Biogeochemical variability in the California Current System

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Abstract

The biological productivity and diversity of the California Current System (CCS) is at the leading edge of major emerging climate trends, including hypoxia and acidification. We present results from a hindcast simulation (reanalysis) of an eddy-resolving oceanic physicalbiogeochemical model of the CCS, to characterize its mean state and its patterns and drivers of variability in marine biogeochemical and ecosystem processes from 1995-2010. This is a companion paper to a physical analysis in Renault et al. (2019). The model reproduces longterm mean distributions of key ecosystem metrics, including surface nutrients and productivity and subsurface O_2 and carbonate undersaturation. The spatial patterns of Net Primary Productivity (NPP) are broadly consistent with measured and remotely sensed rates, and they reflect a predominant limitation by nitrogen, with seasonal and episodic limitation by Fe nearshore in the central CCS, and in the open ocean northern CCS. The vertical distribution of NPP is governed by the trade-off between nutrient and light limitation, a balance that reproduces and explains the observed spatial variations in the depth of the deep Chl maximum. The seasonal to interannual variability of biogeochemical properties and rates is also well captured by model simulations. Because of the prevailing nutrient limitation, fluctuations in the depth of the pycnocline and associated nutricline are the leading single factor explaining interannual variability in the interior biogeochemical state, and the relationships between density and biogeochemical rates and tracers are consistent between model and observations. The magnitude and relationship between density structure and biogeochemical processes is illustrated by the 1997-98 El Niño event, which faithfully reproduces the single largest deviation from the mean state in the simulated period. A slower decadal shoaling of the pycnocline also accounts for the concomitant trends in hypoxic and corrosive conditions on the shelf. The resulting variability is key to understanding the vulnerability of marine species to oceanic change, and to the detection of such changes, soon projected to exceed the range of conditions in the past century.

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1 Introduction

Coastal upwelling systems along the eastern boundaries of the subtropical oceanic basins are some of the most climatically and biologically dynamic regions of the world's oceans (Carr and Kearns, 2003; Kudela et al., 2008). In the California Current system (CCS) of the Eastern North Pacific, the seasonal cycle of alongshore winds and offshore surface currents yields a direct conduit for nutrient-rich water to rise from the deep ocean to the sunlit surface (Chavez and Messie, 2009). The upwelling of cold water near the coast is evidenced in satellite remote sensing of sea surface temperature, expansive offshore marine stratocumulus clouds, and a nearshore ribbon of high chlorophyll from phytoplankton. The resulting cascade of phytoplankton biomass up the food chain supports high biodiversity and productive fisheries (FAO, 2009).

The physical state of the Northeastern Pacific varies on time scales of days to decades. This variability includes mesoscale eddies, seasonal cycle, interannual El Niño Southern Oscillation (ENSO) variability, and lower frequency climate fluctuations characterized by the Pacific Decadal Oscillation (PDO, Mantua et al. (1997)) and the North Pacific Gyre Oscillation (NPGO, Di Lorenzo et al. (2008)). Ecosystems throughout the Eastern Pacific respond strongly to physical forcing at each of these timescales, through the physical influence of winds, light, and heat, and their effects on the supply of nutrients to phytoplankton and oxygen to marine animals. Interannual climate cycles associated with ENSO in particular are major perturbations to these parameters and thus to plankton productivity (Chavez et al., 2002; Bograd and Lynn, 2001). Because thermocline waters entering the CCS are, like other subtropical Eastern Boundary Upwelling Systems, far from the sites of atmospheric ventilation in the west, the rising waters are "old" and bear the signature of decades of biogeochemical process that yield low O_2 , high nutrients, and low pH. The upwelling in eastern boundary systems also generates energetic mesoscale eddy fields (Capet et al., 2008). These eddies can transport the extreme properties of CCS thermocline waters far offshore and down into the subtropical interior, connecting the biogeochemistry of the coastal CCS with the adjacent oceanic gyres (Nagai et al., 2015; Gruber et al., 2011; Renault et al., 2016a; Frenger et al., 2018).

Oceanic acidification and deoxygenation are also emerging trends in the CCS ecosystem (Gruber et al., 2012; Chan et al., 2008). Anthropogenic CO_2 has been detected in coastal subsurface waters off Northern California (Feely et al., 2008). Decadal trends in oxygen have also been observed in the California Current (McClatchie et al., 2010; Pierce et al., 2012) and have altered the proportions of biologically important nutrients (Deutsch et al., 2011). In the Northern California Current, massive benthic die-offs have been attributed to episodes of extreme hypoxia along the Oregon shelf (Chan et al., 2008). Fluctuating abundance of species in upper trophic levels observed over decadal and longer time scales arise from climate variability, but the specific mechanisms remain obscure (Chavez et al., 2003; Rykaczewski and Checkley, 2008). In addition to climate forcing, the coastal ocean is subjected to anthropogenic pollution that could locally exacerbate hypoxia and acidification (Doney et al., 2007), and may contribute to the increasing frequency and toxicity of harmful algal blooms along Californias coast (Andersson et al., 2008).

The continental margin of western North America has a rich endowment of historical and ongoing observational programs aiding evaluation of climate-ecosystem interactions, including one of the most extensive time series programs (the California Cooperative Ocean Fisheries Investigations, CalCOFI) anywhere in the world's oceans (Ducklow et al., 2009). The age of shallow thermocline waters also make them prone to large amplitudes of low-frequency variability (Ito and

Deutsch, 2010) that confound the detection of climate and other anthropogenic changes impinging through the oceanic surface. Thus, even in this relatively well-studied region of the ocean, it is difficult to distinguish long-term trends associated with anthropogenic climate change from the low-frequency variability that pervades oceanic properties in regions of close contact with old, deep waters. For example, it remains unclear whether the changes in hypoxia in the California Current are driven by internal climate variability, or are an early sign of long-term climate warming (McClatchie et al., 2010; Pierce et al., 2012; Long et al., 2016).

Models of this dynamic region often exhibit substantial biases in the mean state and unknown fidelity in representing historical variability and its causal mechanisms. The purpose of this paper is to report a system-wide validation of eddy-resolving, regional model fields and property relationships through comparison to a variety of hydrographic, experimental, and remote sensing observations. Here we focus on biogeochemical aspects of the model solution; the physical dynamics and its model validation are discussed in a companion paper (Renault et al., 2019). In this work, Sec. 2 provides a description of the model, its boundary conditions, the simulations performed, and the datasets used for model evaluation, which constitute the core results described in Sec. 3 and summarized in the final Sec. 4.

2 Methods

2.1 Model description

The ecosystem and biogeochemical cycles are simulated in the Regional Ocean Modeling System (ROMS, Shchepetkin and McWilliams (2005)). As in Renault et al. (2016b), the primary domain extends from 144.7°W to 112.5°W and from 22.7°N to 51.1°N. Its grid is 437 x 662 points with a horizontal resolution of dx = 4 km and 60 vertical levels. Initial and horizontal boundary data for temperature, salinity, surface elevation, and horizontal velocity are taken from the quarter-degree, daily-averaged Mercator Glorys2V3 product (http://www.myocean.eu). In order to maintain a realistic water mass representation, monthly anomalies from the Mercator data are added to the mean monthly climatology from the World Ocean Atlas (WOA, Locarnini et al. (2013); Zweng et al. (2013)) over the period 1995-2004. The freshwater, turbulent heat, and momentum fluxes are estimated using bulk formulae (Large, 2006) and the atmospheric fields derived from an uncoupled simulation with the Weather Research and Forecasting model (WRF). This statistically equilibrated solution, named USW4, is integrated over the period 1995-2010 after a spin up of 1 year (started from a previous ROMS solution with 12 km horizontal resolution). Further details are described in the companion paper (Renault et al., 2019).

The coastal biogeochemical dynamics are simulated using an ecosystem model (the Biogeochemical Elemental Cycling (BEC) model, Fig. 1 and Appendix). This model includes both phytoplankton and zooplankton, and dissolved, suspended, and sinking particulates (Moore et al., 2004). The model includes four phytoplankton functional groups (picoplankton, diatoms, coccolithophores, and diazotrophs) characterized by distinct biogeochemical functions (nutrient recycling, silicification, calcification, and N₂ fixation, respectively). Four nutrient cycles (nitrogen, silicic acid, phosphate, and iron) are simulated and are coupled through a fixed phytoplankton stoichiometry. The ecosystem is linked to an oceanic biogeochemistry module that includes total inorganic carbon (DIC), alkalinity, iron, and dissolved O_2 . Remineralization of sinking organic matter is parameterized according to the mineral ballast model of Armstrong et al. (2001). Gas exchange fluxes for O_2 and CO_2 are based on Wanninkhof (1992). The BEC equations are listed in the Appendix, and the model code, including parameter settings, are available through the GitHub repository (see the remark at the end of the paper).

The iron Fe cycle includes dissolved iron, scavenged iron, and iron associated with organic matter pools and dust particles, but only dissolved iron and organically bound iron are explicitly modeled as state variables. For dissolved iron, four processes are considered: atmospheric deposition, biological uptake and remineralization, scavenging by sinking particles, and release by sediments. Atmospheric iron deposition is based on the dust climatology of Mahowald et al. (2006). We implemented a sedimentary iron source based on benthic flux chamber measurements in the California margin. An equation relating sediment Fe release as a function of bottom water O_2 (log₁₀[F_{fe}] = 2.5 - 0.0165 \cdot O_2 , where O_2 is in mmol m⁻³ and the efflux units are μ mol m⁻² d⁻¹) is derived from data compiled by Severmann et al. (2010). The resulting rates of Fe supply from sediments (Fig. 2) exceed those from atmospheric dust deposition throughout the model domain.

We also added a nitrogen cycle, with losses to the sediments and water column. Bottom water nitrate is removed using a statistical description of sediment denitrification proposed by Middelburg et al. (1996), based on a vertically resolved diagenetic model that predicts the primary dependence of benthic denitrification to be on organic carbon sedimentation rate, with a secondary sensitivity to bottom water oxygen concentration. This statistical description of the complete diagenetic model reproduces basic controls on observed sediment fluxes, without the considerable computational cost of a sedimentary submodel. The predicted rates of N loss from this sedimentary sink (Fig. 2) amount to a small loss of $\approx 3 \times 10^{12}$ gN yr⁻¹. Denitrification in the water column is also modeled, but its integrated removal rate is an order of magnitude smaller than sedimentary losses, and has negligible impact on the results because O_2 in the model domain rarely falls below the threshold (5 mmol m⁻³) assumed for this process. The higher O_2 thresholds associated with anaerobic particle micro-environments could increase the importance of anaerobic processes in the CCS, but they are not represented in this model (Bianchi et al., 2018).

2.2 Biogeochemical forcing and validation data

The physical conditions at the open boundaries are extracted from a global reanalysis as described in the companion paper (Renault et al., 2019) (Fig. 1). The biogeochemical boundary conditions for nutrients $(NO_3^-, PO_4^{3-}, Si(OH)_4)$ and O_2 are taken from monthly climatological observations in the 2013 World Ocean Atlas (WOA) (Garcia-Reyes et al., 2014). Boundary condition data for Fe is taken from global simulations with the Community Earth System Model (CESM) that used an earlier version of the same BEC ecosystem model. The NH_4^+ boundary concentrations, being small in nature, are set to zero, but adjust rapidly to the ecosystem processes in the interior of the domain. Time-dependent carbon cycle parameters, DIC and Alk, are taken from GLO-DAP (Key et al., 2004), with a reference year of 1995. An imposed trend at the boundary scales the anthropogenic component of DIC in proportion to the rise of atmospheric CO_2 since 1995. Time-dependent atmospheric pCO_2 is also used for as surface boundary condition for air-sea gas exchange. Aside from the boundary DIC, the only non-stationary forcing of the model solution comes from the physical boundary conditions and surface forcing (Renault et al., 2019).

In order to ensure the integrity of tracer relationships along isopycnal surfaces, we map the



Figure 1: Schematic structure and physical configuration of ROMS-BEC biogeochemical model. (a) The main ecosystem state variables and fluxes. (b) Geographic scale of simulation, and sources of surface forcing, open boundary condition data and representations of benthic nutrient fluxes. The oceanic lateral boundary conditions are applied in a dx = 12 km outer domain, within which the dx = 4 km subdomain is nested.

biogeochemical boundary conditions from source data to the model grid using density rather than depth as the vertical coordinate, while retaining the mean-seasonal values of T and S (hence density) along the boundary as specified in the physical conditions (Renault et al., 2019). This prevents any errors in the depth of isopycnal surfaces inherited from the physical boundary data (Mercator) from biasing the biogeochemical properties along that surface.

The CCS is among the best-sampled regions of the world's oceans. Hydrographic sampling and biological rate measurements have been conducted repeatedly if not routinely along several sections off the West coast, most notably in the CalCOFI lines in the Southern California Bight, the Newport Line off Oregon, and Line P off British Columbia, the latter being at the northern edge of the 4 km model domain. Despite the abundant datasets from this region, data density is still sparse for much of the central California coast and for many biogeochemical properties of interest (*e.g.*, *Fe*). The total number of profiles in the World Ocean Database are plotted for $NO_3^$ and O_2 over the entire historical data period (1955-2013; Fig. 3). In addition to these climatological databases, we use specific repeat hydrographic lines in the Southern California Bight, off Monterey Bay, California and Newport, Oregon to evaluate the vertical and cross-shelf structure of biogeochemical variables at greater resolution.

