplankton biomass.

225 Loss of phytoplankton due to aggregation is parameterized as

$$A(P'_i) = a_i P_i'^{1.75} \quad (11)$$

226 but constrained to fall between imposed minimum,  $a_i^{min} \cdot P'_i$ , and maximum,  $a_i^{max} \cdot P'_i$ ,  
 227 rates.

228 MARBL includes a representation of phytoplankton calcification as well as opal pro-  
 229 duction by diatoms; these materials play an important role in mediating particulate or-  
 230 ganic carbon export (see below). Calcification in MARBL-CESM2.1 is treated implic-  
 231 itly as a varying fraction of the small phytoplankton NPP, similar to the approach in CESM1  
 232 (Moore et al., 2004). Baseline calcification is 7% of small phytoplankton NPP, but is de-  
 233 creased as nutrient limitation increases by multiplying calcification by the nutrient lim-  
 234 itation term squared ( $V_{sp}^2$ ). This decline in calcification under severe nutrient limita-  
 235 tion aims to represent competition between calcifying coccolithophores and smaller pi-  
 236 coplankton; calcification is thus reduced in the oligotrophic gyres where picoplankton  
 237 have an advantage due to their larger cell surface-area-to-volume ratios (Moore et al.,

238 2002). Calcification is linearly reduced at temperatures below 4°C, reflecting the fact that  
 239 coccolithophores are rare in polar waters (Iglesias-Rodríguez et al., 2002; Holligan et al.,  
 240 2010). When small phytoplankton biomass exceeds 2.5 mmol C m<sup>-3</sup>, calcification is scaled  
 241 up to 40% of small phytoplankton production, providing a representation of enhanced  
 242 calcification associated with blooms of the coccolithophore *Emiliania huxleyi*. Opal for-  
 243 mation by diatoms is computed by multiplying diatom production by Si:C elemental ra-  
 244 tio, which modified by Fe and Si ambient conditions, as described above.

### 245 2.2.2 Zooplankton

246 MARBL-CESM2.1 has one zooplankton class that grazes on phytoplankton accord-  
 247 ing to a Holling Type II relationship

$$G(P'_i) = g_i^{max} \cdot T_f \cdot \left( \frac{P'_i}{P'_i + K_i^P} \right) Z \quad (12)$$

248 The values of half-saturation constants,  $K_i^P$ , are identical for each phytoplankton taxa  
 249 in MARBL-CESM2.1; the maximum grazing rates,  $g_i^{max}$ , vary, however, with diatoms  
 250 experiencing the lowest grazing pressure. The temperature dependence of grazing,  $T_f$ ,  
 251 is the same as for phytoplankton growth and mortality. The source/sink equation for zoo-  
 252 plankton is thus

$$J_Z = \gamma_{diat} G(P'_{diat}) + \gamma_{sp} G(P'_{sp}) + \gamma_{diaz} G(P'_{diaz}) - m_Z T_f Z' - a_Z T_f Z'^{1.5} \quad (13)$$

253 where  $\gamma_i$  is a gross growth efficiency coefficient (Straile, 1997) and the last two terms on  
 254 the right-hand side represent linear and aggregation mortality, respectively (see Table 1).  
 255  $Z'$  is the zooplankton concentration in excess of a depth-dependent threshold.

256 Zooplankton ingestion is partitioned into three fractions: converted to zooplank-  
 257 ton biomass ( $\gamma_i$ ), lost to egestion, and lost to respiration. The fraction of egestion losses  
 258 (roughly 30% of ingestion, 45% for diatom prey) partitioned to sinking detritus varies  
 259 by phytoplankton prey type, such that 10% and 38% of ingested diazotrophs and diatoms  
 260 goes to POC, respectively. The fraction of the grazed small phytoplankton material routed  
 261 to sinking particulate material varies as a function biomass, with reductions in the frac-  
 262 tion exported at low biomass. The remainder of the egestion losses are partitioned to  
 263 DOC (6% of ingestion for all phytoplankton types) and DIC (the remainder). Zooplank-  
 264 ton respiration losses are assumed to be primarily a function of ingestion (40% for small  
 265 phytoplankton and diazotrophs, and 30% for diatoms). The differential routing of zoo-  
 266 plankton grazing is aimed at simulating various types of zooplankton (microzooplank-  
 267 ton, mesozooplankton) within a single, “adaptive zooplankton” class.

268 Following Doney et al. (1996), zooplankton losses include linear and “quadratic”  
 269 (here, “aggregation”) loss terms. The linear losses represent a combination of metabolic  
 270 and reproductive losses, as well as non-predatory mortality, while the aggregation losses  
 271 approximately represent predation by unresolved higher trophic level predators (Fasham,  
 272 1995). The loss coefficients  $m_Z$  and  $a_Z$  are tuned to be consistent with overall mortal-  
 273 ity rates used in previous versions of the model (Doney et al., 1996; Moore et al., 2004,  
 274 2013). The routing of zooplankton losses to DIC, DOC, and POC are computed simi-  
 275 larly for both linear and aggregation mortality losses; though, to simulate the effect of  
 276 various zooplankton types, there are differential losses to POC based on phytoplankton  
 277 prey type. 12% of zooplankton losses originating from grazing on small phytoplankton  
 278 and diazotrophs, and 24% originating from grazing on diatoms, are routed to POC. Of  
 279 the remainder, 6% is partitioned to DOC, which approximates the fraction of the semi-  
 280 labile losses, and the remainder is routed to DIC.

**Table 2.** Remineralization length scales (in meters) for sinking particulate matter as a function of depth. The 100 m value is also used above that depth, and the 1000 m value is also used at deeper depths; for all values in between, the length scale is linearly interpolated from the values in the table.

depth (m)	POC	SiO <sub>2</sub>	CaCO <sub>3</sub>
100	100	650	500
250	360	2340	1800
500	470	3055	2350
1000	480	3120	2400

### 2.2.3 Detrital organic pools

There is no explicit sinking particulate organic matter (POM) tracer in MARBL; rather, sinking POM is simulated implicitly following Armstrong et al. (2002). In this formulation, sinking POM is redistributed in the water column as it is produced in each numerical timestep, with no time lag between surface production and deep remineralization. The redistribution follows the analytical solution to a first-order differential equation expressing POM flux as a function of a sinking velocity and remineralization rate, which together can be described by a remineralization length scale (Sarmiento & Gruber, 2006). Sinking POM is subject to ballasting by mineral dust, biogenic CaCO<sub>3</sub> and Si, which enhance the efficiency of export (Armstrong et al., 2002). Remineralization length-scales vary as a function of depth in MARBL, enabling a representation of the increasingly recalcitrant composition of sinking material at depth (Sarmiento & Gruber, 2006; Lima et al., 2014). In the CMIP6 integrations, the remineralization length scale was specified as 100 m for the upper 100 m, increasing by a factor of 4.8 by 1000 m (see Table 2 for the precise formulation). Remineralization length scales are also increased linearly in the presence of low oxygen ( $< 45 \text{ mmol m}^{-3}$ ). Remineralization rates have no dependence on temperature.

MARBL simulates 6 dissolved organic matter (DOM) pools, including semi-labile (SLDOM) and refractory dissolved organic (RDOM) carbon, nitrogen, and phosphorus (DOP) (Letscher & Moore, 2015; Letscher et al., 2015). MARBL does not include an explicit heterotrophic bacteria pool and thus does not capture the part of the “microbial loop” associated with assimilation of DOM by bacteria that are grazed by microzooplankton (Azam et al., 1983). The parameters controlling DOM cycling were optimized in an offline tracer-transport framework constrained by DOM observations (Letscher et al., 2015). Preferential remineralization of DOP leads to a DOM pool enriched in C and N relative to the composition of phytoplankton (Letscher & Moore, 2015). Both SLDOM and RDOM pools are produced from phytoplankton and zooplankton losses (mortality and aggregation; see above), as well as from grazing due to incomplete assimilation of grazed material. 6% of both the phytoplankton losses and grazing fluxes is routed to DOM and the fractional allocation of this material that flows to semi-labile and refractory pools is controlled via a fixed parameter. The refractory DOM pools also receive 6% of the POM remineralization flux. DOM remineralization has no explicit temperature dependence. DOM pools are remineralized at a rate determined by ambient light levels; remineralization rates for SLDOM are significantly higher in the dark (Letscher et al., 2015). The opposite is true for RDOM, where remineralization is enhanced in the presence of light due to photodegradation by ultraviolet light. SLDOM pools cycle with rates on the order of years; the RDOM pools have remineralization timescales of years at the surface increasing to many millenia at depth. Semi-labile DOP has an additional sink in the up-

319 per ocean associated with its use as a P source sustaining phytoplankton nutrient require-  
 320 ments when phosphate concentrations are low (Letscher et al., 2015).