To facilitate comparison of model outputs to data, and particularly seasonal cycles, we defined 6 regions (Fig. 3), dividing the CCS by distance from the coast into nearshore (0-100 km) and offshore (100-500 km) regions, and by latitude into the Northern, Central and Southern CCS. These designations are somewhat arbitrary, but are based on a combination of topographic delineations and to ensure adequate data coverage in each region. Locations of the primary repeat hydrographic lines and regional boundaries used for model analysis are also shown in Fig. 3. High frequency biogeochemical measurements from moorings are generally available only for more recent periods, and primarily from nearshore environments. We therefore focus our validation efforts on broad-scale measures that can be evaluated from climatological databases, namely the



Figure 2: Parameterized fluxes of iron and nitrate between the water column and sediments. (a) The Fe efflux (in mmol m⁻² yr⁻¹) to the water column from the sediments. (b) The nitrate flux from the water column to the sediments due to net denitrification in sedimentary pore waters. Both fluxes are parameterized as a function of bottom water O_2 (for Fe and N) and organic matter flux to the seafloor (N only). The maps are therefore part of the model solutions, and not prescribed forcings.

2013 World Ocean Database (downloaded from https://www.nodc.noaa.gov/OC5/WOD13/), and its objectively-mapped climatological representation, the World Ocean Atlas (WOA; Garcia-Reyes et al. (2014)). Comparison to mooring data is left for planned downscaling of these simulations better suited to examine high-frequency variability.

3 Results

We describe the spatial patterns and temporal variability of model biogeochemical solutions and their fidelity to observational datasets. Of the numerous properties and rates of the biogeochemical system that are predicted by the model, we focus on those that are most important to the overall elemental cycling of carbon, nitrogen, and oxygen, and are best observed over scales captured by the model. Our analysis begins with surface rates of air-sea CO_2 flux and net primary productivity (NPP), followed by the factors limiting NPP, including nutrients and light. We then evaluate the rates and efficiency of organic matter exported to depth and the accumulated signature of its respiration in deeper waters that lead to hypoxic and corrosive conditions in the thermocline. Aspects of the model solution that are not presented include nutrients that are not limiting, rates that lack large-scale and climatological datasets, and variability that is poorly resolved by a dx = 4 km model (*e.g.*, submesoscale and nearshore phenomena).



Figure 3: Hydrographic data density used for model validation. Observations in the World Ocean Database are binned in a regular 1° latitude/longitude grid for each month over the entire historical data period (1955-2013). Total number of months with a profile are shown for NO_3^- (left) and O_2 (right). Nominal station locations for major repeat hydrographic lines used for validation (see Fig. 17) are shown (red circles), along with the boundaries (black lines) used for regional time series comparisons in Figs. 6, 7, 15, and 21.

3.1 Photic Zone

3.1.1 Chlorophyll and Net Primary Production

We begin with an evaluation of model distributions of Chlorophyll-a (hereafter Chl), and Net Primary Productivity (NPP), both of which are estimated from remote sensing of ocean color. While NPP is of greater biological significance, its estimation is more indirect than for Chl.

The model Chl concentrations are governed by the product of biomass and the C : Chl ratios. Biomass is subject to advection and to ecosystem transformations (see Appendix). The C : Chl ratio is determined by photoacclimation, or the amounts of light-harvesting pigments and photoprotective compounds produced by phytoplankton in response to their growth environment. This process is included in the ROMS-BEC representation of phytoplankton physiology following the model of Geider et al. (1998), which relates changes in chlorophyll synthesis and nutrient uptake in response to changing PAR. The dominant patterns of Chl are also found in biomass (see below), indicating that photoacclimation is not the leading factor in Chl variability. While biomass may be a more ecologically meaningful comparison, we validated model solutions using Chl because it is directly estimated from ocean color. The frequency distribution of Chl in both ROMS and

SeaWIFS is approximately log-normal and is mapped after logarithmic transformation.

The annual mean concentrations of Chl vary most strongly in the cross-shore direction, with relatively weak alongshore gradients, a well-known pattern in observations (Banas and Hickey, 2008) that is well represented by the model (Fig. 4a,b). The offshore drop-off in Chl is somewhat weaker in model simulations, resulting in a wider band of high coastal Chl, a tendency that is not reflected in NPP (discussed immediately below). Kessouri et al. (2019) shows that there is some sensitivity in these distributions to model resolution, with higher resolution increasing the nearshore biomass and productivity. The leading pattern of variability in climatological Chl is characterized by a seasonal cycle that is also largely synchronous along the coast. The leading Empirical Orthogonal Function (EOF) (Fig. 4d,e), has Chl reaching peak values in late summer, and it accounts for the large majority of the climatological variance in both observations (EOF1=64% variance) and model solutions (EOF1 = 69% variance). The second EOF (not shown) also has a similar loading pattern, but with more meridional structure offshore and a minimum in spring but accounts for only 13% and 19% of the variance, respectively. The seasonality of Chl reveals antiphased cycles between nearshore high-Chl band and the lower Chl offshore. While near-shore Chl peaks in late summer, the offshore surface Chl has a minimum. This pattern is not found in the EOFs of depth integrated NPP, indicating that the dipole structure of the Chl pattern results from a vertical redistribution of Chl to greater depths in offshore waters as they become more oligotrophic due to nutrient uptake during summer months. This interpretation is confirmed in the analysis of the vertical *Chl* maximum (Sec. 3.1.2).

In BEC the net primary productivity (NPP) depends on the sum of j model phytoplankton biomasses (B_j) , their maximum growth rates $(\mu_{max}(T) = \mu_0 T^{1.06})$, and the limitation of those rates by light $(0 \le \gamma_j(I) \le 1)$ and the minimum Michaelis-Menten function $\lambda_{i,j}$ $(0 \le \lambda_{i,j}(N_i) \le 1)$, among the *i* nutrients with half-saturation $K_{i,j}$ $(0 \le \lambda_{i,j}(N_i) \le 1)$, written as

$$NPP = \sum_{i,j} \mu_{max\,j}(T) \gamma_j(I) \min[\lambda_{i,j}(N_i)] B_j.$$
(1)

The spatial patterns of modeled NPP fall within the range of satellite-derived estimates (Fig. 5). The two commonly used satellite-based algorithms are the vertically generalized production model (VGPM) (Behrenfeld and Falkowski, 1997) and the carbon-based productivity model (CbPM) (Westberry et al., 2008). The VGPM estimates productivity on the basis of light and chlorophyll concentrations, calibrated to a predominantly coastal radiocarbon incubation dataset. The CbPM additionally incorporates phytoplankton backscattering and growth rate relationships in order to estimate productivity as a function of phytoplankton biomass, and it is calibrated to subtropical gyre radiocarbon incubations. The two algorithms exhibit a relatively wide range for the CCS region, reflecting the considerable uncertainty in "empirical" NPP estimates as well as differences in the measurements underlying each. The VGPM algorithm has a larger offshore gradient, with higher coastal values, and lower values in the open ocean, compared to the carbon-based CbPM. NPP rates from ROMS-BEC generally fall between the two remote sensing products, but are generally closer to the values of the VGPM algorithm, supporting higher near-shore rates, lower offshore rates, and increased seasonality relative to the CbPM. The VGPM has been explicitly calibrated against radiocarbon bottle incubations from the CalCOFI program, and it is therefore likely to be more accurate in this region (Kahru et al., 2009). Indeed, we find that ROMS-BEC rates and distributions of productivity are also consistent with direct estimates from the ship-based data both from CalCOFI (Fig. 5; Munro et al. (2013)) and the broader subtropical Northeast Pacific (Palevsky et al., 2016).

Productivity in the northern CCS has been consistently biased in regional models (Banas and Hickey, 2008), including in our initial simulations. We conducted simulations with and without lateral nitrogen fluxes at the Strait of Juan de Fuca imposed as boundary condition from model simulations by Davis and Martín (2014). Consistent with that study, without the nutrient inputs from the Salish Sea, NPP was biased low by > 50%. The inclusion of the effects of N inputs at the Juan de Fuca Strait brought the model much closer to satellite-based empirical models. The inclusion of these inputs is consistent with the study by Davis and Martín (2014), and it is used in all results reported here. Non-nitrogenous nutrients, including Fe, were not available for inclusion in fluxes from the Strait of Juan de Fuca.

The seasonal cycle of NPP is also well captured by the model (Fig. 6). In all 6 regions of the CCS, the climatological NPP, integrated over the depth of the photic zone and averaged over the regional mask, exhibits an amplitude and phasing that is within the range of satellite-based empirical models. The most notable exception is the offshore Northern domain, where a spring bloom is predicted to be of stronger magnitude than estimated by either satellite product. This model result is consistent with measured geochemical tracers, which also indicate the spring bloom in the offshore Northeast Pacific is greater than estimated from the satellite algorithms (Palevsky et al., 2016). A smaller discrepancy occurs in the southern nearshore region, where ROMS-BEC generates greater summer production than either of the satellite algorithms. Model NPP in the oligotrophic part of the domain is lower than satellite estimates, but is more consistent with the most offshore values in the depth integrated rates based on radiocarbon bottle incubations from CalCOFI. Overall, where and when ROMS-BEC and satellite algorithms for NPP disagree, ROMS-BEC output is generally closer to the available observational data.

3.1.2 Seasonal limitation of productivity by light and nutrients

To evaluate the role of environmental factors shaping the seasonal cycle and regional differences in rates of productivity, we computed monthly mean limitation factors for each of the environmental variables that modulate the maximum growth rates, including macronutrients (NO_3^-, PO_4^{3-}) , $Si(OH)_4$, Fe, and light; see (1). By construction, growth rates are limited by only one nutrient at a time (Liebigs Law of the Minimum), such that only the lowest value has an influence on rates. Light operates as a multiplicative factor on nutrient limitation, while temperature influences maximum growth rate, but is not considered a limitation factor in the upper ocean, so is not analyzed here. Over a climatological seasonal cycle, small plankton growth rates are almost always reduced by light more so than by nutrients, regardless of season or location (Fig. 7). The small plankton are assumed to have a lower half-saturation constant for nutrients, and the resulting higher affinity makes them less prone to nutrient limitation than large plankton are. Thus, in the inshore regions where nutrients are high, light is always limiting. Offshore, nutrients can limit small phytoplankton to a similar degree as light in the summer. However, large phytoplankton make up $\approx 90\%$ of modeled net primary productivity on average, and large phytoplankton ("diatoms") are primarily limited by nutrient availability. In the model, this results from a nutrient supply by combined windinduced upwelling and eddy-induced subduction (Gruber et al., 2011; Nagai et al., 2015; Renault et al., 2016a) that are unable to saturate the potential net uptake of nitrogen by phytoplankton at prevailing light levels.² Large phytoplankton only experience light limitation in the northern CCS,

² Significant eddy fluxes also occur due to submesoscale eddies and fronts in the CCS (Kessouri et al., 2019).

where the seasonal cycle alternates between winter light limitation and nutrient limitation for the rest of the year.

While on the regional scales used for this analysis N limitation appears more stringent than Fe limitation throughout the CCS, significant Fe limitation occurs on smaller scales and shorter durations (see below). The seasonal amplitude of plankton growth rates is relatively small ($\approx 10\%$), indicating that the amplitude of seasonal production ($\approx 100\%$) is governed by seasonal controls on biomass rather than growth rates. The trade-off between light and nutrient limitation spatially and seasonally is a ubiquitous feature of phytoplankton distributions and phenology. Moving deeper in the water column, light becomes more limiting as photosynthetically active radiation (PAR, light of 400 - 700 nm wavelengths) is absorbed and scattered, while nutrient concentrations are greatest below the surface mixed layer and rapidly decrease upward across the seasonally variable pycnocline; *i.e.*, there is a well-defined nutricline.

The competing influences of nutrient and light limitation on depth of optimal plankton growth are reflected in the depth of the deep chlorophyll maximum (DCM). In the CCS, the depth of the DCM deepens from the coast to the open ocean, suggesting that the optimal combination of light and nutrients is found deeper offshore, consistent with a deepening nutricline that intensifies nutrient limitation at the surface and light limitation where nutrients are abundant for both small and large phytoplankton. We use the observed pattern of the DCM depth as an indicator of whether the model achieves a realistic trade-off between these two countervailing growth condition gradients (Fig. 8). However, for DCM depth to be a reliable indicator of phytoplankton growth rate trade-offs, we must rule out two simpler interpretations relating to isopycnal advection and photoacclimation. First, the offshore deepening of the DCM can have a physical origin. Because it closely follows the plunging of isopycnal surfaces offshore, the vertical peak in biomass and associated chlorophyll could be caused by eddy subduction carrying high surface chlorophyll away from the coast along deepening isopycnals (Gruber et al., 2011; Nagai et al., 2015; Renault et al., 2016a). To evaluate this possibility, we compared the depth of plankton biomass to the depth at which the product of light and nutrient limitation terms are maximized (Fig. 8). The two maps are virtually indistinguishable, suggesting the DCM follows growth rates rather than advection. As a more stringent test, we performed a short (5-year) simulation in which surface PAR was reduced by 10%. The results revealed a significant shoaling of both the biomass and Chl peaks, but no detectable change in isopycnal depths, confirming that these depths do in fact reflect a nutrientlight trade-off rather than advection along density surfaces. Second, the peak depth of Chl may also be decoupled from that of biomass and growth rates due to photoacclimation, or a shift in the amounts of light-harvesting pigments and photoprotective compounds produced by phytoplankton in response to the light-environment. This process is included in the ROMS-BEC representation of phytoplankton physiology following the model of Geider et al. (1998), which relates changes in chlorophyll synthesis and nutrient uptake in response to changing PAR. Indeed, we find that the depth of the DCM is slightly deeper than that of the maximum plankton biomass. However, the offshore and latitude gradients of the depth of peak biomass and chlorophyll are very similar. DCM deepening offshore is consistent with optimized growth conditions in the model, and reproduce the pattern observed in the available CalCOFI data (Fig. 8).

Thus, we take this correspondence to indicate that the model adequately captures the essential trade-off between light and nutrient limitation. Growth rates are modulated by a complex and evolving pattern of nutrient limitation by reactive nitrogen $(NO_3^- + NH_4^+)$ and soluble dissolved Fe, with no appreciable limitation by $Si(OH)_4$ and PO_4^{3-} in the CCS. The limitation factors are

mapped as a biomass-weighted fraction of time that each of the nutrients is most limiting (Fig. 9). The spatial pattern among nutrients largely reflects the areas where NO_3^- supply routinely exceeds maximum potential uptake seasonally. Thus, the waters entering the CCS from the subarctic High Nutrient – Low Chlorophyl (HNLC) region are most frequently Fe limited. Along the coast, the seasonal upwelling of excess nitrogen leads to significant periods of Fe limitation as well. In the coastal zone off Monterey Bay, Fe limitation has been diagnosed via incubation experiments, in a band of water slightly offshore, with nitrogen limitation both in more shoreward and open coastal zones (Firme et al., 2003). This pattern is consistent with that predicted by the model (Fig. 9b inset). Most of the rest of the model domain is perpetually nitrogen limited.

3.1.3 Nutrient concentrations

Surface nutrient concentrations provide an important measurable test of system behavior. For nutrients that limit phytoplankton growth, accurately simulating their distributions is a necessary condition for a mechanistic prediction of NPP. Moreover, they provide an integrative measure of net community production (equal to NPP less community heterotrophic respiration) and export of organic matter to the thermocline. We therefore compared model predicted distributions of the two primary limiting nutrients, NO_3^- and Fe, to available observations. Coastal measurements of dissolved Fe reveal a spatially patchy distribution, reflecting its short residence time with respect to removal by plankton uptake and particle-active scavenging. Existing data are too sparse to yield a clear climatological pattern for model validation. However, the primary coastal region where the model predicts most frequent Fe limitation, in the central CCS, has been relatively well sampled, including on two cruises off Monterey Bay that also tested for Fe limitation (Firme et al., 2003); see Fig. 9, inset). Given the lack of a clear large-scale pattern of surface Fe levels, we used a more statistically-based validation metric, focusing on the relative frequency of Fe measurements versus concentration and distance from the shore (Fig. 10). On average, the data and model both show a decline in the mean and median Fe levels with offshore distance. This reduction is driven largely by the decreasing frequency of high concentrations, while the most commonly observed Felevels remain consistent at $\approx 0.5 \ 10^{-6} \ \text{mol m}^{-3}$ regardless of distance from shore. The thinning tail of high concentrations in Fe distribution with cross-shelf distance occurs in both modeled and observed fields, but is more pronounced in the measurements.

In the CCS, the nutrient most often limiting NPP is reactive nitrogen, of which by far the largest and most commonly measured pool is nitrate ([NO_3^-], hereafter simply N). We therefore compare the model simulated patterns of NO_3^- to climatological values from the World Ocean Database (Fig. 11). A depth of 50 m is chosen because it approximates the average depth of maximum biomass and NPP (see Fig. 8). We included all historical measurements for this analysis because the data density in the model period (1994-2010) was much sparser, and no significant differences were found between the average N in this period relative to 1955-2013. ROMS-BEC captures regional patterns well for NO_3^- (Fig. 11). Annual mean concentrations of ≈ 15 mmol m⁻³ along the coast decline to values below the half-saturation level for model diatom growth (N < 1 mmol m⁻³) within 500 km from shore. The offshore gradient is similar throughout most of the CCS, except the Southern California Bight (SCB) where coastal surface values are much lower. Similar model fidelity was found for other macronutrients ($Si(OH)_4$, PO_4^{3-}), but not shown because they do not reach limiting concentrations. The coastal zone exhibits strong variability in N at the base of the photic zone (50 m), with standard deviations of 5-10 mmol m⁻³ throughout most of the

coastal zone, but with a slight northward increase in variance (Fig. 11c). The variability of N in the climatology (WOA) exhibits a similar spatial pattern, but with $\approx 50\%$ of the magnitude. Thus, approximately half of the variation in surface nutrients most commonly limiting NPP is associated not with the seasonal cycle, but with interannual variability. The model also reproduces observed magnitudes and patterns of NO_3^- variability (Fig. 11d). We use the interannual anomaly fields in the model and in observations to test the importance of nutrient supply as a mechanism driving changes in NPP over time.

3.1.4 Interannual variability in NPP

Factors that limit NPP during the mean seasonal cycle may drive interannual and longer term productivity changes. We examined correlations between NPP and both light and nutrient concentrations in model simulations and observations, where available. Interannual anomalies in NPP in ROMS-BEC are found to be well correlated ($R^2 \approx 0.5$) to the density of water below the photic zone (100 m) (Fig. 12). This reflects the role of pycnocline heave and vertical mixing in supplying macronutrients, and alleviating local nutrient limitation. Observations show a similar magnitude of correlation in the southern and central CCS, but a weaker correlation to the north. This may reflect the role of nutrient supply processes that are either missing or represented only climatologically in our model, and not connected to pycnocline heave. Because the weaker correlations are in the northern domain where nutrients can enter from subarctic surface waters, the climatological NO_3^- used for the boundary conditions is a likely culprit. However, the presence of river N sources or a variable Juan de Fuca flux could also weaken the correlation in the observations relative to the model.

Predicted correlations between NPP and density can be tested directly from hydrographic observations in the southern CCS. Relationships between nutrient and density anomalies (subtracting the mean seasonal cycle), are of similar strength, accounting for $\approx 50\%$ of the variability in both CalCOFI observations and model simulations (Fig. 13). Interannual anomalies in NPP in ROMS-BEC are also significantly correlated with surface PAR due to variable cloudiness, though it accounts for a smaller fraction of the variance ($\approx 20\%$). The role of light is confined closer to the coastal upwelling where surface NO_3^- is high, and light availability thus limits phytoplankton growth.

3.1.5 Remineralization and export of carbon from the photic zone

Of the net production of organic matter by phytoplankton, a substantial fraction can be respired by zooplankton and higher trophic levels. The remainder is available to be transported away, by particle sinking and transport of dissolved organic matter, *e.g.*, via eddy subduction (Sec. 3.1.2). The fraction of NPP that is regenerated within the surface ocean depends on food web processes, such as grazing rates. Although data is not available to evaluate large-scale patterns of grazing rates, an indirect comparison can be made through the export flux and the fraction of NPP that is exported in sinking particles rather than recycled (the so-called pe-ratio (Dunne et al., 2005; Murray et al., 1196)).

The fraction of NPP that is exported varies from 5-25%, consistent with the range of values inferred in field studies (Fig. 15). The model predicts highest pe-ratios in the coastal zone, where productivity is high and sea surface temperature relatively low. These dependencies are also

consistent with those inferred from a global analysis of rate measurements for NPP and Net Community Production (NCP, assumed equal to total export of both particulate and dissolved organic carbon) (Dunne et al., 2005). For a more quantitative comparison, we compared modeled pe-ratios to those predicted by a statistical model fit to global observations by Dunne et al. (2005), over the seasonal cycle in each of our 6 standard CCS regions (Fig. 15). In both the empirical model and in ROMS-BEC, the mean value, phasing, and seasonal amplitude of changes in pe-ratio are similar. Empirically based estimates generate a similar result even when sea surface temperature is held constant, suggesting that the trophic mismatch between phytoplankton growth and zooplankton grazing is the primary reason for variable pe-ratios, a prediction that is also consistent with model behavior.

Annual mean export flux represents the transfer of biogenic material from the surface to depth, and thus the influence of CCS productivity on air-sea CO_2 flux and thermocline properties. Model simulated export production ranges from $\approx 1-10 \mod C \mod r^{-2} y^{-1}$ (Fig. 15b). Few measurements of export or net community production are available to evaluate the overall pattern of this flux. In the SCB region, measurements of O_2 have been used to estimate NCP rates of 3 - 17 (mean 6.5) mol C $m^{-2} y^{-1}$ (Munro et al., 2013), in line with modeled rates and with similar spatial patterns of export (greatest along the Northern coast of the SCB). Munro et al. (2013) also combine information from ¹⁵N uptake experiments and nitrate based new production ratios (Dugdale et al., 1992; Eppley et al., 1992) with radiocarbon incubations to generate very similar estimates of the magnitude and spatial variability of production in the SCB. Particle export based estimates of export (neglecting the role of dissolved carbon) are lower than observed NCP as well as ROMS-BEC export, 2 - 4 mol C $m^{-2} y^{-1}$ (Collins et al., 2011; Stukel et al., 2011), though enhanced particle export associated with mesoscale fronts (2 - 3 times greater rates over smaller spatial and temporal scales (Stukel et al., 2017)) highlights the potentially important role of subduction by eddies and fronts in explaining mismatches between observations. Importantly, ROMS-BEC generates such mesoscale features which contribute to model export estimates. Model export production is also similar to regional nutrient budget analyses which suggest NCP averaging 7 - 9 mol $C \text{ m}^{-2} \text{ y}^{-1}$ over the broader CalCOFI region (Roemmich and McCallister, 1989; Bograd and Lynn, 2001). Similar nutrient budget analysis indicates that annual NCP should be $\approx 17 \text{ mol } C \text{ m}^{-2} \text{ y}^{-1}$ off Monterrey Bay, again in line with ROMS-BEC estimates of export for that region (Fig. 15b). As noted above, the fidelity of modeled nitrate distributions across the model domain provides a critical broad-scale measure of net biological drawdown, and thus of net community and export production.

Surface ocean CO_2 partial pressure and sea-to-air flux is reduced by net community production, increased by surface heat flux, reduced by freshwater fluxes, and modulated by upwelling and lateral circulation. It thus represents another important metric of overall system function. The role of the CCS and its sub-regions in the atmosphere-ocean balance of CO_2 has previously been investigated in several studies, both empirically (*e.g.*, Hales et al. (2012)) and in models (*e.g.*, Fiechter et al. (2014); Turi et al. (2014) and references therein). We evaluated the patterns of annual mean and summertime surface pCO_2 in the model hindcast simulation against observations in the SOCATv6 database from 1995-2010 (Bakker and coauthors, 2016), and the associated airsea fluxes over the modeled coastal region. Similar to observations over the simulated period, strong CO_2 supersaturations are simulated in a narrow band of coastal water within about 100 km of the shore. The values are highest along the central coast (35N-43N), lowest in the northern CCS, and intermediate in the southern domain. The highest values are associated with major topographic features, as previously noted by (Fiechter et al., 2014). The lack of apparent regional or seasonal

bias in the the surface pCO_2 , together with the good model-data agreement in surface buoyancy fluxes (Renault et al., 2019) and biological rates in these simulations (above), suggests that the balance of processes regulating the model's surface CO_2 fluxes is reliably captured. Consistent with previous studies, the net integrated CO_2 flux to the atmosphere (an uptake of 1.4 TgC/yr;) is found to be a relatively small residual of larger compensating outgassing and ingassing fluxes. The fact that our model predicts a larger net uptake is likely a result of the fact that its domain extends farther to the northern CCS where the combination of cold water and fresh water forcing suppresses surface pCO_2 . A detailed accounting of factors driving pCO_2 and air-sea flux variability has been described by (Fiechter et al., 2014; Turi et al., 2014). An extension of that analysis to the northern CCS domain with stronger freshwater forcing is left to future work.



Figure 4: Mean annual Chlorophyll (Chl) and its seasonal cycle. Mean annual concentrations (upper panels) are shown in mg m⁻³ averaged over the simulation period in both model output and SeaWIFS remote sensing level 3 product. The seasonal cycle is shown as the spatial loading patterns (middle panels) and time series (bottom panel) of the first EOF of climatological values of $\log 10[Chl]$. In both the mean annual and seasonal variations of Chl, the dominant variations are cross-shore and at Point Conception, which separates the central CCS from the SCB. High coastal Chl extends further offshore in model solutions than in observations, a bias that is not found in productivity (Fig. 5).



Figure 5: Spatial distribution of annual Net Primary Productivity (mol $C \text{ m}^{-2} \text{ yr}^{-1}$) integrated over the depth of the photic zone, from (top left) ROMS, (top right) CalCOFI, (bottom left) VGPM, and (bottom right) CbPM.



Figure 6: Seasonal cycle of annual Net Primary Productivity. The NPP rate (mol $C \text{ m}^{-2} \text{ yr}^{-1}$) from ROMS-BEC (black), and two satellite algorithms (VGPM, blue; CbPM, red) are integrated over the depth of the photic zone, and averaged over 6 regions (see Fig. 3) from northern (top row), central (middle row), and southern (bottom row) of the CCS, and divided by distance from the coast into a nearshore (0-100 km; right column) and offshore (100-500 km; left column).



Figure 7: Seasonal cycle of growth limitation factors for light (γ term in (1), blue) and for nutrients (λ term in (1): NO_3^- (green) and Fe (red) for diatoms (dashed) and small phytoplankton (dotted)). Factors are NPP-weighted and averaged over the photic zone for each region shown in Fig. 3. Limitation factors close to 1 mean no limitation; values close to 0 mean complete limitation.



Figure 8: Depth of the vertical maximum of Chlorophyll in CalCOFI (panel a) and ROMS (panel b), and of model diatom biomass (panel c) and nutrient limitation factor (panel d). The correspondence between model fields demonstrates that the model DCM tracks the nutrient limitation, while the fidelity to observations implies the model captures a realistic trade-off between nutrients and light limitation.



Figure 9: Frequency of limitation by N (left) and Fe (right) for the models dominant primary producer, diatoms. The limitation factors are weighted by biomass (as in Fig. 7), using 5-day output, and plotted as the fraction of time that each nutrient limitation factor is the lowest among nutrients. The inset shows offshore band of relatively frequent Fe limitation along the central CCS, as observed by (Firme et al., 2003).



Figure 10: Histogram of surface Fe concentrations (10^{-6} mol m⁻³) from observations (left) and ROMS (right). Because Fe concentrations are patchy in nature and sparsely observed, values are binned by distance from shore (0-20 km, upper row; 20-40 km middle row; 40-60 km lower row) to reveal a cross-shelf gradient, in the same latitude band off Monterey Bay for which observations were made on summer cruises (Firme et al., 2003).