#### 321 **2.2.4 Nitrogen cycle**

322 MARBL simulates the marine nitrogen cycle with inputs from rivers, atmospheric  
 323 deposition, and prognostic N fixation; losses of N include water column and sedimentary  
 324 denitrification as well as ammonia emissions from the sea surface. Nitrogen fixation is  
 325 simulated based on a fixed ratio of 1.25 between diazotroph N fixation and C fixation,  
 326 which depends on adequate light and Fe availability. Diazotrophs are not limited by N,  
 327 but do assimilate nitrate and ammonium when available—though given their much slower  
 328 growth rates, they are at a competitive disadvantage relative to other taxa where in re-  
 329 gions where N is not limiting.

330 Nitrification (the oxidation of ammonium to nitrite) is simulated as a first-order  
 331 rate process dependent on the concentration of ammonium with a rate constant of  $0.06 \text{ d}^{-1}$ .  
 332 The model does not explicitly simulate nitrite or  $\text{N}_2\text{O}$ : nitrification in the model thus  
 333 represents both ammonium and nitrite oxidation and nitrate is the only product; there  
 334 are no losses to  $\text{N}_2$  via nitrification—therefore, nitrification does not result in losses of  
 335 fixed N from the model ocean. Nitrification is light-inhibited and only occurs in MARBL  
 336 where PAR is below  $1 \text{ W m}^{-2}$ . The subgrid-scale treatment of light (see above) is ap-  
 337 plied to the nitrification computation, such that nitrification is computed for each sub-  
 338 column and the ultimate grid-cell mean is an area-weighted average across sub-columns.  
 339 CESM simulates a diurnal light cycle, thus there is nitrification in the surface ocean at  
 340 night.

341 Denitrification is the oxidation of organic matter via nitrate. Water column deni-  
 342 trification is simulated as a function of organic matter remineralization and local oxy-  
 343 gen concentrations. Where oxygen declines below  $10 \text{ mmol m}^{-3}$ , the fraction of organic  
 344 matter oxidation accounted for by denitrification is linearly increased until oxygen reaches  
 345  $5 \text{ mmol m}^{-3}$ , where denitrification is assumed to account for 100% of organic matter ox-  
 346 idation. MARBL also simulates sedimentary denitrification on the basis of an empiri-  
 347 cal relation depending on POC flux to the seafloor (Bohlen et al., 2012). Water column  
 348 and sedimentary denitrification are reduced where nitrate concentrations approach zero.  
 349 Denitrification does lead to fixed N loss from the model, to balance N fixation, but the  
 350  $\text{N}_2$  product is not explicitly tracked. Oceanic emission of ammonia is simulated prog-  
 351 nostically following Paulot et al. (2015).

#### 352 **2.2.5 Iron cycle**

353 MARBL includes a representation of oceanic iron cycling that is an extension of  
 354 the formulation described in Moore and Braucher (2008). Sources of dissolved iron to  
 355 the ocean specified via forcing include dissolved iron inputs from sediments ( $\sim 20 \text{ Gmol yr}^{-1}$ ),  
 356 hydrothermal vents ( $\sim 5.0 \text{ Gmol yr}^{-1}$ ), and rivers ( $0.37 \text{ Gmol yr}^{-1}$ ). The sedimentary  
 357 iron source is applied using subgrid-scale bathymetry; this results in a vertical distribu-  
 358 tion of iron input in the water column, relative to simply applying the source at the model  
 359 bottom. The sedimentary iron source is a temporally-static field; it is computed offline  
 360 using a parameterization that depends on POC fluxes and bottom-current velocity simu-  
 361 lated by CESM. The source of iron from oxic sediments is parameterized via a constant,  
 362 low background value; this source is increased in regions of high bottom horizontal cur-  
 363 rent speed (sediment re-suspension) according to the current velocity squared by up to  
 364 a factor of 100. The source of iron from reducing sediments is linearly related to the sink-  
 365 ing POC flux where the POC flux exceeds  $3 \text{ g C m}^{-2} \text{ yr}^{-1}$ ; below this threshold, the re-  
 366 ducing sediment source is zero. This puts a source on the shelf, and along productive  
 367 slope/margins, but has little source in the deep ocean, where almost all the remineral-  
 368 ization is oxic right on the sediment surface. The two iron source types were combined

369 into one Fe input field for CESM2. Atmospheric deposition of soluble iron is computed  
 370 prognostically as a function of dust and black carbon deposition provided by the atmo-  
 371 spheric model. Dust also contributes sources of phosphate and silicate, following Krishnamurthy  
 372 et al. (2010). The iron cycle includes a representation of scavenging (Moore & Braucher,  
 373 2008) and complexation by an explicit ligand tracer. The ligand tracer has sources due  
 374 to remineralization and dissolved organic matter production.

### 375 **2.2.6 Riverine forcing**

376 Riverine nutrient (N, P, Si, Fe), dissolved inorganic carbon, alkalinity, and DOM  
 377 fluxes are supplied to the CESM2 ocean model from a dataset, which includes nutrient  
 378 loading estimates from GlobalNEWS (Mayorga et al., 2010) and the Integrated Model  
 379 to Assess the Global Environment-Global Nutrient Model (IMAGE-GNM) (Beusen et  
 380 al., 2016; Beusen et al., 2015). Nutrient inputs are provided for dissolved inorganic ni-  
 381 trogen (DIN), phosphorus (DIP), Si and Fe, as well as dissolved organic nitrogen and  
 382 phosphorus. Carbon inputs are provided in inorganic and organic forms. Organic river-  
 383 ine inputs are partitioned into MARBL’s corresponding semi-labile and refractory or-  
 384 ganic matter tracers; the fractions routed into the refractory tracers are 0.2, 0.1, and 0.025  
 385 for carbon, nitrogen, and phosphorus, respectively. Riverine DIC inputs are assumed to  
 386 be comprised of 100% bicarbonate and thus alkalinity fluxes are equal to DIC fluxes. Par-  
 387 ticulate constituents are assumed to be removed in estuaries and therefore are not in-  
 388 cluded in the riverine forcing supplied to the model. GlobalNEWS does not include dis-  
 389 solved iron inputs to the oceans; for the riverine Fe source we assumed a constant river  
 390 concentration of 10 nM in a climatological runoff for the current era. Riverine nutrients  
 391 and carbon fluxes are held constant using the GlobalNEWS data with the exception of  
 392 DIN and DIP fluxes, which are taken from IMAGE-GNM and vary from 1900 through  
 393 2000; outside of this period, the fluxes are held constant using the closest temporal value.

394 In CESM2, the handling of riverine freshwater inputs was changed relative to pre-  
 395 vious versions of the model. In particular, rather than spreading riverine freshwater in-  
 396 puts out over a relatively large prescribed region of the ocean surface, CESM2 routes river-  
 397 ine freshwater input into the ocean column closest to location of the riverine input. A  
 398 box-model framework is used to parameterize vertical estuary exchange flow and asso-  
 399 ciated mixing of river and seawater prior to applying the input to the ocean column (Sun  
 400 et al., 2017). This estuary box-model framework is also used in CESM2 to distribute river-  
 401 ine biogeochemical inputs, ensuring consistent treatment of freshwater and biogeochem-  
 402 ical riverine inputs, and also avoiding the unrealistic spreading of inputs over a large re-  
 403 gion of the surface ocean.

### 404 **2.2.7 Benthic processes**

405 Riverine nutrient and carbon fluxes in CESM2 introduce material into the coupled  
 406 system without a direct compensating sink from the land model. To avoid drift in ocean  
 407 nutrient inventories and spurious accumulation of carbon in the atmosphere, ocean losses  
 408 must balance riverine inputs. The marine nitrogen cycle in MARBL is open, capable of  
 409 achieving a dynamic equilibrium as nitrogen fixation and denitrification come into bal-  
 410 ance with the other supply terms. Carbon, phosphorus, and silica inputs, by contrast,  
 411 are balanced with the process of burial at the seafloor.

412 MARBL computes burial and denitrification losses of material at the seafloor ac-  
 413 cording to empirical relationships. Particulate organic carbon burial is computed using  
 414 a relationship between burial efficiency and POC flux from Dunne et al. (2007), with an  
 415 imposed maximum burial efficiency of 80%. Burial of SiO<sub>2</sub> at the seafloor is based on  
 416 observations in Ragueneau et al. (2000). In MARBL, 4% of Si incident on the seafloor  
 417 is buried, except where the incident flux of Si to the seafloor exceeds 2 mmol m<sup>-2</sup> d<sup>-1</sup>;  
 418 then, 20% of Si is buried. As described above, sedimentary denitrification depends on

419 the incident POC flux and is computed based on an empirical relationship from Bohlen  
 420 et al. (2012). Burial of  $\text{CaCO}_3$  on the ocean floor occurs where  $\Omega > \Omega_{crit}$  in the model's  
 421 bottom layer; where  $\Omega < \Omega_{crit}$ , all  $\text{CaCO}_3$  reaching the model's bottom layer is dissolved.  
 422 All  $\text{CaCO}_3$  is assumed to be calcite, thus ignoring the distinction between the mineral  
 423 forms calcite and aragonite, which may be important in modulating dissolution depths  
 424 (Gangstø et al., 2008).