Figure 11: Long-term mean and historical variability of nitrate (mmol m^{-3}) near the base of the photic zone (50 m) in a) ROMS, and b) WOD. Variability is mapped as the standard deviation and consists of roughly equal contributions from seasonal and interannual variability (see text). The full period of the World Ocean Database (1955-2013) is used to yield the most robust estimate of variance.



Figure 12: Correlation (R^2) between NPP and density, from ROMS-BEC (left) and CalCOFI measurements. The NPP rate is integrated over depth from ¹⁴C bottle incubation data, and density is interpolated to 100 m as an index of nutrient supply (see Fig. 12). In both variables, the mean seasonal cycle is removed leaving interannual variations. Relationships have similar strength in data and model, and indicate that $\approx 50\%$ of interannual variability in NPP can be attributed to anomalous nutrient supply due to pycnocline heaving.



Figure 13: Correlation (R^2) between NO_3^- and density at 100 m depth in ROMS-BEC (left) and WOD (right). For both variables, the mean seasonal cycle is removed leaving interannual variations. Interannual variations in subsurface (100 m) NO_3^- highly correlated with density ($R^2 \ge 0.8$) in most of CCS. Correlation is weaker in northern CCS in data than in model.





Figure 14: Surface ocean CO_2 partial pressure (ppm) and air-sea flux (mol m⁻² yr⁻¹). (a) Annual surface pCO_2 from ROMS. (b) Surface pCO_2 from SOCATv6 gridded coastal dataset, averaged over all months from 1995-2010. (c) Annual air gea CO_2 flux from ROMS. Negative values indicate outgassing to the atmosphere.



Figure 15: Annual mean and seasonal cycle of fraction of NPP that is exported in sinking particles (*i.e.*, pe-ratio; colored lines), and the export flux (mol $C \text{ m}^{-2} \text{ yr}^{-1}$; black line). Values diagnosed from model simulations (red line) are compared to an empirically derived algorithm (Dunne et al., 2005) based on *Chl* and SST (blue line), and from the same algorithm applied with constant SST (green line). Observed annual net community production, which should approximate export on an annual basis are measured to 5-10 mol $C \text{ m}^{-2} \text{ yr}^{-1}$ (Munro et al., 2013).

3.2 Thermocline

Here we test the model's representation of biogeochemical properties below the mixed layer and photic zone, *i.e.*, in the thermocline. We focus on distributions and variability of O_2 and aragonite saturation state (Ω_A) as these properties influence habitability for calcification and aerobic respiration by marine animals. The values of Ω_A are calculated from model dissolved inorganic carbon (DIC) and total alkalinity (Alk) using carbonate system equilibrium equations (CO_2SYS) (van Heuven et al., 2011). The variability of O_2 has been analyzed in greater detail using previous simulations of this model with different atmospheric and physical boundary conditions (Durski et al., 2017).

To evaluate the vertical structure of biogeochemical tracers, we turn to repeat hydrographic sections (Fig. 17). Transects through three cross-sections spanning the Southern California Bight (CalCOFI line 80), the central California coast (MBARI line 67), and the central Oregon coast (Newport line) show the typical vertical and cross-shelf gradients of NO_3^- and O_2 . The downward enhancement of NO_3^- and depletion of O_2 is a signal of the broad-scale vertical redistribution of these elements by the formation and degradation of organic matter within the CCS, as well as the gradients imported from the Pacific basin through the boundary conditions. The shoaling of the isopleths of both quantities follows that of the isopycnal surfaces by upwelling along the coast. The distributions of NO_3^- and O_2 are generally well reproduced by ROMS-BEC. The model somewhat underestimates the slope of these isopleths very nearshore. This tendency is also reflected in, and likely derived from, the same underestimate in the zonal tilt of isopycnal surfaces (Renault et al., 2019).

Along isopycnal surfaces, O_2 generally increases with latitude and with distance from shore, reflecting the contrasting properties carried by the broad offshore California current from the O_2 -rich subarctic, and the narrow near-shore California Undercurrent that transports low- O_2 waters of tropical origin northward along the slope. Both northern and southern end-member water types can be seen on the isopycnal surface 26.5 (Fig. 18), which also comprises the source of water upwelling onto the continental shelf along much of the US west coast (Pierce et al., 2012). On this and other density surfaces, the distribution of O_2 in ROMS-BEC is consistent with climatological observations, suggesting that the balance of distinct water masses and the respiratory modifications they experience in the interior of the domain are relatively well represented in the model. Thermocline nutrient distributions exhibit a similar model skill (not shown).

The O_2 in the thermocline of the CCS is highly variable, with standard deviations of ≈ 20 mmol m⁻³ that are on average 15-20% of the mean O_2 across the historical measurements (Fig. 18). The magnitudes and patterns of variance are well captured by the ROMS hindcast. In both the model hindcast and in observations, the standard deviation of monthly O_2 is ≈ 5 times larger than that of the climatological seasonal cycle. Thus, a large majority of the O_2 variability is explained by non-seasonal time-scales, including large eddy-driven fluctuations (Frenger et al., 2018) and low-frequency climate variability (Buil and Lorenzo, 2017; McClatchie et al., 2010). Both model and observations indicate that O_2 variation peaks slightly offshore, in a pattern resembling that of eddy kinetic energy. The high interannual to decadal O_2 variability observed in the central North Pacific, which reaches its maximum on the isopycnal 26.5, is not included in the climatological boundary conditions, and thus likely accounts for the model bias toward low O_2 variance in the north of the domain.

Variations in O_2 on depth surfaces in the thermocline are also dominated by interannual anoma-

lies. The fluctuating O_2 at depths of 100-200 m are highly correlated with density ($R^2 \ge 0.5$) throughout the CCS, reflecting the importance of isopycnal heaving of the background O_2 gradient. Similar correlations are observed in ROMS and World Ocean Database (Fig. 19). In the model, the largest such anomaly is associated with the ENSO event in 1997-98, in which deepened isopycnals yield high O_2 conditions that last for ≈ 1 year. The signal is recorded in the central CCS as well, although the magnitude of the anomalies is reduced by $\approx 50\%$ relative to the better-sampled event in the SCB. The variability of O_2 within an isopycnal surface can be used to account for this portion of variance, leaving only lateral circulation and respiration. We find that along $\sigma_{\theta} = 26.5$ kg m⁻³, a large fraction of O_2 variability is correlated with salinity (Fig. 16), commonly used as a proxy for tropical low-oxygen and high-salinity water transported poleward in the coastal undercurrent (Meinvielle and Johnson, 2013). Interannual variability in respiration rates on this surface also account for $\approx 20\%$ of isopycnal O_2 variance, and is in turn correlated with the depth of the density layer (Deutsch et al., 2011). Historical observations show a declining strength of correlation between S and O_2 with latitude, suggesting that variability from dynamics other than the CUC become an increasingly important source of O_2 variability to the north. Indeed, the variability of O_2 in central mode water from the open North Pacific is most pronounced on this density surface, but is not represented in the model boundary conditions. Thus the relatively constant correlation between S and O_2 across latitude in the model may stem from the lack of isopycnal variance entering the domain from the open North Pacific. A complete attribution of the observed magnitudes of O_2 variance in source waters to the CCS will require inclusion of anomalies entering from the broader North Pacific and is left for future work.

In addition to low coastal O_2 , the CCS is characterized by shallow depth horizons for aragonite saturation (Fig. 20). Below ≈ 100 m depth, CO_2 concentrations are commonly undersaturated with respect to aragonite mineral formation, and thus inhibit shell formation by calcifying organisms. The aragonite saturation state, Ω_A , is predicted to fall below saturation ($\Omega_A < 1$) along most of the coast in summer, consistent with observations in NOAA coastal surveys (*e.g.*, Feely et al. (2008)). Coastal hydrographic surveys reveal a strong mesoscale patchiness to the carbonate saturation state, likely reflecting mesoscale eddies and submesoscale features. In the multi-annual mean distribution of Ω_A , the model hindcast captures the scale and intensity of undersaturated conditions well.

Both low O_2 and low Ω_A have been implicated as primary factors mediating the influence of climate on organism fitness and species habitat in the CCS. We compare decadal trends in both these properties from the hindcast simulations to the observed changes over time. For Ω_A , the measurements are too sparse and the distribution too patchy to define a robust trend, even over the short model period. For each property, time series are shown for the regions with the most data coverage (Fig. 21): northern CCS for aragonite saturation, and southern CCS for O_2 . In both cases, the trend in the data is within the uncertainty in the measurements.

As a metric of variability in these habitat constraints, we computed the volume of water subject to hypoxic or corrosive conditions. We use a constant O_2 level of 100 mmol m⁻³ as a simple indicator of hypoxic constraints, recognizing that this value varies among species, and depends on other factors, including temperature (Deutsch et al., 2015). Corrosive conditions are defined by simple thermodynamic undersaturation ($\Omega_A < 1$), though biological sensitivities may begin at higher thresholds. Water volumes are computed as the sum of grid cell volumes with $O_2 < 100$ mmol m⁻³ or $\Omega_A < 1$ that are on the continental shelf (-z < 200 m).

The volume of hypoxic and corrosive water in the CCS varies strongly over latitude and time

(Fig. 22). For both properties, restriction of putative habitat volume is stronger to the north of Pt. Conception, opposite the latitudinal gradient of O_2 and Ω_A . The corrosive volumes are much larger than hypoxic volumes, exceeding 90% of water volume in northern latitudes during the summer upwelling season, consistent with NOAA survey data. On an annual basis, waters with a more stringent criterion for calcification ($\Omega_A < 2$) are about twice as voluminous still, primarily because the length of the season with low carbonate is broadened. Hypoxic conditions occupy a smaller fraction of shelf waters, but reach 30% of shelf water volume over a broad latitude range. The fractional coverage by hypoxia peaks around 45°N, on the Oregon coast, and it is quite small in the Southern California Bight, where O_2 declines most sharply below 200 m rather than on the shelf.

An analogous figure is shown in Renault et al. (2019) for the along-coast and temporal variability of the sea-surface height and depth of the $\sigma_{\theta} = 26.5$ kg m⁻³ depth anomalies. Relative to the quantities shown in Fig. 22, they exhibit more along-coast coherence and a more dominant seasonal cycle, with less evident interannual variability than shown here, apart from the 1997-98 ENSO event. This indicates somewhat smoother physical fields than biogeochemical ones, reflective of non-conservative biogeochemical processes acting on top of the broader patterns of physical circulation influence.

Variability in both habitat constraints is largely synchronous (Fig. 22c,d), reflecting the strong control on both O_2 and Ω_A by the effects of cumulative organic matter respiration. For both volumes, the fractional variation is similar and substantial, reaching $\approx 50\%$ of the mean across much of the latitude range. Variability of corrosive volumes is greatly attenuated south of Pt. Conception. Simulated hypoxic volumes increased sharply during the early 2000s off Oregon, when major ecosystem die-offs were attributed to the onset of extreme hypoxia there (Chan et al., 2008).



Figure 16: Correlation coefficient between O_2 and salinity anomalies in (left) ROMS, and (right) WOD along the isopycnal surface $\sigma_{\theta} = 26.5$ kg m⁻³. Prevailing negative values indicate that high S occurs when O_2 is low. To maximize data availability, the observational correlation is based on monthly anomalies in WOD from the period 1955-2013. The correlations are of similar magnitude when confined to the simulated period, 1995-2010, but are only available in the Southern California Bight.



Figure 17: Vertical sections of annual mean NO_3^- (left) and O_2 (right) (mmol m⁻³), from ROMS (left) and WOD (right) at the latitudes with regular observations by repeat hydrographic surveys. The lines span the northern CCS (upper row, ~ 44.5°N, nearest the Newport OR), the central CCS (middle row, MBARI line 67), and the southern CCS (bottom row, CalCOFI line 80). Locations of observations are shown in Fig. 3.



Figure 18: Thermocline O_2 in model simulations and observations of the climatological mean (upper) and variance (lower). All maps are interpolated to a potential density surface (26.5) surface chosen as the density class of waters upwelling onto the shelf in summer (a.k.a. source water). The mean O_2 maps are from summer months (JJA), however other seasons reveal similar model fidelity. Observed mean summer O_2 is from objectively mapped climatology (WOA). Variability is mapped as the standard deviation of monthly values from the World Ocean Database covering 1955-2013 (lower right), and it is predominantly due to interannual variability rather than the seasonal cycle (see text).



Figure 19: Correlation between O_2 and density anomalies at 100 m, in ROMS-BEC (left) and WOD (right). Interannual variations in subsurface O_2 are highly correlated with density ($R^2 \ge 0.5$) throughout the CCS, and the strength of the relationship is similar in model and observations.



Figure 20: Aragonite saturation state (Ω_A) at 100 m from observations (left) and ROMS-BEC (right). Observations are from large-scale objectively analyzed fields (color field, GLODAP2 (Lauvset and coauthors, 2016)), and from NOAA coastal surveys in the summer of 2007 (circles, Feely et al. (2008)). Model distribution is averaged over summer (JJA) from 2004-2010 climatology.



Figure 21: Trends in thermocline O_2 and Ω_A over the simulated period. Maps of the linear trend are shown in upper panels. Time series (lower panels) are shown for the regions with the most data coverage for each tracer: northern CCS for carbonate, and southern CCS for O_2 . For Ω_A all available profiles are shown. For O_2 , the mean value and standard deviation are plotted as box-whisker for each month in the WOD.



Figure 22: Top row: mean temperature and temporal trend (C year⁻¹) of water on the shelf (-z < 200 m). Middle and bottom rows: volume of corrosive and hypoxic water over time and latitude. Water volumes are computed as sum of grid cell volumes with $O_2 < 100 \text{ mmol m}^{-3}$ (bottom) and $\Omega_A < 1$ (middle) that are on the shelf (-z < 200 m). Mean values for each month are shown in the left column, and anomalies, computed as a fractional deviation from the climatological mean monthly volume at each latitude, in the middle column. Trends over time are shown in the right column.