425 In order to achieve balanced global ocean tracer inventories, we impose global co-  
 426 efficients that scale burial of carbon, phosphorus, and silicon linearly following applica-  
 427 tion of the initial empirical relationships. The burial scaling factor for particulate organic  
 428 nitrogen is the same as for particulate organic carbon. These coefficients enable enforc-  
 429 ing a match between globally-integrated burial and global riverine inputs, thus enforc-  
 430 ing equilibrium in the preindustrial climate. The global burial coefficients were tuned  
 431 online in a spin-up configuration (see below), adjusting the coefficient to force burial to  
 432 match inputs on a 10-year timescale. To enable this, we compute an exponentially-weighted  
 433 moving average of each term online in the model, thus filtering out temporal variabil-  
 434 ity below the 10-year timescale, and adjusting the coefficients at each time step.  $\Omega_{crit}$   
 435 was also tuned in the spin-up runs, to ensure a balanced alkalinity inventory.

### 436 **2.2.8 Dissolved oxygen**

437 During the ocean-tracer spin-up of the CESM2 model, we found that ventilation  
 438 of the deep North Pacific was very sluggish in the coupled model, leading to severe oxy-  
 439 gen depletion over a large portion of the interior water column. The problem is evidenced  
 440 by simulated natural radiocarbon age in the deep North Pacific being twice that of ob-  
 441 servations (see Results). We were not able to alleviate the oxygen depletion by compen-  
 442 sating for the circulation bias via tuning of MARBL parameters and it was too late in  
 443 the CMIP6-driven development cycle to explore changes in the coupled model config-  
 444 uration to improve the ventilation. While ventilation biases are common in coarse res-  
 445 olution OGCMs, the North Pacific oxygen depletion in the spin up was so intense and  
 446 widespread that we were concerned about large-scale denitrification leading to extensive  
 447 loss of fixed nitrogen, which would perturb other aspects of the ocean biogeochemical  
 448 simulation. Therefore, we were forced to address the problem via ad hoc means: we im-  
 449 plemented a scale factor to reduce oxygen consumption in the North Pacific; oxygen con-  
 450 sumption is multiplied by this scale factor, which was set to 0.3 in the deep Pacific (be-  
 451 low 1500 m and north of 20°S) and 1.0 elsewhere; the scale factor changes linearly from  
 452 1.0 at 40°S to 0.3 at 20°S and similarly between 750 m and 1500 m depth. This ad hoc  
 453 scaling of oxygen consumption breaks stoichiometric relationships between oxygen and  
 454 other biogeochemical tracers, invalidating assumptions commonly made in the analysis  
 455 of biogeochemical simulations (e.g., computing preformed nutrients). In order to avoid  
 456 confusion from potential users of CESM2 CMIP6 output, we opted to withhold publi-  
 457 cation of oxygen-related fields from CESM2 CMIP6 experiments.

### 458 **2.2.9 MARBL features not enabled in CMIP6**

459 The CESM2-CMIP6 integrations include calcification simulated by the implicit cal-  
 460 cification treatment described above. Recent developments have parameterized a prog-  
 461 nostic phytoplankton calcifier in MARBL that is modeled on coccolithophore physiolo-  
 462 gy (Krumhardt et al., 2019). The ratio of calcification to photosynthesis ( $r_{\text{CaCO}_3:\text{C}}$ ) by  
 463 the coccolithophore functional type is responsive to environmental conditions, where  $r_{\text{CaCO}_3:\text{C}}$   
 464 is a function of temperature, nutrients, and  $\text{CO}_2$ . This enables a calcification response  
 465 to ocean acidification, among other environmental changes (for further details see Krumhardt  
 466 et al., 2019).

467 MARBL includes a representation of carbon isotopes, which follows on the imple-  
 468 mentation in POP by Jahn et al. (2015). The carbon isotope tracers in MARBL were

469 not enabled for the CESM2 CMIP6 integrations. An abiotic radiocarbon tracer imple-  
470 mented in POP was enabled for these runs, however (see below).

471 As mentioned above, the number and definition of plankton functional types in MARBL  
472 is flexible and can be configured at runtime via an input file. Ecosystem models with ad-  
473 ditional resolved plankton groups may be useful for coupling with models of higher trophic  
474 levels, providing a framework for understanding climate-driven variations in potential  
475 fisheries yield, for instance. The Size-based Plankton Ecological Traits (SPECTRA) ver-  
476 sion of MARBL employs allometric, i.e., size-based, scaling for various aspects of organ-  
477 ismal physiology such as metabolic rates, resource acquisition, mortality, and predator-  
478 prey interactions, while maintaining important “trait-based” functions that are impor-  
479 tant for elemental cycles (e.g., opal production by diatom groups). The resulting MARBL-  
480 SPECTRA configuration has nine phytoplankton and six zooplankton PFTs represent-  
481 ing various planktonic taxa within the 0.5  $\mu\text{m}$  to 20mm size range, allowing explicit sim-  
482 ulation of food resources for higher trophic levels.

### 483 *2.2.10 Ancillary tracers*

484 The simulated circulation of an ocean model plays a critical role in the ability of  
485 the ocean model to skillfully simulate biogeochemical tracers (Doney et al., 2004). In CESM2  
486 experiments for CMIP6, the ocean model was run with several ocean diagnostic tracers:  
487 abiotic radiocarbon, chlorofluorocarbons (CFCs), and sulfur hexafluoride ( $\text{SF}_6$ ). These  
488 tracers provide information about the ocean model’s circulation that is relevant to the  
489 simulation of biogeochemical tracers. In particular, the natural component of abiotic ra-  
490 diocarbon provides information about the ocean model’s circulation on multi-centennial  
491 and longer timescales, due to its 5730-year half-life. This information is complemented  
492 on decadal time-scales by the bomb-spike component of abiotic radiocarbon and the CFC  
493 and  $\text{SF}_6$  tracers. While these diagnostic tracers are not included in the MARBL library,  
494 we analyze some aspects of their simulated values to put the analysis of the MARBL trac-  
495 ers in the appropriate context of the simulated flow. The implementation of these diag-  
496 nostic tracers in CESM2 follows the protocols described in Orr et al. (2017). The abi-  
497 otic radiocarbon implementation is largely based on the implementation described in Jahn  
498 et al. (2015).

499 Following Orr et al. (2017), the abiotic radiocarbon tracer is implemented as two  
500 abiotic tracers,  $\text{DIC}^{\text{abio}}$  and  $^{14}\text{DIC}^{\text{abio}}$ . Because these tracers are abiotic, they are not  
501 directly comparable to observations individually. We instead compare to  $\Delta^{14}\text{C}$ , the iso-  
502 topic fractional abundance of  $^{14}\text{C}$  compared to  $^{12}\text{C}$ , corrected for biological fractiona-  
503 tion and normalized by dividing  $^{14}\text{C}/^{12}\text{C}$  by  $^{14}r_{\text{std}}$ , which is  $^{14}\text{C}/^{12}\text{C}$  from a pre-bomb  
504 standard sample. Following Orr et al. (2017), the modeled tracer  $^{14}\text{DIC}^{\text{abio}}$  is normal-  
505 ized by dividing by  $^{14}r_{\text{std}}$ . Because it is abiotic and is not linked to biological carbon cy-  
506 cling, the modeled tracer  $^{14}\text{DIC}^{\text{abio}}$  does not require a fractionation correction; there-  
507 fore, given this implementation,  $\Delta^{14}\text{C}$  for the model is computed as

$$\Delta^{14}\text{C} = 1000 \cdot \left( ^{14}\text{DIC}^{\text{abio}} / \text{DIC}^{\text{abio}} - 1 \right). \quad (14)$$

## 508 **2.3 Numerical experiments**

509 As mentioned above, the experiments described here are from the CESM2 contri-  
510 bution to CMIP6 (Eyring et al., 2016). We analyze 3 types of experiments: a preindus-  
511 trial control experiment, experiments spanning the recent historical past, and future sce-  
512 nario experiments. In the preindustrial control experiment, referred to as piControl, pre-  
513 scribed forcings used by the model repeatedly cycle through values representative of the  
514 year 1850. The initialization of the piControl experiment is described below. The exper-  
515 iments of the recent historical past, referred to as historical, were run using prescribed  
516 forcings for years 1850–2014. These experiments were initialized from the piControl ex-  
517 periment, using the model’s state at 1 January, taken from different years. We analyze

518 11 ensemble members of this type of experiment. The only difference between these en-  
 519 semble members is the year of the piControl experiment that their initial state came from.  
 520 The future scenario experiments follow the protocols of Scenario Model Intercompari-  
 521 son Project (ScenarioMIP) (O'Neill et al., 2016). The scenarios, referred to as Shared  
 522 Socioeconomic Pathways (SSPs), were generated using integrated assessment models, based  
 523 on a combination of different assumptions about societal development and target radiative  
 524 forcings. We analyze 3 ensemble members of 4 different SSPs that span a range of  
 525 anthropogenic impacts on the climate system. These experiments were initialized from  
 526 the end of different historical experiments and were run, and used prescribed forcings,  
 527 for years 2015-2100.