4 Conclusions

We present model simulations of ecosystem and biogeochemical cycles in the CCS that reproduce the broad patterns of processes and states observed in this region over the past couple of decades. Our results demonstrate that productivity of the CCS reflects a complex interplay of factors. The limitation by the physical supply and removal of macronutrients (nitrate) provides the dominant seasonal and spatial pattern of NPP, but with significant constraints from light and Fe on a seasonal basis, especially in the northern CCS. Interannual variations in NPP are reasonably well predicted by fluctuations in pycnocline depth that modulate the rates of surface nutrient supply. Expanded datasets on near-surface Fe concentrations are needed to better establish its role as a limiting factor for growth in the CCS. A significant correlation between model NPP and surface irradiance suggests that changes in light are also influential. Together, these results suggest that an index of NPP that accounts for both regional pycnocline structure and cloud cover would be more skillful than one based only on coastal winds (*e.g.*, Bakun (1990); Jacox et al. (2018)). Our results highlight the value of continued measurements of the depth of the chlorophyll maximum.

Biogeochemical properties of subsurface waters in the CCS are also well reproduced by model simulations. The amplitude of interannual variability in NO_3^- at the base of the photic zone and of O_2 in the thermocline are also both strongly correlated to undulations of the pycnocline. The largest such anomalies in our simulation period were associated with the 1997-98 ENSO event, whose amplitude of density and O_2 anomalies remains coherent over a wide latitude band, albeit with declining magnitude. For NO_3^- , the overall variance is somewhat lower, and the strength of density correlations is somewhat higher in model output than in observations in the northern domain. This suggests an important role for anomalies entering the CCS from the subarctic North Pacific, an HNLC region. Basin-scale changes in biogeochemical properties are known to be exceptionally high at the gyre boundary $\approx 45^{\circ}$ N (Mecking et al., 2008), and these remote anomalies are likely to play an important but uncertain role in the variability observed in the CCS. Evaluating these remote influences using empirically-based time-dependent biogeochemical boundary conditions is an important avenue for future research.

The variability of biogeochemical properties leads to significant changes in the volume of waters characterized by biologically stressful conditions of hypoxia and carbonate undersaturation. In the volume anomalies for both habitat constraints, there is strong coherence across the CCS. Years with unusually large volumes of hypoxic or corrosive water offer few obvious latitudinal refuges. The onset of these conditions tends to propagate from the central CCS ($\approx 40^{\circ}$ N), arriving in the northern CCS with a 2-3 month delay. Thus, monitoring hypoxia and CO_2 system parameters in the central CCS may offer some seasonal predictability for northern ecosystem impacts. **Codes and Simulation Data** The physical and biogeochemical codes used for our simulations are at https://github.com/UCLA-ROMS/Code. Simulation model output archive data can be made available by email requests to the Corresponding Authors.

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5 Appendix: Biogeochemical Model

Here, for completeness, we summarize the equations of the Biogeochemical Elemental Cycling (BEC) model in the implementation used for this work. This formulation is based on the original version presented in Moore et al. (2004).

5.1 Variables and parameters

Name	Description	Units
N_{NH_4}	ammonium	mmol N/m^3
N_{NO_2}	nitrite	mmol N/m^3
N_{NO_3}	nitrate	mmol N/m^3
N_{N_2O}	nitrous oxide	mmol N/m ³
N_{NO_3}	nitrate	mmol N/m ³
P_{PO_4}	Phosphorus	mmol P/m^3
O_2	dissolved oxygen	mmol O_2/m^3
Fe	iron	nmol Fe/m^3
Si_{SiO_2}	Silicate	mmol SiO_2/m^3
C_{DIC}	dissolved inorganic carbon	mmol C/m^3
Alk	total alkalinity	mmol/m ³
C_{sp}	small phytoplankton carbon	mmol C/m^3
C_{diat}	diatom carbon	mmol C/m^3
C_{diaz}	diazotroph carbon	mmol C/m^3
Chl_{sp}	small phytoplankton carbon	mmol Chl/m^3
Chl_{diat}	diatom chlorophyll	mmol Chl/m^3
Chl_{diaz}	diazotroph chlorophyll	mmol Chl/m^3
C_{zoo}	zooplankton chlorophyll	mmol Chl/m^3
Fe_{sp}	small phytoplankton iron	mmol Fe/m^3
Fe_{diat}	diatom iron	mmol Fe/m^3
Fe_{diaz}	diazotroph iron	mmol Fe/m^3
Ca_{sp}	small phytoplankton calcium carbonate	mmol $CaCO_3/m^3$
Si_{diat}	diatom silicate	mmol SiO_2/m^3
N_{don}	dissolved organic nitrogen	mmol N/m^3
P_{dop}	dissolved organic phosphorus	mmol P/m^3
C_{doc}	dissolved organic carbon	mmol C/m^3
Fe_{dofe}	dissolved organic iron	mmol Fe/m ³

Prognostic variables

Local variables

Name	Description	Units
$J^u_{sp,no3}$	nitrate uptake by small phytoplankton	mmol N/m ³ /s
$J^{u}_{sp.nh4}$	ammonium uptake by small phytoplankton	mmol N/m ³ /s
$J^u_{diat.no3}$	nitrate uptake by diatoms	mmol N/m ³ /s
$J^u_{diat.nh4}$	ammonium uptake by diatoms	mmol N/m ³ /s
J_{sp}^{grz}	grazing loss for small phytoplankton	mmol C/m ³ /s
$J_{sn.dic}^{grz}$	grazed small phytoplankton routed to DIC	mmol C/m ³ /s
J_{sndoc}^{grz}	grazed small phytoplankton routed to DOC	mmol C/m ³ /s
$J_{sn \ noc}^{grz}$	grazed small phytoplankton routed to POC	mmol C/m ³ /s
$J^{grz}_{sn\ zoo}$	grazed small phytoplankton routed to new zooplankton	mmol C/m ³ /s
39,200	biomass	
J_{diat}^{grz}	grazing loss for diatoms	mmol C/m ³ /s
$J_{diat.dic}^{grz}$	grazed diatoms routed to DIC	mmol C/m ³ /s
$J_{diat\ doc}^{grz}$	grazed diatoms routed to DOC	mmol C/m ³ /s
$J_{diat\ noc}^{grz}$	grazed diatoms routed to POC	mmol C/m ³ /s
$J_{diat\ zoo}^{grz}$	grazed diatoms routed to new zooplankton biomass	mmol C/m ³ /s
J_{diaz}^{grz}	grazing loss for diazotrophs	mmol C/m ³ /s
$J_{diaz\ dic}^{grz}$	grazed diazotrophs routed to DIC	mmol C/m ³ /s
$J_{diaz\ doc}^{grz}$	grazed diazotrophs routed to DOC	mmol C/m ³ /s
$J_{diaz \ poc}^{grz}$	grazed diazotrophs routed to POC	mmol C/m ³ /s
$J_{diaz\ zoo}^{grz}$	grazed diazotrophs routed to new zoo biomass	mmol C/m ³ /s
J_{sp}^{agg}	aggregation loss of small phytoplankton	mmol C/m ³ /s
J_{diat}^{agg}	aggregation loss of diatoms	mmol C/m ³ /s
J_{sp}^{l}	non-grazing mortality of small phytoplankton	mmol C/m ³ /s
$J^{\hat{p}rod}_{sp.caco3}$	$CaCO_3$ production by small phytoplankton	mmol $CaCO_3/m^3/s$
$Q_{P:C}^{diaz}$	Diazotroph phosphorus to carbon ratio	no units
$J^l_{sp,dic}$	non-grazing mortality of small phytoplankton routed to DIC	mmol C/m ³ /s
$J^{l}_{sp,doc}$	non-grazing mortality of small phytoplankton routed to	mmol C/m ³ /s
	DOC	
$J^l_{sp,poc}$	non-grazing mortality of small phytoplankton routed to	mmol C/m ³ /s
	POC	
J_{diat}^l	non-grazing mortality of diatoms (mmol C/m ³ /s)	2
$J^l_{diat,dic}$	non-grazing mortality of diatoms routed to DIC	mmol C/m ³ /s
$J^l_{diat,doc}$	non-grazing mortality of diatoms routed to DOC	mmol C/m ³ /s
$J^l_{diat,poc}$	non-grazing mortality of diatoms routed to POC	mmol C/m ³ /s
J^l_{diaz}	non-grazing mortality of diazotrophs	$(\text{mmol C/m}^3/\text{s})$
$J^l_{diaz,dic}$	non-grazing mortality of diazotrophs routed to DIC	mmol C/m ³ /s
$J^l_{diaz,doc}$	non-grazing mortality of diazotrophs routed to DOC	mmol C/m ³ /s
$J^{l}_{diaz,poc}$	non-grazing mortality of diazotrophs routed to POC	mmol C/m ³ /s
J^{ι}_{zoo}	zooplankton mortality	mmol C/m ³ /s
$J_{zoo,dic}^{\iota}$	zooplankton mortality routed to DIC	mmol C/m ³ /s)
$J^{l}_{zoo,doc}$	zooplankton mortality routed to DOC	mmol C/m ³ /s

PAR	photosynthetically active radiation	W/m^2
J^{ammox}	rate of NH_4 oxidation to NO_2	mmol N/m ³ /s
J^{nit}	rate of NO_2 oxidation to NO_3	mmol N/m ³ /s
J^{WCD}	amount of NO_3 converted to N_2 by water column denitrif.	mmol N/m ³ /s
J^{BD}	amount of NO_3 converted to N_2 by benthic denitrif.	mmol N/m ³ /s
J_{remin}^{other}	reminineralization in the sediments by processes other than	
	oxic remin.	
	and denitrif.	mmol C/m ³ /s
J_{poc}^{prod}	amount of particulate organic C produced	mmol C/m ³ /s
J_{pic}^{prod}	amount of particulate inorganic C produced	mmol C/m ³ /s
J_{PSi}^{prod}	amount of particulate organic Si produced	mmol Si/m ³ /s
J_{pfe}^{prod}	amount of particulate organic Fe produced	mmol Fe/m ³ /s
J_{poc}^{remin}	remineralized particulate organic C	mmol Fe/m ³ /s
\dot{J}_{pic}^{remin}	remineralized particulate inorganic C	mmol Fe/m ³ /s
\dot{J}_{PSi}^{remin}	remineralized particulate organic Si	mmol Fe/m ³ /s
J_{pfe}^{remin}	remineralized particulate organic Fe	mmol Fe/m ³ /s
$\Phi_{poc}^{mineral}$	incoming mineral associated particulate C flux	mmol $C \text{ cm/m}^3/\text{s}$
Φ_{pcaco3}^{soft}	incoming soft $CaCO_3$ particulate flux	mmol $CaCO_3$ cm/m ³ /s
Φ_{pcaco3}^{hard}	incoming hard $CaCO_3$ particulate flux	mmol $CaCO_3$ cm/m ³ /s
$\Phi^{soft}_{SiO_2}$	incoming soft SiO_2 particulate flux	mmol $PSiO_2$ cm/m ³ /s
$\Phi_{SiO_2}^{hard}$	incoming hard SiO_2 particulate flux	mmol $PSiO_2$ cm/m ³ /s
Φ_{dust}^{soft}	incoming soft dust flux	mmol $C \text{ cm/m}^3/\text{s}$
Φ_{dust}^{hard}	incoming hard dust flux	mmol $C \text{ cm/m}^3/\text{s}$
J_{poc}^{bury}	particulate C-flux buried in the sediments	mmol $C \text{ cm/m}^3/\text{s}$
λ_{poc}	remineralization length scale for soft poc	no units
J_{diaz}^{Nfix}	total N_2 -fixation by diazotrophs	mmol N/m ³ /s
$J_{diaz}^{\tilde{Nexcrete}}$	N excreted by diazotrophs	mmol N/m ³ /s
J_{diaz}^{photoN}	N fixed by diazotroph (non-excreted)	mmol $N/m^3/s$

5.2 Ecosystem parameters

Parameters	Description	Values	Units
Model grid			
<i>.</i>	Small value	10-8	mma1/m ³
e	Small value	10	
Δz	Model layer thickness	variable	meters
$\mathbf{k}_{ ho}$	Index of model vertical	1-60	no units
	level at tracer points		
Carbon	Ĩ		

PC^{sp}_{ref}	max phyto C-specific growth rate at Tref (GD98) for small phytoplankton	3.0	1/d
PC_{ref}^{diat}	max phyto C-specific growth rate at Tref (GD98) for diatoms	3.0	1/d
PC_{ref}^{diaz}	max phyto C-specific growth rate at Tref (GD98) for diazotrophs	0.4	1/d
$lpha_{chl}$	chlorophyll-specific initial slope of P vs. I curve for diatoms and	0.3	mmol $C \text{ m}^2/(\text{mg} Chl \text{ W day})$
$lpha_{chl}^{diaz}$	chlorophyll-specific initial slope of P vs. I curve for diazotrophs	0.036	(mmol C m ² /(mg Chl W day))
Nutrient limita- tion			
$\mathbf{k}_{NO_3}^{sp}$	Small phyto. half saturation constant for NO_3	0.5	mmol/m ³
$\mathbf{k}_{NO_{3}}^{diat}$	diatom half saturation constant for NO_3 up- take	2.5	mmol/m ³
$k_{NH_4}^{sp}$	Small phyto. half saturation constant for NH_4 uptake	0.01	mmol/m ³
$\mathbf{k}_{NH_4}^{diat}$	Diatom half saturation constant for NH_4 up-take	0.1	mmol/m ³
k_{Fe}^{sp}	Small phyto. half sat- uration constant for <i>Fe</i> uptake	0.035e-3	mmol/m ³
\mathbf{k}_{Fe}^{diat}	Diat half saturation constant for Fe uptake	0.08e-3	mmol/m ³
\mathbf{k}_{Fe}^{diaz}	Diazotroph half satura- tion constant for Fe up- take	0.1e-3	mmol/m ³
k_{PO4}^{sp}	Small phyto. half saturation constant for PO_4 uptake	0.01	mmol/m ³

\mathbf{k}_{PO4}^{diat}	Diatom half saturation constant for PO_4 up-	0.1	mmol/m ³
\mathbf{k}_{PO4}^{diaz}	Diazotroph half satura- tion constant for PO_4	0.005	mmol/m ³
\mathbf{k}^{diat}_{SiO2}	Diatom half saturation constant for SiO_32 uptake	1	mmol/m ³
Fixed Stoi- chiometry			
$\mathbf{Q}_{N:C}$	Nitrogen to Carbon ra- tio	0.137	no units
$Q_{P:C}$	Small phyto. and di- atom P:C ratio	0.00855	(mmol <i>P</i> /mmol <i>C</i>)
$\mathbf{Q}_{P:C}^{diaz}$	Diazotroph P:C ratio	0.002735	(mmol <i>P</i> /mmol <i>C</i>)
Dissolved or-			
$ au_{DOM}$	Dissolved organic mat- ter remineralization in- verse timescale	0.01	1/d
N ₂ fixation			
$lpha_{ex}^{diaz}$	Ratio of N excreted by diazotrophs to total N fixed	0.3	
Chl stoichiome- try and produc- tion			
$\mathbf{Q}_{N:chl,max}^{sp,growth}$	Max ratio of Chl produced to N photosynthesized for small phyto.	2.5	mg Chl/mmol N

$\mathbf{Q}_{N:chl,max}^{diat,growth}$	Max ratio of Chl pro- duced to N photosyn- thesized for diatom	4.0	mg Chl/mmol N
$\mathbf{Q}_{N:chl,max}^{diaz,growth}$	Max ratio of Chl pro- duced to N photosyn- thesized for diaz.	3.4	mg Chl/mmol N
Light			
\mathbf{f}_{Isw}	Fraction of incoming radiation used for pho- tosynthesis	0.45	
μ_w	Radiation attenuation coefficient per unit chlorophyll	0.03	1/m/(mg <i>Chl</i> /m ³)
μ_{Chl}	Radiation attenuation coefficient for water	0.04	1/m
<i>Fe</i> stoichiome- try and cycling			
$Q^{sp,g0}_{Fe:C}$	Ratio used in the calculation of the Fe to C ratio of uptake for small phyto	6e-6	mmol Fe/mmol C
$Q^{sp,g1}_{Fe:C}$	Maximum Fe:C ratio of uptake by small phyto. if Fe $< 2 \cdot k_{T}^{sp}$.	2.5e-6	mmol Fe/mmol C
$Q_{Fe:C}^{diat,g0}$	Ratio used in the calculation of the Fe to C ratio of uptake for diatoms	6e-6	mmol Fe/mmol C
$\mathbf{Q}_{Fe:C}^{diat,g1}$	Maximum Fe:C ratio of uptake by diatoms if Fe $< 2 \cdot k_{Ee}^{diat}$	2.5e-6	mmol Fe/mmol C
$Q^{diaz,g0}_{Fe:C}$	Ratio used in the calculation of the Fe to C ratio of uptake for diazotrophs	42e-6	mmol Fe/mmol C
$Q^{diaz,g1}_{Fe:C}$	Maximum Fe:C ratio of uptake by diazotrophs if Fe $< 2 \cdot k_{Fe}^{diaz}$	14e-6	mmol Fe/mmol C

$\beta_{fe}^{max,scale1}$	Scaling parameter used in the calculation of Fe	3.0	no units
$\beta_{fe}^{thresh,1}$	Fe threshold parameter used in the calculation of Fe scavenging	0.6e-3	mmol/m ³
$\beta_{fe}^{thresh,2}$	Fe threshold parameter used in the calculation of Fe scavenging	0.5e-3	mmol/m ³
CaCO ₃ stoi-			
chiometry and production			
$\mathbf{Q}^{sp,max}_{CaCO_3:C}$	Maximum calcification to C photosynthesis ratio	0.4	mmol CaCO3/mmol C
${ m f}^{prod}_{CaCO_3}$	Initial calcification to C photosynthesis ratio	0.026	mmol CaCO3/mmol C
$T_1^{CaCO_3}$	Temperature parameter used to modify the ini- tial calcification rate	1.	°C
$\mathrm{T}_{2}^{CaCO_{3}}$	Temperature parameter used to modify the ini- tial calcification rate	-2.	°C
Si stoichiome-			
try			
$\mathrm{Q}^{diat,max}_{Si:C}$	Maximum $Q_{Si:C}$ ratio for diatoms	0.685	mmol Si/mmol C
${ m Q}^{diat,g0}_{Si:C}$	Default Si:C ratio of growth for diatoms	0.137	mmol Si/mmol C
$Q^{diat,g1}_{Si:C}$	Ratio used to calculate the Si to C multiplica- tive ratio of growth for diatoms	2.5	no units
Grazing			
β_z^{grz}	grazing coefficient, used in density de- pendent grazing modification	1.05	mmol C/m ³

$\beta_{thres}^{grz,sp,0}$	Small phytoplankton threshold concentration for grazing	0.001	mmol C/m ³
$\beta_{thres}^{grz,diat,0}$	Diatom threshold con- centration for grazing	0.02	mmol C/m^3
$\beta_{thres}^{grz,diaz,0}$	Diazotroph thresh- old concentration for grazing	0.01	mmol C/m ³
$\alpha_{sp}^{grz,zoo}$	Fraction of small phyto. grazing going to zoo- plankton	0.3	no units
$\alpha_{diat}^{grz,zoo}$	Fraction of diatom grazing going to zooplankton	0.3	no units
$\alpha_{diaz}^{grz,zoo}$	Fraction of diazotroph grazing going to zoo- plankton	0.21	no units
$\alpha_{sp}^{grz,poc}$	Default fraction of small phytop. grazing going to POC	0.22	no units
$\alpha_{diat}^{grz,poc}$	Fraction of diatom grazing going to POC	0.26	no units
$\alpha^{grz,poc}_{diaz}$	Fraction of diazotroph grazing going to POC	0.0	no units
$lpha_{sp}^{grz,doc}$	Fraction of small phyto. grazing going to DOC	0.34	no units
$lpha_{diat}^{grz,doc}$	Fraction of diatom grazing going to DOC	0.13	no units
$lpha_{diaz}^{grz,doc}$	Fraction of diazotroph grazing going to DOC	0.24	no units
$\alpha_{sp}^{grz,dic}$	Fraction of small phy- top. grazing going to DIC	0.36	no units
$lpha_{diat}^{grz,dic}$	Fraction of diatom grazing going to DIC	0.31	no units
$\alpha_{diaz}^{grz,dic}$	Fraction of diazotroph grazing going to DIC	0.55	no units
Losses			
λ_{sp}^{mort}	Small phyto. mortality 1	0.15	1/d
λ_{diat}^{mort}	Diatom mortality 1	0.15	1/d
λ_{diaz}^{mort}	Diazotroph mortality 1	0.16	1/d

$\alpha_{diat}^{l,poc}$	Fraction of diatom loss going to POC	0.05	no units
$lpha_{diaz}^{l,poc}$	Fraction of diazotroph	0	no units
\mathbf{f}_{labile}	Fraction of Labile dis- solved organic matter for loss calculations	0.70	no units
Aggregation			
$ au^{agg,min}_{diat}$	Minimum aggregation rate for diatoms	0.01	1/d
$ au^{agg.min}_{diaz}$	Minimum aggregation rate for diazotrophs	0.01	1/d
$ au_{sp}^{agg,max}$	Maximum aggregation rate for small phyto.	0.75	1/d
$ au^{agg,max}_{diat}$	Maximum aggregation rate for diatoms	0.75	1/d
$ au^{agg,max}_{diaz}$	Maximum aggregation rate for diazotrophs	0.75	1/d
λ_{sp}^{mort2}	Small phyto. quadratic mortality	0.0035	$1/(\text{mmol } \text{m}^3 \text{ d})$
λ_{diat}^{mort2}	diatom quadratic mor- tality	0.0035	$1/(\text{mmol } \text{m}^3 \text{ d})$
λ_{diaz}^{mort2}	diazotroph quadratic mortality	0.16	1/(mmol m ³ d)
N cycle rates			
$ au^{ammox}$	NH_4 oxidation inverse timescale	0.06	1/d
$ au^{nit}$	NO_2 oxidation inverse timescale	0.33	1/d
$\operatorname{PAR}_{max}^{nitrif}$	Light threshold for on- set of nitrification	4.0	W / m^2
$Q^{denit}_{N:C}$	Ratio of inorganic N consumed to POC rem- ineralized during deni- trification	104/106	mmol N / mmol POC
${ m f}_{n2o}^{denit}$	Fraction of denitrifica-	0.95	no units

 τ^{cons}_{n2o}

timescale

tion that goes to N_2O

 N_2O reduction inverse 0.33333

1/d

a_0^{bohl}	Parameter used to calculate sedimentary denitrification (Bohlen	0.06	no units
a_1^{bohl}	et al., 2012) Parameter used to calculate sedimentary denitrification (Bohlen et al. 2012)	0.19	no units
a_2^{bohl}	Parameter used to calculate sedimentary denitrification (Bohlen et al., 2012)	0.99	no units
Particle cycling ρ_{pic}	Organic carbon to inor- ganic carbon mass ratio in PIC	$0.07 \cdot \frac{100.09}{12.01}$	no units
$ ho_{psi}$	Organic carbon to inor- ganic Si mass ratio in <i>PSiO</i> ₂	$0.033 \cdot \frac{60.08}{12.01}$	no units
$ ho_{dust}$	Organic carbon to dust mass ratio in dust	$0.07 \cdot \frac{10^6}{12.01}$	no units

5.3 Model equations

5.3.1 Tracer equations

Here the symbol d/dt denotes the sum of the local time derivative and the physical transport.

$$\frac{d}{dt}(N_{no3}) = J^{nit} - J^{denit}_{no3,no2} - J^{denit}_{sed} - (J^u_{diat,no3} + J^u_{sp,no3})$$
(A1)

$$\frac{d}{dt}(N_{nh4}) = Q_{N:C}J_{poc}^{remin} + \tau_{dom}N_{don} + Q_{N:C}(J_{diat,dic}^{l} + J_{sp,dic}^{l} + J_{diaz,dic}^{l}) + Q_{N:C}J_{zoo,dic}^{l} + Q_{N:C}(J_{diat,dic}^{grz} + J_{sp,dic}^{grz} + J_{diaz,dic}^{grz}) - J^{nit} - (J_{diat,nh4}^{u} + J_{sp,nh4}^{u})$$
(A2)

$$\frac{d}{dt}(N_{no2}) = J^{ammox} - J^{nit} - J^{denit}_{no2,n2o}$$
(A3)

$$\frac{d}{dt}(N_{n2o}) = 0.5J_{no2,n2o}^{denit} - J_{n2o,n2}^{denit}$$
(A4)

$$\frac{d}{dt}(N_{n2}) = J_{n2o,n2}^{denit} + 0.5 J_{sed}^{denit}$$
(A5)

$$\frac{d}{dt}(Fe_{fe}) = J_{DOfe,fe} - J_{Fe}^{scav} + (Q_{Fe:C}^{zoo}J_{zoo,dic}^{l}) + J_{remin}^{Fe} - J_{diaz,Fe}^{u}
+ Q_{Fe:C}^{sp}(J_{sp,dic}^{l} + J_{sp,dic}^{grz}) + Q_{Fe:C}^{diat}(J_{diat,dic}^{l} + J_{diat,dic}^{grz})
+ Q_{Fe:C}^{diaz}(J_{diaz,dic}^{l} + J_{diaz,dic}^{grz}) - J_{sp,Fe}^{u} - J_{diat,Fe}^{u}
+ J_{sp,zoo}^{grz}(Q_{Fe:C}^{sp} - Q_{Fe:C}^{zoo}) + J_{diat,zoo}^{grz}(Q_{Fe:C}^{diat} - Q_{Fe:C}^{zoo})
+ J_{diaz,zoo}^{grz}(Q_{Fe:C}^{diaz} - Q_{Fe:C}^{zoo}) + J_{Fe}^{Sed}$$
(A6)

$$\frac{d}{dt}(P_{po4}) = \frac{1}{Q_{P:C}} (J_{poc}^{remin} + J_{zoo,dic}^{l} + J_{sp,dic}^{l} + J_{diat,dic}^{l} + J_{sp,dic}^{grz} + J_{diat,dic}^{grz})
+ J_{diaz,dip}^{l} - (J_{sp,P}^{u} + J_{diat,P}^{u} + J_{diaz,P}^{u}) + J_{DOP,po4}$$
(A7)

$$\frac{d}{dt}(Si_{sio2}) = \frac{1}{Q_{Si:C}} (J_{DOC,dic} + 0.5 \quad J_{diat,dic}^{grz} + 0.95 \quad J_{zoo}^{l})
- J_{diat,Si}^{u} + J_{remin}^{Si}$$
(A8)

$$If (O_{2} > O_{2}min)$$

$$\frac{d}{dt}(O_{2}) = (J_{sp,C}^{u} + J_{diat,C}^{u} + J_{diaz,C}^{u})/R_{C:O}^{d} + ((-J_{poc}^{remin} - J_{DOC,dic} - J_{zoo,dic}^{l} - J_{sp,dic}^{l} - J_{sp,dic}^{grz} - J_{diat,dic}^{l} - J_{diaz,dic}^{grz} - J_{diaz,dic}^{l} - J_{diaz,dic}^{grz})/R_{C:O}^{d})$$

$$- 1.5 J^{ammox} - 0.5 J^{nit} - J_{remin}^{poc}/(R_{C:O}^{d})$$
(A9)