### 528 **2.3.1 Initialization of piControl**

529 To initialize the piControl, we applied a spin-up methodology to equilibrate bio-  
 530 geochemical tracers, including abiotic carbon and ideal age, to the simulated circulation.  
 531 The spin-up was conducted using forcing and physical state extracted from a twenty-one  
 532 year segment of a previous fully-coupled CESM2 experiment. The forcing was applied  
 533 cyclically to the ocean and sea-ice component models for spin-up, which yields a much  
 534 lower computational cost than the fully coupled system. The objective of the spin-up  
 535 was to find a quasi-steady-state tracer distribution; this entails minimizing the differ-  
 536 ence in the tracer distributions between the beginning and end of the selected twenty-  
 537 one year forcing period. The ocean physical state was reset at the beginning of each twenty-  
 538 one year cycle, keeping it synchronized with the surface forcing and eliminating drift in  
 539 temperature and salinity, for example. The spin-up was run for 1029 years. In the spin-  
 540 up, the  $\Omega_{crit}$  value (the threshold for  $\text{CaCO}_3$  burial; see above) was manually adjusted  
 541 at several points during the spin-up to ensure that loss of alkalinity from burial of  $\text{CaCO}_3$   
 542 balanced riverine input of alkalinity. The final threshold value was 0.89, which was the  
 543 value used in all subsequent experiments. Scaling coefficients applied to the burial of POM  
 544 and silica at the seafloor (see above) were automatically adjusted in order to balance burial  
 545 of carbon, phosphorus, and silicon with corresponding riverine inputs. All subsequent  
 546 experiments used the values of the scaling factors determined in the spin-up. At several  
 547 points during the spin-up, a Newton-Krylov based solver, based on (Lindsay, 2017), was  
 548 used to more completely spin up a subset of the biogeochemical tracers. This Newton-  
 549 Krylov based solver was applied to dissolved organic matter tracers (semi-labile and re-  
 550 fractory), DIC, alkalinity, abiotic carbon tracers, and ideal age. The solver was not ap-  
 551 plied to other biogeochemical tracers because it had not yet been successfully extended  
 552 to them. For each of tracers where it is applied, the Newton-Krylov approach solved di-  
 553 rectly for tracer equilibrium, assuming a fixed cyclo-stationary productivity field. The  
 554 tracers to which the Newton-Krylov solver was applied are more equilibrated than one  
 555 would expect from the duration of the spin-up. The globally-integrated air-sea  $\text{CO}_2$  flux  
 556 at the end of the spin-up was  $\sim 0.02 \text{ Pg C yr}^{-1}$ . The implication of this small air-sea  $\text{CO}_2$   
 557 flux is that riverine inputs of carbon are nearly completely balanced by sediment burial.

## 558 **3 Observational datasets**

559 We used several observationally-based datasets for model validation. Model fields  
 560 were averaged over the period 1990–2014, unless noted otherwise, and averaged across  
 561 ensemble members to assess the simulated mean-state in comparison with observations.  
 562 Model chlorophyll fields were compared to climatological seasonal chlorophyll means de-  
 563 rived from the Sea-viewing Wide Field-of-view Sensor (SeaWiFS) satellite over the pe-  
 564 riod September 1997 to December 2010, calculated using the GSM (Garver-Siegel-Maritorena  
 565 model) algorithm (Maritorena et al., 2002; Maritorena & Siegel, 2005). We compared  
 566 these to chlorophyll concentrations from the upper level of the ocean model (top 10 m).  
 567 Modeled inorganic nutrient fields were compared those from the World Ocean Atlas ver-  
 568 sion 2018 (WOA; Garcia et al., 2018). Observationally-based CFC and radiocarbon ( $\Delta^{14}\text{C}$ )

569 distributions were taken from the Global Ocean Data Analysis Project, version 1 (GLO-  
570 DAPv1) database (Key et al., 2004). The data comprising GLODAPv1 were collected  
571 over approximately the 1990s; thus, when comparing model output to these data, we use  
572 11-yr, ensemble-mean averages centered on 1995. GLODAP reports constituent concen-  
573 trations per mass of seawater. Since the CESM ocean component is volume conserving,  
574 we convert observation-based concentrations to per-volume units using a constant ref-  
575 erence density,  $\rho = 1026 \text{ kg m}^{-3}$  (no volume correction is applied to the radiocarbon  
576 isotope ratio). The simulated oceanic anthropogenic  $\text{CO}_2$  ( $C_{ant}$ ) inventory at year 2007  
577 was compared to values derived from GLODAPv1 (Key et al., 2004; Sabine et al., 2004),  
578 with a correction for carbon uptake between 1994 and 2007 from Gruber et al. (2019).  
579 We compare 1990–2014 air-sea  $\text{CO}_2$  flux to the gridded flux product of Landschützer et  
580 al. (2017), which we refer to as SOM-FFN, reflecting the two-step method described in  
581 Landschützer et al. (2016): first a self organizing map (SOM) is used to cluster the global  
582 ocean into biogeochemical provinces; second, a feed forward neural-network (FFN) is used  
583 to estimate  $p\text{CO}_2$  from driver variables (i.e. sea surface temperature) after training us-  
584 ing observations from the Surface Ocean  $\text{CO}_2$  Atlas (SOCAT; Bakker et al., 2016). The  
585 SOM-FFN  $\text{CO}_2$  fluxes are estimated from  $p\text{CO}_2$  using the same gas exchange param-  
586 eterization employed by MARBL (Sweeney et al., 2007) and wind speed derived from  
587 the National Centers for Environmental Prediction (Kalnay et al., 1996). To evaluate  
588 simulated dissolved iron (dFe) distributions, we make use of a data compilation extended  
589 from that in Moore and Braucher (2008) and including data from Tagliabue et al. (2012)  
590 and the GEOTRACES Intermediate Data Product compilation (Schlitzer et al., 2018).  
591 We compare simulated DOM distributions to a compilation of observations from Letscher  
592 and Moore (2015).

## 593 4 Results and discussion

594 In this section, we present and discuss several representative diagnostics of the CESM2  
595 solutions submitted to CMIP6. Our objectives are not to provide a comprehensive anal-  
596 ysis of these solutions, but rather we aim for a broad overview, illustrating key aspects  
597 of the ocean biogeochemical simulation and documenting important patterns in the model  
598 biases. We include a brief treatment of future projections, noting that output from CESM2  
599 is also included in several model intercomparison papers (e.g., Séférian et al., 2020; Arora  
600 et al., 2020; Kwiatkowski et al., 2020).

### 601 4.1 Mixed layer depth

602 Surface mixed layer depths are a key control on the upper ocean habitat and im-  
603 portant mediator of water mass ventilation affecting transient tracer uptake. The mixed  
604 layer depth simulation in CESM2 is therefore of interest in the context of understand-  
605 ing large-scale biogeochemical distributions and the strength of the biological pump. No-  
606 tably, the mixed layer depth in the model manifests as a result of interactions between  
607 the vertical mixing scheme (Large et al., 1994) and both parameterized (e.g., Danaba-  
608 soglu et al., 2010; Gent & McWilliams, 1990; Fox-Kemper et al., 2008) and resolved trans-  
609 port controlling stratification (Small et al., 2020). Figure 1 shows winter and summer  
610 distributions of mixed layer depth in CESM2 historical simulations compared with an  
611 observational estimate. To approximate the mixed layer depth, we use monthly-mean  
612 salinity and temperature to compute potential density; the mixed layer depth is calcu-  
613 lated as the depth at which potential density changes by  $0.125 \text{ kg m}^{-3}$  from its surface  
614 value. The same procedure is applied to the World Ocean Atlas observationally-based  
615 product (Locarnini et al., 2019; Zweng et al., 2019). The model shows broad agreement  
616 with the observations in terms of the large-scale distribution of mixed layer depth (Fig-  
617 ure 1 left two columns), but includes some important biases. Notably, high-latitude sum-  
618 mertime mixed layers tend to be too deep in the model (Figure 1C), including regions  
619 with very deep biases in the North Atlantic and Southern Ocean. The wintertime mixed

**Table 3.** Global means fluxes.

<b>Flux or Concentration</b>	<b>preindustrial (CESM1)</b>	<b>1990 - 2014 (CESM1)</b>	<b>RCP 8.5 2090s (CESM1)</b>	<b>preindustrial (CESM2)</b>	<b>1990-2014 (CESM2)</b>	<b>SSP5-8.5 2090s (CESM2)</b>
Net primary production, full depth (PgC/yr)	55.9 <sup>a</sup>	56.1 <sup>a</sup>	54.0 <sup>a</sup>	48.2	48.8	49.8
Sinking POC at 100 m (PgC/yr)	8.07	7.98	7.20	6.98	7.05	6.69
Sinking CaCO <sub>3</sub> at 100 m (PgC/yr)	0.757	0.748	0.723	0.767	0.767	0.808
Sinking SiO <sub>2</sub> at 100 m (Tmol/yr)	-	-	-	77.6	78.2	69.8
Rain ratio (CaCO <sub>3</sub> /POC) at 100 m	0.094	0.094	0.100	0.110	0.109	0.121
Nitrogen fixation (TgN/yr)	175	169	144	241	243	285
Nitrogen deposition (TgN/yr)	6.6	29.6	30.0	13.3	37.2	38.3
Water Column Denitrification (TgN/yr)	190	194	188	185	192	256
Sediment Denitrification (TgN/yr)	-	-	-	67	71	68
Nitrogen Burial to Sediment (TgN/yr)	-	-	-	24	27	22
Nitrogen surface emissions (TgN/yr)	-	-	-	6	5	3
Nitrogen River Flux (TgN/yr)	-	-	-	13	25	25
N cycle imbalance <sup>b</sup> (TgN/yr)	-8.0	5.2	-13.8	-14.1	10.3	-0.8
Air-sea CO <sub>2</sub> flux (PgC/yr)	-0.02	2.03	4.71	-0.04	2.04	5.33
Diatom primary production, top 100m (%)	34	34	32	35	37	31
Diatom primary production, full depth (%)	35 <sup>a</sup>	35 <sup>a</sup>	32 <sup>a</sup>	36	37	31

<sup>a</sup> NPP was only reported to 150m depth in CESM1

<sup>b</sup> Deposition + Fixation - Denitrification - Burial - Emissions + River Flux

620 layer depth distributions are characterized by heterogeneous biases in the North Atlantic  
 621 (Figure 1F), though these biases are improved relative to older model versions (i.e., CCSM3)  
 622 that lacked a parameterization of density driven overflows representing Denmark Strait  
 623 and Faroe Bank Channel (Danabasoglu et al., 2012). Mixed layer depths are too shal-  
 624 low in the eastern Subpolar North Atlantic and Greenland, Iceland and Norwegian (GIN)  
 625 Seas, but too deep in the Labrador Sea. Wintertime mixed layer depths are too shallow  
 626 in the Subantarctic along the northern flank of the Antarctic Circumpolar Current (ACC)  
 627 (Figure 1F). These biases are likely attributable to the representation of horizontal ad-  
 628 vection and insufficient transport of warm, salty subtropical waters into the ACC region  
 629 (Small et al., 2020). The biases in the Subantarctic likely restrict Southern Ocean up-  
 630 take of transient tracers (next section). Furthermore, the mixed layer depth simulation  
 631 affects the seasonal evolution of NPP and air-sea CO<sub>2</sub> fluxes. For example, since iron  
 632 is a critical control on Southern Ocean phytoplankton blooms, substantial biases in the  
 633 mixed layer depth may result in poor simulation of iron supply, which is thought to be  
 634 mediated largely by seasonal entrainment (Tagliabue et al., 2014).

## 635 4.2 Ventilation tracers

636 We examined the simulated distribution of three tracers: CFC-11, total radiocar-  
 637 bon (i.e., natural plus the “bomb-spike”;  $\Delta^{14}\text{C}$ ), and anthropogenic CO<sub>2</sub> ( $C_{ant}$ ). CFC-  
 638 11 and  $\Delta^{14}\text{C}$  provide a means to assess the simulated circulation and potential biases  
 639 in ventilation; as noted above, these tracers are simulated in the CESM2 ocean compo-  
 640 nent, not within MARBL. However, we include them here as circulation biases provide  
 641 an important context for understanding biogeochemical distributions. Since CFC-11 up-  
 642 take is concentrated in water masses with ventilation ages of up to several decades (e.g.,  
 643 Dutay et al., 2002), biases in CFC-11 uptake are predominately informative of ventila-  
 644 tion in thermocline waters. In order to isolate the influence of ventilation processes on  
 645 CFC-11, we examine the partial pressure of CFC-11 ( $p\text{CFC-11}$ ), thereby removing the  
 646 influence of temperature and salinity biases on the tracer distribution.  $\Delta^{14}\text{C}$ , by con-  
 647 trast, provides an indication of the fidelity of deep ocean overturning circulation on cen-  
 648 tennial timescales (e.g., Matsumoto, 2007). Rather than examine radiocarbon-derived  
 649 estimates of circulation age, we simply present  $\Delta^{14}\text{C}$  distributions directly as a qualita-  
 650 tive indication of ventilation biases.

651  $p\text{CFC-11}$  highlights significant biases in thermocline ventilation in CESM2, with  
 652 deficits in thermocline waters in both the Pacific and Atlantic basins (Figure 2C, F) and  
 653 evidence of too much uptake in North Atlantic Deep Water (NADW; Figure 2C). These  
 654 biases are largely similar to those found in CESM1 simulations (Long et al., 2013). Larger  
 655 than observed values of  $\Delta^{14}\text{C}$  are evident in the North Atlantic, confirming indications  
 656 of vigorous NADW formation evident in  $p\text{CFC-11}$  (Figure 3C). The most dramatic as-  
 657 pect of the  $^{14}\text{C}$  simulation, however, is the very large  $\Delta^{14}\text{C}$  deficit in the deep North Pa-  
 658 cific (Figure 3F); this illustrates the sluggish circulation simulated by the coupled model  
 659 in this region. As described above, the CESM2 simulation of  $^{14}\text{C}$  is abiotic, so does not  
 660 represent the vertical transfer of  $^{14}\text{C}$  accomplished by sinking organic matter. The in-  
 661 clusion of biology would cause the simulated deep ocean  $\Delta^{14}\text{C}$  to be less depleted, re-  
 662 ducing the magnitude of the apparent bias; however, this effect is expected to quite small  
 663 (e.g., 25‰) relative to the magnitude of the bias (Jahn et al., 2015). These deep cir-  
 664 culation biases are much worse than in previous versions of the model (i.e., CESM1). No-  
 665 tably, testing indicates that running the CESM2 ocean component with reanalysis forc-  
 666 ing restores the deep ocean circulation. Since changes to the physical ocean component  
 667 between CESM1 and CESM2 were quite minimal, we thus attribute this bias to changes  
 668 in the atmosphere component, though the specific mechanism remains under investiga-  
 669 tion. Oddly, Heuzé (2021) determined that CESM2 was one of the best performing mod-  
 670 els in terms of AABW and NADW formation; this study, however, was based on met-  
 671 rics of deep convection and watermass properties; it did not include an examination of  
 672 transient tracers.

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### 4.3 Anthropogenic CO<sub>2</sub> inventory

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Ventilation tracers are useful in the context of understanding biases in the uptake of  $C_{ant}$ , since  $C_{ant}$  uptake is largely mediated by overturning circulation. To compute  $C_{ant}$ , we make use of the two parallel carbonate systems simulated by MARBL. In the CESM2 CMIP6 integrations, these tracer systems are subject to identical transport and source/sink terms; they differ only in their atmospheric CO<sub>2</sub> boundary conditions: the primary DIC tracer is exposed to increasing CO<sub>2</sub> according to historical or scenario forcing, while the secondary DIC tracer, “DIC\_ALT\_CO2”, is forced with a constant preindustrial (1850) value for atmospheric CO<sub>2</sub> (284.7 ppm). We thus define anthropogenic CO<sub>2</sub> as

$$C_{ant} = DIC - DIC\_ALT\_CO2$$

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Note that this definition differs subtly from subtracting a DIC field obtained from an 1850-control integration: changes in climate impact both DIC and DIC\_ALT\_CO2, so the resulting  $C_{ant}$  does not include the impact of climate on natural CO<sub>2</sub> as it would if the baseline DIC field were taken from an 1850-control integration. While climate impacts on natural CO<sub>2</sub> are significant under future scenarios with strong radiative forcing, this feedback is modest over the historical period.

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CESM2 simulates weaker  $C_{ant}$  uptake than suggested by observational estimates (Figure 4). The model has  $C_{ant}$  deficiencies of order 10 mmol m<sup>-3</sup> evident broadly in the thermocline waters. The total GLODAP-based observational estimate for the  $C_{ant}$  inventory at year 2007 is 152±19 Pg C (Gruber et al., 2019; Sabine et al., 2004); the CESM2-simulated inventory at this time is 114 Pg C, or roughly 75% of the observations. Note that the GLODAP based observations to which we are comparing omit coverage in some oceanic regions (i.e., the Gulf of Mexico, the Arctic Ocean). The CESM2 simulated inventory for the portion of the ocean included in the gridded observational product is 106 Pg C at year 2007, compared with ~137 Pg C for the observationally-based  $C_{ant}$  estimates. A caveat with this comparison is that the model  $C_{ant}$  field only includes carbon uptake since 1850. As noted in Lindsay et al. (2014), Figure 3 of Khatiwala et al. (2009) indicates pre-1850 anthropogenic ocean uptake to be about 12±3 Pg C. Taking this into account reduces, but does not eliminate the model’s low-uptake bias.