If $(O_2 \leq O_2 min)$

$$\frac{d}{dt}(O_2) = (J^u_{sp,C} + J^u_{diat,C} + J^u_{diaz,C})/R^d_{C:O}$$
(A10)

$$\frac{d}{dt}(DIC) = J_{DOC,dic} + 0.33 \quad J_{sp}^{grz} \quad Q_{C:caco3}
+ J_{zoo,dic}^{l} + J_{sp,dic}^{l} + J_{sp,dic}^{grz} + J_{diat,dic}^{l} + J_{diat,dic}^{grz} - J_{sp,C}^{u} - J_{diat,C}^{u}
- J_{sp,caco3}^{prod} + J_{diaz,dic}^{grz} + J_{diaz,dic}^{l} - J_{diaz,C}^{u} + (J_{poc}^{remin} + J_{pic}^{remin})$$
(A11)

$$\frac{d}{dt}(Alk) = -\frac{dN_{no3}}{dt} + \frac{dN_{nh4}}{dt} + 2 \quad J_{pic}^{remin} + 0.33 \quad J_{sp}^{grz} \quad Q_{C:caco3} - J_{sp,caco3}^{prod} + 2 \quad J_{pic}^{remin} \quad (A12)$$

$$\frac{d}{dt}(C_{sp}) = J_{sp,C}^{photo} - (J_{sp}^{grz} + J_{sp}^{l} + J_{sp}^{agg})$$
(A13)

$$\frac{d}{dt}(Chl_{sp}) = J_{sp,chl}^{pa} - Q_{Chl:C}^{sp}(J_{sp}^{grz} + J_{sp}^{l} + J_{sp}^{agg})$$
(A14)

$$\frac{d}{dt}(Fe_{sp}) = gQ_{Fe:C}^{sp}J_{sp,C}^{photo} - Q_{Fe:C}^{sp}(J_{sp}^{grz} + J_{sp}^{l} + J_{sp}^{agg})$$
(A15)

$$\frac{d}{dt}(Ca_{sp}) = J_{sp,caco3}^{prod} + J_{sp,nh4}^{u}) - Q_{caco3}(J_{sp}^{grz} + J_{sp}^{l} + J_{sp}^{agg})$$
(A16)

$$\frac{d}{dt}(C_{diat}) = J_{diat,C}^{photo} - (J_{diat}^{grz} + J_{diat}^{l} + J_{diat}^{a})$$
(A17)

$$\frac{d}{dt}(Fe_{diat}) = gQ_{Fe:C}^{diat}J_{diat,C}^{photo} - Q_{Fe:C}^{diat}(J_{diat}^{grz} + J_{diat}^{l} + J_{diat}^{agg})$$
(A18)

$$\frac{d}{dt}(Chl_{diat}) = J^{pa}_{diat,chl} - Q^{diat}_{Chl:C}(J^{grz}_{diat} + J^{l}_{diat} + J^{agg}_{diat})$$
(A19)

$$\frac{d}{dt}(Si_{diat}) = gQ_{Si:C}J_{diat,C}^{photo} - Q_{Si:C}^{diat}(J_{diat}^{grz} + J_{diat}^{l} + J_{diat}^{agg})$$
(A20)

$$\frac{d}{dt}(C_{diaz}) = J^{photo}_{diaz,C} - \left(J^{grz}_{diaz} + J^{l}_{diaz}\right)$$
(A21)

$$\frac{d}{dt}(Fe_{diaz}) = gQ_{Fe:C}^{diaz}J_{diaz,C}^{photo} - Q_{Fe:C}^{diaz}(J_{diaz}^{grz} + J_{diaz}^{l} + J_{diaz}^{agg})$$
(A22)

$$\frac{d}{dt}(Chl_{diaz}) = J^{pa}_{diaz,chl} - Q^{diaz}_{Chl:C}(J^{grz}_{diaz} + J^l_{diaz} + J^{agg}_{diaz})$$
(A23)

$$\frac{d}{dt}(C_{zoo}) = (J_{sp,zoo}^{grz} + J_{diat,zoo}^{grz} + J_{diaz,zoo}^{grz}) - J_{zoo}^{l}$$
(A24)

$$\frac{d}{dt}(C_{doc}) = J_{sp,doc}^{l} + J_{diat,doc}^{l} + J_{diaz,doc}^{l} + J_{zoo,doc}^{l} + (J_{sp,doc}^{grz} + J_{diat,doc}^{grz} + J_{diaz,doc}^{grz}) - \tau_{dom} C_{doc}$$
(A25)

$$\frac{d}{dt}(N_{don}) = Q_{N:C} \left(J_{sp,doc}^{l} + J_{diat,doc}^{l} + J_{diaz,doc}^{l} + J_{zoo,doc}^{l} + J_{sp,doc}^{grz} + J_{diat,doc}^{grz} + J_{diaz,doc}^{grz}\right) - \tau_{dom} N_{don} + J_{diaz,N}^{excrete}$$
(A26)

$$\frac{d}{dt}(P_{dop}) = Q_{P:C} \left(J_{sp,doc}^{l} + J_{diat,doc}^{l} + J_{sp,doc}^{grz} + J_{diat,doc}^{grz} + J_{zoo,doc}^{l}\right)
+ J_{diaz,dop}^{l} - \tau_{dom} P_{dop}$$
(A27)

$$\frac{d}{dt}(Fe_{dofe}) = Q_{Fe:C}^{sp}\left(J_{sp,doc}^{l} + J_{sp,doc}^{grz}\right) + Q_{Fe:C}^{diat}\left(J_{diat,doc}^{l} + J_{diat,doc}^{grz}\right) + Q_{Fe:C}^{diaz}\left(J_{diaz,doc}^{l} + J_{diaz,doc}^{grz}\right) + Q_{Fe:C}^{zoo}\left(J_{zoo,doc}^{l} - \tau_{dom} Fe_{dofe}\right)$$
(A28)

5.3.2 Treatment of particulate organic matter

General model

Particulate organic matter is produced and instantaneously distributed over the depth of the water column following the exponential solution to the steady-state 1-dimensional production-remineralization equation:

$$\frac{\partial \Phi(z)}{\partial z} = J^{remin}(z) - J^{prod}(z) = -\frac{\Phi(z)}{\lambda} + J^{prod}(z), \tag{A29}$$

where Φ is a flux, λ is the remineralization length-scale, J^{prod} is the known production rate within the layer, and J^{remin} the remineralization rate within the layer, which needs to be determined. For a single layer, assuming the flux at the top of the layer $\Phi(k)$ is known, and the production $J^{prod}(k)$ is constant within the layer, the solution to equation A29, can be cast to determined the flux out of the layer, $\Phi(k-1)$, for each element *i*:

$$\Phi_i(k-1) = \Phi_i(k) e^{-\frac{\Delta z}{\lambda_i}} + J^{prod}(k) \left(1 - e^{-\frac{\Delta z}{\lambda_i}}\right) \lambda,$$
(A30)

Particulate organic carbon (POC) is partitioned between a free and mineral associated component:

$$\Phi_{poc}(z) = \Phi_{poc}^{free}(z) + \Phi_{poc}^{mineral}(z)$$
(A31)

 $\Phi_{poc}^{mineral}$ can be associated with CaCO₃, SiO₂ or dust. Each mineral-associated POC flux is further partitioned into a "soluble" component, which remineralizes with the length-scale of the associated mineral, and a "non-soluble" component which remineralizes with a length-scale of 40,000 m.

$$\Phi_{poc}^{mineral}(z) = \rho_{pic} \left(\Phi_{pic}^{sol}(z) + \Phi_{pcaco3}^{nonsol}(z) \right) + \rho_{psi} \left(\Phi_{psi}^{sol}(z) + \Phi_{psi}^{nonsol}(z) \right) + \rho_{dust} \left(\Phi_{dust}^{sol}(z) + \Phi_{dust}^{nonsol}(z) \right)$$
(A32)

For all components of the fluxes except Fe, the flux out of the layer, $\Phi(k-1)$, is computed first, with knowledge of the source within the layer, $J^{prod}(k)$, and of the remineralization length-scale. Remineralization in the layer is then calculated from conservation, *i.e.*, from (A29). Below, for each component, we list the equations used to determine the production terms $J^{prod}(k)$, followed by the fluxes out, $\Phi(k-1)$, and finally the remineralization terms, $J^{remin}(z)$, which enter the tracer conservation equations.

Production

$$J_{poc}^{prod} = \left(J_{sp,poc}^{grz} + J_{diat,poc}^{grz} + J_{diaz,poc}^{grz}\right) + \left(J_{sp}^{agg} + J_{diat}^{agg}\right) \\ + \left(J_{sp,poc}^{l} + J_{diat,poc}^{l} + J_{diaz,poc}^{l}\right) + f_{zoo}^{d} J_{zoo}^{l}$$
(A33)

$$J_{pic}^{prod} = \left(0.67 J_{sp}^{grz} + J_{sp}^{agg} + J_{sp}^{l}\right) Q_{caco3:C}^{sp}$$
(A34)

$$J_{psi}^{prod} = \left(0.5 J_{sp}^{grz} + 0.05 J_{sp}^{l}\right) Q_{Si:C}^{diat}$$
(A35)

Available production for free POC is then:

$$J_{poc,avail}^{prod} = J_{poc}^{prod} - \rho_{pic} J_{pic}^{prod} - \rho_{psi} J_{psi}^{prod}$$
(A36)

Fluxes out

Temperature dependency is used to modify the remineralization length scales of particulate organic carbon, POC, and opal, SiO₂:

$$T_{poc} = (Q_{10,poc})^{\frac{T-30}{10}} \tag{A37}$$

$$T_{Psi} = (Q_{10,psi})^{\frac{T-30}{10}}$$
(A38)

Free POC flux equation:

$$\Phi_{poc}^{free}(k-1) = \Phi(k)_{poc}^{free} e^{-\frac{\Delta z T_{poc}}{\lambda_{poc}}} + J_{poc,avail}^{prod}(k) \left(1 - e^{-\frac{\Delta z T_{poc}}{\lambda_{poc}}}\right) \frac{\lambda_{poc}}{T_{poc}}$$
(A39)

Soluble mineral-associated POC flux equation:

$$\Phi_{pic}^{sol}(k-1) = \Phi(k)_{pic}^{sol} e^{-\frac{\Delta z}{\lambda_{pic}}} + J_{pic}^{prod}(k) \left(1 - \gamma_{pic}\right) \left(1 - e^{-\frac{\Delta z}{\lambda_{pic}}}\right) \lambda_{pic}$$
(A40)

$$\Phi_{psi}^{sol}(k-1) = \Phi(k)_{psi}^{sol} e^{-\frac{\Delta z \, T_{psi}}{\lambda_{psi}}} + J_{poc}^{prod}(k) \left(1 - \gamma_{psi}\right) \left(1 - e^{-\frac{\Delta z \, T_{psi}}{\lambda_{psi}}}\right) \frac{\lambda_{psi}}{T_{psi}}$$
(A41)

$$\Phi_{dust}^{sol}(k-1) = \Phi(k)_{dust}^{sol} e^{-\frac{\Delta z}{\lambda_{dust}}}$$
(A42)

Non-soluble mineral-associated POC flux equation:

$$\Phi_{pic}^{nonsol}(k-1) = \Phi(k)_{pic}^{sol} e^{-\frac{\Delta z}{\lambda_{nonsol}}} + J_{pic}^{prod}(k) \gamma_{pic} \Delta z$$
(A43)

$$\Phi_{psi}^{nonsol}(k-1) = \Phi(k)_{psi}^{sol} e^{-\frac{\Delta z \, T_{psi}}{\lambda_{nonsol}}} + J_{poc}^{prod}(k) \,\gamma_{psi} \,\Delta z \tag{A44}$$

$$\Phi_{dust}^{nonsol}(k-1) = \Phi(k)_{dust}^{nonsol} e^{-\frac{\Delta z}{\lambda_{nonsol}}}$$
(A45)

Remineralization

Remineralization is computed from conservation, *i.e.*, (A29):

$$J^{remin} = J^{prod} + \frac{\partial \Phi(z)}{\partial z}$$
(A46)

Numerically, for each individual layer, we have:

$$J_{poc}^{remin}(k) = J_{poc}^{prod}(k) + \frac{\Phi_{poc}^{free}(k) - \Phi_{poc}^{free}(k-1) + \Phi_{poc}^{mineral}(k) - \Phi_{poc}^{mineral}(k-1)}{\Delta z}$$
(A47)

$$J_{pic}^{remin}(k) = J_{pic}^{prod}(k) + \frac{\Phi_{pic}^{sol}(k) - \Phi_{pic}^{sol}(k-1) + \Phi_{pic}^{nonsol}(k) - \Phi_{pic}^{nonsol}(k-1)}{\Delta z}$$
(A48)

$$J_{psi}^{remin}(k) = J_{psi}^{prod}(k) + \frac{\Phi_{psi}^{sol}(k) - \Phi_{psi}^{sol}(k-1) + \Phi_{psi}^{nonsol}(k) - \Phi_{psi}^{nonsol}(k-1)}{\Delta z}$$
(A49)

$$J_{dust}^{remin}(k) = \frac{\Phi_{dust}^{sol}(k) - \Phi_{dust}^{sol}(k-1) + \Phi_{dust}^{nonsol}(k) - \Phi_{dust}^{nonsol}(k-1)}{\Delta z}$$
(A50)

Particulate Fe

Production is as follows:

$$J_{pfe}^{prod} = (J_{sp}^{agg} + J_{sp,poc}^{grz} + J_{sp,poc}^{l})Q_{Fe:C}^{sp} + (J_{diat}^{agg} + J_{diat,poc}^{grz} + J_{diat,poc}^{l})Q_{Fe:C}^{diat} + (J_{diaz}^{agg} + J_{diaz,poc}^{grz})Q_{Fe:C}^{diaz} + 0.1 J_{Fe}^{scav}$$
(A51)