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### 4.4 Macronutrients

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A primary objective of MARBL is to represent the structure and function of the biological pump. The biological pump is fueled by nutrients; export of these constituents from the surface ocean via sinking and dissolved organic matter plays a dominant role in structuring nutrient distributions. Macronutrients are reasonably well-observed in the ocean and thus provide a good constraint on model performance. Figure 5 presents a comparison of surface NO<sub>3</sub>, PO<sub>4</sub>, and SiO<sub>3</sub> distributions from CESM2 with WOA observations. The overall geographical patterns of these macronutrients are well represented in CESM2, with increased surface nutrients at the high latitudes and equatorial regions and low nutrient concentrations in subtropical gyre regions. However, the simulated surface nutrient concentrations are too high in the subtropical oligotrophic gyres and too low in the Subarctic North Pacific; furthermore, there are substantial biases in the Southern Ocean. Simulated surface NO<sub>3</sub> and PO<sub>4</sub> are both too low in the Southern Ocean, by ~4–8 mmol m<sup>-3</sup> and ~0.25–0.55 mmol m<sup>-3</sup>, respectively. Conversely, SiO<sub>3</sub> concentrations are too low in the Antarctic zone and too high in the Subantarctic. Surface nutrients reflect the balance of net community production (NCP) and physical supply, which together determine the extent of nutrient utilization. Thus, negative biases in surface nutrients in the Southern Ocean suggest that the model may over estimate NCP and the simulated algal community composition includes an insufficient contribution from diatoms—or the Si to N stoichiometric ratio of diatom production, which is higher under extensive Fe limitation (Moore et al., 2004), may be too low. Alternatively, the nutrient content of water masses upwelling in the Southern Ocean may already be too low—indicative

724 of a large-scale bias in the nutrient simulation that may be related to excessive trapping  
725 of nutrient in the North Pacific.

726 A latitude-depth view of nutrient distributions lends credence to this latter hypoth-  
727 esis. Figures 6–8 show zonal-mean, depth-latitude plots of macronutrients in the Atlantic  
728 and Pacific Ocean basins, indicating how the vertical structure of the simulated macronu-  
729 trients compares with observations. While the simulated nutrient distributions show over-  
730 all structure that is similar to the observations, several key biases indicate deficiencies  
731 in the CESM2 solutions. Most notably, these include  $\text{NO}_3$  depletion in the tropical ther-  
732 mocline (Figure 6C, F), which is driven by denitrification in overly extensive oxygen de-  
733 ficient zones, and excessive accumulation of macronutrients in the deep North Pacific,  
734 attributable to sluggish deep circulation in this region.

735 The vertical structure of  $\text{PO}_4$  provides an indication of the functioning of the bi-  
736 ological pump without the complications of denitrification; the zonal means of simulated  
737  $\text{PO}_4$  indicate excessive nutrient concentrations in the tropical thermocline, particularly  
738 in the Atlantic, and nutrient trapping the deep North Pacific (Figure 7). The negative  
739 bias in nitrate and phosphate in the Southern Ocean surface waters (Figure 5) is also  
740 evident in the ocean interior over much of the Southern Hemisphere water column (Fig-  
741 ure 7C, F). This pattern demonstrates that the whole Southern Ocean nutrient inven-  
742 tory is too low in the model, suggesting that too weak supply of nutrients to the surface  
743 ocean via upwelling in this region is partially responsible for negative surface nutrient  
744 biases.

745 The situation is subtly different for silicate, which displays a positive surface bias  
746 over much of the Southern Ocean (Figure 5I)—but a dipole bias pattern in the zonal-  
747 mean column view (Figure 8C, F). This pattern is characterized by negative biases in  
748 the region associated with AABW and the deep overturning cell, but positive biases in  
749 upper, equatorward portion of the column associated with the upper cell and where Antarc-  
750 tic Intermediate Water and Subantarctic Mode water are formed. These patterns indi-  
751 cate that while  $\text{SiO}_3$  supply via upwelling to Southern Ocean surface waters may be too  
752 weak, opal production is also too weak, resulting in excessive leakage of  $\text{SiO}_3$  from the  
753 Southern Ocean (*sensu* Sarmiento et al., 2004). Sarmiento et al. (2007) demonstrated  
754 that the Southern Ocean effectively traps silica (see also Primeau et al., 2013; Moore et  
755 al., 2018), a phenomena we have also demonstrated operates in POP for  $\text{CaCO}_3$  and al-  
756 kalinity (Krumhardt et al., 2020). In this vein, excessive silica leakage from the South-  
757 ern Ocean in CESM2 may help explain why upper-ocean  $\text{SiO}_3$  concentrations are too  
758 high at the surface over much of the rest of the global ocean, with the exception of the  
759 North Pacific (Figure 8).

#### 760 4.5 Nutrient limitation

761 Nutrients and temperature play important roles in structuring phytoplankton pro-  
762 ductivity throughout the global ocean. The most limiting nutrients for each phytoplank-  
763 ton functional type are shown in Figure 9; these fields are computed as phytoplankton-  
764 biomass-weighted means of the upper-ocean limitation terms, thereby providing a pic-  
765 ture of resource limitation relevant to understanding vertically integrated production.  
766 Small phytoplankton in CESM2 are limited by N in much of the low to mid-latitudes,  
767 except in the South Pacific, where Fe is primarily limiting growth. Indeed Fe limits pro-  
768 duction of small phytoplankton and diatoms in most oceanic regions south of  $15^\circ\text{S}$ . These  
769 major patterns of phytoplankton N and Fe limitation are generally supported by obser-  
770 vations (e.g., Moore et al., 2013). Diatoms are limited by  $\text{SiO}_3$  in regions bordering the  
771 continents in the Southern Hemisphere, as well as in the North Pacific and North At-  
772 lantic in CESM2. Diazotrophs fix N and are therefore not limited by N availability; they  
773 are limited by Fe and P in the tropics and subtropics (Sañudo-Wilhelmy et al., 2001; Letscher  
774 & Moore, 2015); temperature limits the geographic distribution of diazotrophs to warmer

775 waters ( $>15^{\circ}\text{C}$ ; Figure 9C). P limitation is mainly evident in the low latitude North At-  
 776 lantic for all three phytoplankton functional types; here increased Fe deposition from Sa-  
 777 haran aeolian fluxes stimulate N fixation by diazotrophs making P the limiting nutri-  
 778 ent (Wu et al., 2000; Sañudo-Wilhelmy et al., 2001).

#### 779 4.6 Surface chlorophyll

780 Satellite-derived observations of chlorophyll provide a proxy for phytoplankton dis-  
 781 tribution and biomass. Here, we compare seasonal (DJF and JJA) mean chlorophyll con-  
 782 centrations from SeaWiFS to CESM2 chlorophyll concentrations in the surface ocean (Fig-  
 783 ure 10). While CESM2 simulates the overall patterns of chlorophyll distribution during  
 784 the Southern Hemisphere growing season, it is markedly too high for certain regions (Fig-  
 785 ure 10A–C). Strong positive biases are evident in the Subantarctic region of the South-  
 786 ern Ocean, especially in the Atlantic sector and south of Australia. Chlorophyll concen-  
 787 trations in the eastern equatorial Pacific are also overestimated by CESM2. Despite a  
 788 geographic pattern that is similar to the observations, Northern Hemisphere summer chloro-  
 789 phyll concentrations are also too high in the model for the North Pacific and North At-  
 790 lantic, according to the SeaWiFS observations (Figure 10D–F). Another prevalent bias  
 791 apparent in the model is in the coastal regions (Figure 10). Coastal phytoplankton pro-  
 792 duction and chlorophyll concentrations are commonly underestimated in nominal 1-degree  
 793 (or lower) resolution GCMs (Laufkötter et al., 2015)—though it is also the case that the  
 794 satellite observations may overestimates chlorophyll in the coastal zone (e.g., Gregg &  
 795 Casey, 2004).

#### 796 4.7 Net primary productivity and export

797 Primary production by marine phytoplankton is the ultimate constraint on the strength  
 798 of the biological pump and also forms the ecological base of the ocean food web. Satel-  
 799 lite observation-based estimates of globally integrated net primary production (NPP)  
 800 typically fall within the range of 43–67 Pg C year<sup>-1</sup> (Behrenfeld et al., 2005; Behren-  
 801 feld & Falkowski, 1997). Globally integrated NPP in CESM2 is 48.9 Pg C yr<sup>-1</sup> over the  
 802 period 1990–2014 (Table 3, Figure 11), within the range of satellite-based estimates. The  
 803 distribution of NPP in CESM2 follows a familiar pattern, with the highest rates of NPP  
 804 in equatorial upwelling regions (Figure 11A); however, the NPP difference between the  
 805 subtropics and extra-tropic appears less pronounced than that evident in satellite-based  
 806 estimates (e.g., Behrenfeld et al., 2005). Simulated globally-integrated particulate ex-  
 807 port at 100 m for the same period is 7.1 Pg C yr<sup>-1</sup> (Table 3, Figure 11), which is also  
 808 broadly consistent with observationally-based estimates, considering uncertainty (e.g.,  
 809 Henson et al., 2011; Siegel et al., 2014; Boyd & Trull, 2007). The distribution of partic-  
 810 ulate export at 100 m has greater spatial variability than NPP (Figure 11B), as it re-  
 811 flects a combination of NPP and controls on export mediated by phytoplankton com-  
 812 munity composition. Indeed, the global mean particulate export ratio (pe-ratio = sink-  
 813 ing export/NPP) in CESM2 is about 0.15, but varies by more than a factor of 3 (Fig-  
 814 ure 11C). Mineral ballasting by CaCO<sub>3</sub> and opal contributes to high pe-ratios (Armstrong  
 815 et al., 2002; Lima et al., 2014), with the regions of high opal export, indicative of diatom-  
 816 dominated assemblages, corresponding to the locations of high pe-ratio (Figure 12). No-  
 817 tably, the elemental composition of exported organic matter varies in MARBL. N:P con-  
 818 centrations are elevated below the subtropical oligotrophic gyres, which is broadly con-  
 819 sistent with observations (e.g., Martiny et al., 2013) and inverse model results (Wang  
 820 et al., 2019). These patterns are enabled by the variable P:C uptake ratios.

#### 821 4.8 Air-sea CO<sub>2</sub> fluxes

822 Figure 13 shows a comparison of CESM2 simulated air-sea CO<sub>2</sub> (1990–2014), as  
 823 well as an observationally-based flux estimate (Landschützer et al., 2017). In general,

824 the simulated flux field compares quite well with the observations (Figure 13A, B). The  
 825 globally-integrated flux is larger in magnitude ( $-2.0 \text{ Pg C yr}^{-1}$ ) than in the observational  
 826 product ( $-1.4 \text{ Pg C yr}^{-1}$ ), though this discrepancy is not significant if riverine carbon  
 827 inputs are properly accounted for in the comparison. In nature, riverine carbon fluxes  
 828 to the ocean induce net outgassing due to an imbalance between inputs and burial at  
 829 the seafloor (Gruber et al., 2009); recent estimates suggest this flux is about  $0.45\text{--}0.78 \text{ Pg C yr}^{-1}$   
 830 (Resplandy et al., 2018). Recall from above, however, that the CESM2 spin-up method-  
 831 ology aimed to achieve a balance between riverine inputs and burial of carbon at the seafloor  
 832 and near-zero net air-sea flux in the preindustrial state. Considering this methodology  
 833 then, a correction for riverine fluxes would suggest that the model’s simulated globally-  
 834 integrated air-sea  $\text{CO}_2$  flux is indistinguishable from that implied by the observations.  
 835 The model simulates outgassing in the tropics, most notably in the equatorial Pacific,  
 836 and  $\text{CO}_2$  uptake at mid- and high-latitudes (Figure 13A, B). The seasonal cycle of zonal-  
 837 mean  $\text{CO}_2$  flux is also well simulated—though notable differences in the seasonal evo-  
 838 lution of the fluxes are evident in the Southern Ocean poleward of  $45^\circ\text{S}$  (Figure 13C, D).  
 839 This region is challenging to model, in part because the net air-sea  $\text{CO}_2$  flux manifests  
 840 as the residual between opposing thermal and biologically-driven tendencies (e.g., Mongwe  
 841 et al., 2018).

#### 842 **4.9 Dissolved organic matter**

843 A fraction of plankton loss terms are routed to the DOM pool, which is subsequently  
 844 transported by circulation, and degraded by microbial activity. DOM thus provides a  
 845 transport pathway for the export of biogeochemical constituents to the deep ocean. In  
 846 particular, it is thought to account for  $\sim 20\%$  of total carbon export (Hansell, 2013). In-  
 847 deed, in CESM2 DOC export across 100 m is  $1.91 \text{ Pg C yr}^{-1}$  during 1990–2014, which  
 848 accounts for 21% of the total organic carbon flux across 100 m (particulate flux is  $7.1 \text{ Pg C yr}^{-1}$ ,  
 849 Figure 11B). Figure 14 shows upper 100 m total DOM (i.e., semi-labile plus refractory)  
 850 concentrations simulated by CESM2 as well as observations of these constituents com-  
 851 piled by Letscher and Moore (2015). The distribution of DOM partially reflects surface  
 852 water residence times, with high concentrations accumulating within the permanently  
 853 stratified oligotrophic ocean and lower surface concentrations found within the equato-  
 854 rial and Southern Ocean upwelling regions. The simulated global maxima in surface DOC  
 855 ( $76\text{--}96 \mu\text{M}$ ) and DON ( $5.5\text{--}6.5 \mu\text{M}$ ) concentrations are found in the tropical to subtrop-  
 856 ical Atlantic and Indian Ocean basins, in agreement with the observations. Simulated  
 857 global maxima in surface DOP concentrations ( $0.23\text{--}0.28 \mu\text{M}$ ) are found in the subtrop-  
 858 ical Pacific Ocean and northern Indian basin, while the global minimum is found in the  
 859 subtropical North Atlantic Ocean ( $<0.08 \mu\text{M}$ ), also in agreement with the observations.  
 860 Important to capturing the global minimum in surface DOP within the subtropical North  
 861 Atlantic is the ability for phytoplankton to use semi-labile DOP as a P source for growth  
 862 when inorganic phosphate is scarce—as is the case for the North Atlantic subtropical gyre  
 863 (Figure 5). Figure 15 presents two-dimensional histograms, showing the relationship be-  
 864 tween simulated and observed DOM. Very high observed DOM concentrations are likely  
 865 to come from regions with significant riverine influence that may not be adequately cap-  
 866 tured by the model and the supplied riverine DOM forcing. The global-mean elemen-  
 867 tal composition of total DOM in CESM2 is 385:29:1 (C:N:P) in the upper 100 m, sam-  
 868 pled at the same locations as the DOM observations, which have a mean elemental ra-  
 869 tio of 735:60:1 (Letscher & Moore, 2015). This indicates that the MARBL DOM stoi-  
 870 chiometry is P-enriched relative to the observations, while the simulated C:N stoichiom-  
 871 etry is relative close to the observed values (MARBL C:N = 13.3; observed C:N = 12.3).

#### 872 **4.10 Nitrogen cycle**

873 Globally-integrated nitrogen fixation over the 1990–2014 period averaged  $243 \text{ Tg}$   
 874  $\text{N yr}^{-1}$  (Table 3, Figure 16A); this number is larger than observationally-based estimates,

875 which range from about 100–230 Tg N yr<sup>-1</sup> (e.g., Zehr & Capone, 2021; Wang et al.,  
 876 2019). Simulated water column and sedimentary denitrification over this period were 192  
 877 and 71 Tg N yr<sup>-1</sup>, respectively. Burial of N at the sea floor and surface emissions of am-  
 878 monia account for additional losses from the model, while atmospheric deposition and  
 879 riverine inputs provide additional sources. Collectively, these terms lead to an N cycle  
 880 imbalance of about 10 Tg N yr<sup>-1</sup> (Table 3). Unfortunately, the CESM2 simulation of  
 881 dissolved oxygen is inadequate (see above) strongly influencing simulated water-column  
 882 denitrification; therefore, CESM2 cannot be reliably used to study changes to N cycle  
 883 processes with climate.

#### 884 4.11 Iron cycle

885 Figure 17 illustrates the spatial distribution of the dominant terms in the global  
 886 ocean iron budget as simulated in CESM2. Iron is supplied to the ocean via prognos-  
 887 tic atmospheric deposition (Figure 17A), with a pattern reflecting proximity to conti-  
 888 nental dust sources. The dominant source of iron to the ocean is from marine sediments  
 889 (Figure 17B) with additional contributions from hydrothermal vents (Figure 17C). River-  
 890 ine input of iron is small, accounting for only 0.37 Gmol yr<sup>-1</sup>. Iron is removed from the  
 891 ocean via burial at the seafloor (Figure 17D). The sources and sinks of dissolved iron in  
 892 the ocean are highly uncertain and model intercomparison activities have demonstrated  
 893 that it is possible to produce realistic dissolved iron concentration fields with very dif-  
 894 ferent inputs and loss terms (Tagliabue et al., 2016). Despite this caveat, there is use-  
 895 ful information in a comparison of simulated dissolved iron concentrations to observa-  
 896 tions (Figures 18 and 19).

897 CESM2 captures the dominant structure of dissolved iron reasonably well, with el-  
 898 evated surface concentrations in the tropical Atlantic and much lower concentrations in  
 899 the Pacific. Surface concentration of dissolved iron in the North Pacific appear to be too  
 900 high in the model, which might be attributable to overly diffusive flow and unrealistic  
 901 transport of sedimentary sources offshore. Indeed, we have found that iron supply in the  
 902 North Pacific changes dramatically when integrating the model at high resolution (Harrison  
 903 et al., 2018). Figure 19 shows global histograms of the model and observations; these plots  
 904 indicate that CESM2 does a good job simulating the range of iron concentrations ob-  
 905 served in the ocean, though does not have sufficient representation of waters with very  
 906 low concentrations in the upper ocean. This bias is exacerbated at depth; below 500 m,  
 907 the observations indicate that iron concentrations should be approximately normally-  
 908 distributed with a median concentration of about 0.6 nM—but CESM2 simulates some-  
 909 thing closer to a uniform distribution (Figure 19C), which is likely tied to scavenging rates  
 910 and ligand dynamics.

#### 911 4.12 Transient simulations

912 Here we include a brief discussion of future scenario integrations conducted with  
 913 CESM2. Space limitations preclude a comprehensive analysis, so we present only a few  
 914 key metrics. Figure 20A shows the prescribed atmospheric CO<sub>2</sub> forcing for each of the  
 915 future scenario integrations and Figure 20B illustrates the associated change in sea sur-  
 916 face temperature (SST). The scenarios range from SSP1-2.6—under which CESM2 projects  
 917 about 1.7°C of SST warming at 2100 relative to preindustrial—to SSP5-8.5, under which  
 918 CESM2 projects SST warming of about 4.6°C. CESM2 has an equilibrium climate sensi-  
 919 tivity (ECS, the change in surface temperature for a doubling of atmospheric CO<sub>2</sub> above  
 920 preindustrial at equilibrium) of 5.2°C and—more relevant to the simulations at hand—  
 921 a transient climate sensitivity (TCS, the surface temperature warming around the time  
 922 of CO<sub>2</sub> doubling in a 1% per year CO<sub>2</sub> increase simulation) of 2.0°C. The CMIP6 multi-  
 923 model mean for these quantities is 3.7±1.1 (standard deviation) for ECS and 2.0±0.4  
 924 for TSC (Meehl et al., 2020) and, indeed, CESM2 simulates transient warming close to  
 925 the multi-model mean (Brunner et al., 2020). The transient behavior of globally-integrated

926 air-sea CO<sub>2</sub> flux is shown in Figure 20C as a function of time and Figure 20D as a func-  
927 tion of atmospheric CO<sub>2</sub>. Importantly, for the two high-CO<sub>2</sub> scenarios (SSP3-7.0 and  
928 SSP5-8.5), the ocean CO<sub>2</sub> uptake saturates (and under SSP5-8.5 even begins to decline),  
929 in spite of continually increasing atmospheric CO<sub>2</sub>. This behavior is indicative of feed-  
930 back; in particular, as the ocean carbon inventory increases, so does the Revelle Factor,  
931 limiting additional uptake (e.g., Schwinger et al., 2014). Furthermore, climate warming  
932 increases buoyancy stratification in the upper ocean, thereby reducing vertical exchange  
933 and the transfer of excess carbon into the ocean interior. Air-sea CO<sub>2</sub> flux in the two  
934 lower CO<sub>2</sub> scenarios (SSP1-2.6 and SSP2-4.5) shows somewhat distinct behavior, as these  
935 scenarios both include a reduction in the atmospheric CO<sub>2</sub> growth rate—in SSP1-2.6,  
936 there is actually a negative trend in atmospheric CO<sub>2</sub> beyond year 2070 (Figure 20A).  
937 As the atmospheric CO<sub>2</sub> forcing is relaxed, ocean CO<sub>2</sub> uptake declines; notably, in SSP1-  
938 2.6, this decline means that air-sea CO<sub>2</sub> uptake is weaker for the same atmospheric CO<sub>2</sub>  
939 mixing ratio than in the historical period (Figure 20D). This phenomenon has impor-  
940 tant implications for managing climate mitigation; CO<sub>2</sub> previously absorbed by the ocean  
941 drives hysteresis, meaning that the ocean sink will decline in efficiency as emissions-reductions  
942 slow the atmospheric growth rate.

943 The CESM2 transient integrations indicate contrasting projections for globally in-  
944 tegrated NPP and export (Figure 21). NPP is simulated to increase over the historical  
945 period and continue increasing in the future, particularly under the high emission sce-  
946 nario SSP5-8.5. However, CESM2 projects a reduction in globally integrated POC flux  
947 at 100 m for all future scenarios (Figure 21B). These global changes reflect spatially het-  
948 erogeneous patterns. The North Atlantic is projected to undergo strong reductions in  
949 NPP and export production, while NPP is projected to increase over much of the rest  
950 of the ocean (except in the western tropical Pacific and Indian basins) (Figure 22). Crit-  
951 ically, shifts in algal community composition are a significant driver of changes in the  
952 pe-ratio. Globally, there is a decline in the pe-ratio under all warming scenarios, as well  
953 as a shift in algal community composition leading to diminished diatom prevalence  
954 relative to small phytoplankton. These changes are also spatially heterogeneous, however,  
955 as diatoms decline over most of the ocean except, most notably, in the Southern Ocean,  
956 where their relative prevalence increases dramatically (diatom fractions also increase in  
957 the equatorial Pacific) (Figure 22C, D). These patterns are broadly consistent with re-  
958 sults from CMIP5, in which models capable of transitioning production from large phy-  
959 toplankton (diatoms) to small phytoplankton projected only weak reductions in NPP,  
960 but significant changes in export (Fu et al., 2016; Bopp et al., 2013). As noted in Laufkötter  
961 et al. (2015), model differences in NPP reflect different choices in treatment of temper-  
962 ature sensitivity and model nutrient dynamics, linked to the sensitivity of nutrient sup-  
963 ply to stratification. Most CMIP5 models projected decreasing globally integrated NPP  
964 ranging from  $-1\%$  to  $-15\%$  under RCP8.5 (Bopp et al., 2013); CMIP6 models do not  
965 show as much of a decline in NPP (Kwiatkowski et al., 2020).

## 966 5 Conclusion

967 We have presented a technical description of MARBL, which is the ocean biogeo-  
968 chemistry component for the CESM2. MARBL is a state-of-the-art global ocean biogeo-  
969 chemistry model, with a prognostic representation of the coupled cycles of nitrogen, phos-  
970 phorus, silicon, iron, carbon and oxygen. The model is built on a plankton functional  
971 type paradigm, and supports flexible ecosystem configuration. In addition to document-  
972 ing MARBL, we have presented and discussed diagnostics to evaluate the ocean biogeo-  
973 chemistry simulations in the fully-coupled CESM2 integrations submitted to CMIP6. Our  
974 analysis highlights challenges associated with the development and application of ocean  
975 biogeochemical models in the context of coarse resolution OGCMs. The CESM2 solu-  
976 tions suffer from weak thermocline ventilation, which produces overly extensive oxygen  
977 minimum zones and weakens uptake of transient tracers. Deep ocean circulation in CESM2

978 is also sluggish—so much so, in fact, that we had to artificially reduce oxygen consumption  
979 in the North Pacific to prevent widespread anoxia in the model. Unfortunately, the  
980 severity of this bias required several hundred years of integration to become clear, making  
981 it a challenging issue to address in the context of model development. While biases  
982 in circulation impose limits on the fidelity of model solutions, in general MARBL captures  
983 large-scale biogeochemical distributions reasonably well, and provides a platform  
984 for researching interactions between climate, nutrient and carbon cycling in the ocean.

985 MARBL has been explicitly designed to facilitate coupling with multiple OGCMs,  
986 an effort motivated in part by interest in engaging a broad research community. This  
987 capacity is being exercised currently: we presented solutions from MARBL integrated  
988 in POP2, and we have implemented MARBL in MOM6, the ocean component for CESM  
989 version 3; the Department of Energy has implemented MARBL in MPAS-O, the ocean  
990 component for the Energy Exascale Earth System Model (E3SM) (Burrows et al., 2020);  
991 efforts are underway to implement MARBL in the Regional Oceanographic Model (ROMS)  
992 (Shchepetkin & McWilliams, 2005), enabling high-resolution regional configurations; and  
993 finally, MARBL has been coupled to the Ocean Circulation Inverse Model (OCIM) (DeVries  
994 & Primeau, 2011; DeVries, 2014) leveraging an interface layer suitable for Matlab and  
995 Python applications. Future goals include establishing a one-dimensional test-bed frame-  
996 work, enabling more comprehensive parameter exploration and an educational resource.  
997 Our goals explicitly include building a diverse, inclusive community of researchers involved  
998 in the development and application of MARBL across a range of use-cases. Managing  
999 such a development process imposes some challenges, but also has the potential to yield  
1000 significant benefits derived from synergies across diverse applications.

1001 As mentioned above, the ecosystem configuration in MARBL is flexible, and im-  
1002 provements in the simulated phytoplankton and zooplankton diversity is currently an  
1003 area of active development. Part of the motivation for this work is to improve the ca-  
1004 pacity for CESM to address critical questions related to the impacts of climate variabil-  
1005 ity and change on marine ecosystems. Another area of interest involves climate inter-  
1006 vention strategies, including the efficacy and potential impacts associated with ocean car-  
1007 bon dioxide removal (CDR) strategies.

1008 In summary, we aim to continue to develop MARBL as a community resource and  
1009 cutting-edge research tool. Work continues to improve MARBL, building toward a com-  
1010 prehensive treatment of ocean biogeochemical cycles, capable of robust assessments of  
1011 climate impacts and the effect of human manipulations.

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