Particulate Fe remineralization is assumed to be proportional to POC remineralization:

$$J_{pfe}^{remin}(k) = J_{poc}^{remin}(k) \frac{\Phi_{pfe}^{sol} + \Phi_{pfe}^{nonsol}}{\Phi_{poc}^{free} + \Phi_{poc}^{mineral}}$$
(A52)

The flux out can then be computed from conservation:

$$\Phi_{pfe}(k-1) = \Phi_{pfe}(k) + \Delta z \left(J_{pfe}^{prod}(k) - J_{pfe}^{remin}(k) \right)$$
(A53)

5.3.3 Biogeochemical rates

Carbon

$$J_{sp,C}^{photo} = \operatorname{PC}_{\operatorname{ref}}^{\operatorname{sp}} f_{nut}^{sp} T_{func} \left(1 - e^{-\alpha_{chl}(Chl_{sp}/C_{sp})} \right) C_{sp}$$
(A54)

$$J_{diat,C}^{photo} = \operatorname{PC}_{\operatorname{ref}}^{\operatorname{diat}} f_{nut}^{diat} T_{func} \left(1 - e^{-\alpha_{chl}(Chl_{diat}/C_{diat})} \right) C_{diat}$$
(A55)

$$J_{diaz,C}^{photo} = \operatorname{PC}_{\operatorname{ref}}^{\operatorname{diaz}} f_{nut}^{diaz} T_{func} \left(1 - e^{-\alpha_{chl}(Chl_{diaz}/C_{diaz})} \right) C_{diaz}$$
(A56)

Nutrient limitation

$$f_{nut}^{sp} = \min[V_{NO_3}^{sp} + V_{NH_4}^{sp}, V_{Fe}^{sp}, V_{PO_4}^{sp}]$$
(A57)

$$f_{nut}^{diat} = \min[V_{NO_3}^{diat} + V_{NH_4}^{diat}, V_{Fe}^{diat}, V_{SiO_3}^{diat}, V_{PO_4}^{diat}]$$
(A58)

$$f_{nut}^{sp} = \min[V_{Fe}^{diaz}, V_{PO_4}^{diaz}]$$
(A59)

$$V_{NO_3}^{sp} = \frac{N_{NO_3}/k_{no3}^{sp}}{1 + N_{NO_3}/k_{no3}^{sp} + N_{NH_4}/k_{nh4}^{sp}}$$
(A60)

$$V_{NO_3}^{diat} = \frac{N_{NO_3}/k_{no3}^{diat}}{1 + N_{NO_3}/k_{no3}^{diat} + N_{NH_4}/k_{nh4}^{diat}}$$
(A61)

$$V_{NH_4}^{sp} = \frac{N_{NH_4}/k_{nh4}^{sp}}{1 + N_{NO_3}/k_{no3}^{sp} + N_{NH_4}/k_{nh4}^{sp}}$$
(A62)

$$V_{NH_4}^{diat} = \frac{N_{NH_4}/k_{nh4}^{diat}}{1 + N_{NO_3}/k_{no3}^{diat} + N_{NH_4}/k_{nh4}^{diat}}$$
(A63)

$$V_{Fe}^{sp} = \frac{Fe}{Fe + k_{fe}^{sp}} \tag{A64}$$

$$V_{Fe}^{diat} = \frac{Fe}{Fe + k_{fe}^{diat}}$$
(A65)

$$V_{Fe}^{diaz} = \frac{Fe}{Fe + k_{fe}^{diaz}} \tag{A66}$$

$$V_{PO_4}^{sp} = \frac{PO_4}{PO_4 + k_{po4}^{sp}}$$
(A67)

$$V_{PO_4}^{diat} = \frac{PO_4}{PO_4 + k_{po4}^{diat}} \tag{A68}$$

$$V_{PO_4}^{diaz} = \frac{PO_4}{PO_4 + k_{po4}^{diaz}} \tag{A69}$$

$$V_{SiO_2}^{diat} = \frac{Si_{sio3}}{Si_{sio2} + k_{sio2}^{diat}}$$
(A70)

NO_3 and NH_4 uptake

$$J_{sp,no3}^{u} = Q_{N:C} \frac{V_{NO_3}^{sp}}{V_{NO_3}^{sp} + V_{NH_4}^{sp}} J_{sp,C}^{photo}$$
(A71)

$$J_{sp,nh4}^{u} = Q_{N:C} \frac{V_{NH_4}^{sp}}{V_{NO_3}^{sp} + V_{NH_4}^{sp}} J_{sp,C}^{photo}$$
(A72)

$$J_{diat,no3}^{u} = Q_{N:C} \frac{V_{NO_3}^{diat}}{V_{NO_3}^{diat} + V_{NH_4}^{diat}} J_{diat,C}^{photo}$$
(A73)

$$J_{diat,nh4}^{u} = Q_{N:C} \frac{V_{NH_4}^{diat}}{V_{NO_3}^{diat} + V_{NH_4}^{diat}} J_{diat,C}^{photo}$$
(A74)

N_2 fixation

$$J_{diaz,N}^{photo} = Q_{N:C} J_{diaz,C}^{photo}$$
(A75)

$$J_{diaz}^{nfix} = J_{diaz,N}^{photo} / (1 - \alpha_{ex}^{diaz})$$
(A76)

$$J_{diaz,N}^{ex} = \alpha_{ex}^{diaz} J_{diaz}^{nfix}$$
(A77)

Chl stoichiometry and production

$$Q_{Chl:C}^{sp} = Chl_{sp} / (C_{sp} + \epsilon) \tag{A78}$$

$$Q_{Chl:C}^{diat} = Chl_{diat} / (C_{diat} + \epsilon)$$
(A79)

$$Q_{Chl:C}^{diaz} = Chl_{diaz} / (C_{diaz} + \epsilon)$$
(A80)

$$J_{sp,chl}^{pa} = Q_{N:chl,max}^{sp,growth} \frac{J_{sp,C}^{photo}/C_{sp}}{\alpha_{chl} Q_{Chl:C}^{sp} \text{ PAR }} \frac{Q_{N:C} J_{sp,C}^{photo}}{C_{sp} Q_{Chl:C}^{sp}}$$
(A81)

$$J_{diat,chl}^{pa} = Q_{N:chl,max}^{diat,growth} \frac{J_{diat,C}^{photo}/C_{diat}}{\alpha_{chl} Q_{Chl:C}^{diat} \operatorname{PAR}} \frac{Q_{N:c} J_{diat,C}^{photo}}{C_{diat} Q_{Chl:C}^{diat}}$$
(A82)

$$J_{diaz,chl}^{pa} = Q_{N:chl,max}^{diaz,growth} \frac{J_{diaz,C}^{photo}/C_{diaz}}{\alpha_{chl} Q_{Chl:C}^{diaz} \operatorname{PAR}} \frac{Q_{N:c} J_{diaz,C}^{photo}}{C_{diaz} Q_{Chl:C}^{diaz}}$$
(A83)

Light

$$\operatorname{PAR}_{B}(k-1) = \begin{cases} \max(0, f_{Isw} I_{SW}), & \text{if } k = k_{surf} \\ \operatorname{PAR}_{B}(k) e^{(\mu_{chl}(Chl_{sp}+Chl_{diat}+Chl_{diaz})+\mu_{w})\Delta z} & \text{elsewhere} \end{cases}$$
(A84)

$$PAR(k) = \frac{PAR_B(k+1) - PAR_B(k)}{\mu_{chl}(Chl_{sp} + Chl_{diat} + Chl_{diaz}) + \mu_w)\Delta z}$$
(A85)

Fe stoichiometry and cycling

$$Q_{Fe:C}^{sp} = Fe_{sp}/C_{sp} \tag{A86}$$

$$Q_{Fe:C}^{diat} = Fe_{diat}/C_{diat} \tag{A87}$$

$$Q_{Fe:C}^{diaz} = Fe_{diaz}/C_{diaz} \tag{A88}$$

$$Q_{Fe:C}^{sp,g} = \begin{cases} Q_{Fe:C}^{sp,g0}, & \text{if } Fe \ge 2 \, k_{fe}^{sp} \\ \max(Q_{Fe:C}^{sp,g1}, \frac{Q_{Fe:C}^{sp,g0} Fe}{2 \, k_{fe}^{sp}}), & \text{otherwise} \end{cases}$$
(A89)

$$Q_{Fe:C}^{diat,g} = \begin{cases} Q_{Fe:C}^{diat,g0}, & \text{if } Fe \ge 2 \, k_{fe}^{diat} \\ \max(Q_{Fe:C}^{diat,g1}, \frac{Q_{Fe:C}^{sp,g0} \, Fe}{2 \, k_{fe}^{diat}}), & \text{otherwise} \end{cases}$$
(A90)

$$Q_{Fe:C}^{diaz,g} = \begin{cases} Q_{Fe:C}^{diaz,g0}, & \text{if } Fe \ge 2 \, k_{fe}^{diaz} \\ \max(Q_{Fe:C}^{diaz,g1}, \frac{Q_{Fe:C}^{diaz,g0} \, Fe}{2 \, k_{fe}^{diaz}}), & \text{otherwise} \end{cases}$$
(A91)

$$J_{Fe}^{scav} = \begin{cases} (J_{Fe}^{scav,0} \min[(\Phi_{poc} + 8.33 \cdot 10^4 \, \Phi_{dust}^{hard}) / \Phi_{poc}^{ref}, \, \beta_{fe}^{max,scale1}] \\ + (Fe - \beta_{fe}^{thresh,1})) Fe, \\ (J_{Fe}^{scav,0} \min[(\Phi_{poc} + 8.33 \cdot 10^4 \, \Phi_{dust}^{hard}) / \Phi_{poc}^{ref}, \, \beta_{fe}^{max,scale1}] \\ + \frac{Fe}{\beta_{fe}^{thresh,2}})) Fe, \end{cases}$$
(A92)

$$J_{Fe}^{Sed} = \begin{cases} 10^{(2.5-0.0165 \max[1,O_2])} (0.001/86400) \Delta z, & \text{if } k = k_{bottom} \\ 0, & \text{elsewhere} \end{cases}$$
(A93)

75% of J_{Fe}^{Sed} is released in the bottom layer and 25 % in the layer above.

$CaCO_3$ stoichiometry production

$$Q_{caco3:C}^{sp} = \begin{cases} CaCO_{3\,sp}/C_{sp} & \text{if } CaCO_{3\,sp}/C_{sp} \le Q_{caco3:C}^{sp,max} \\ Q_{caco3:C}^{sp,max} & \text{otherwise} \end{cases}$$
(A94)

$$J_{sp,caco3}^{prod,max} = f_{caco3}^{prod} J_{sp,C}^{photo} / C_{sp} f_{nut}^{sp}$$
(A95)

$$J_{sp,caco3}^{prodT} = \begin{cases} J_{sp,caco3}^{prodmax} & \text{if } T \ge T_1^{caco3} \\ J_{sp,caco3}^{prodmax} \frac{\max[T - T_1^{caco3}, 0]}{T_1^{caco3} - T_2^{caco3}} & \text{if } T < T_1^{caco3} \text{and } C_{sp} < 3.0 \end{cases}$$
(A96)

$$J_{sp,caco3}^{prod} = \begin{cases} \min[(J_{sp,caco3}^{prodT}C_{sp}/3, 0.4J_{sp,C}^{photo}] & \text{where } C_{sp} > 3.0\\ J_{sp,caco3}^{prod,T} & \text{elsewhere} \end{cases}$$
(A97)

Si stoichiometry

$$Q_{Si:C}^{diat} = \max(Si_{diat}/C_{diat}, Q_{Si:C}^{diat,max})$$
(A98)

$$Q_{Si:C}^{diat,g} = \begin{cases} [Q_{Si:C}^{diat,g0}(2Q_{Si:C}^{diat,g1}k_{fe}^{diat}/Fe - Q_{Si:C}^{diat,g1} + 1), Q_{Si:C}^{diat,max}] \\ & \text{if } 0 < Fe < 2k_{fe}^{diat} \text{and } Si_{sio2} > 2k_{sio2}^{diat} \\ Q_{Si:C}^{diat,max} & \text{if } Fe = 0 \\ Q_{Si:C}^{diat,g0} & \text{otherwise} \end{cases}$$
(A99)

Grazing

$$\beta_{thres}^{grz,sp} = \begin{cases} \beta_{thres,0}^{grz,sp} \cdot (200 - z)/100 & z \le 100m \\ \beta_{thres,0}^{grz,sp} \cdot (200 - z)/100 & 100 \text{ m} < z < 200 \text{ m} \\ 0 & z \ge 200 \text{ m} \end{cases}$$
(A100)

$$P'_{sp} = \max(C_{sp} - \beta^{sp}_{grz}, 0) \tag{A101}$$

$$\beta_{thres}^{grz,diat} = \begin{cases} \beta_{thres,0}^{grz,diat} & z \le 100m \\ \beta_{thres,0}^{grz,diat} \cdot (200 - z)/100 & 100 \text{ m} < z < 200 \text{ m} \\ 0 & z \ge 200 \text{ m} \end{cases}$$
(A102)

$$P'_{diat} = \max(C_{diat} - \beta^{diat}_{grz}, 0) \tag{A103}$$

$$\beta_{thres}^{grz,diaz} = \begin{cases} \beta_{thres,0}^{grz,diaz} & z \le 100m\\ \beta_{thres,0}^{grz,diaz} \cdot (200-z)/100 & 100 \text{ m} < z < 200 \text{ m}\\ 0 & z \ge 200 \text{ m} \end{cases}$$
(A104)

$$P'_{diaz} = \max(C_{diaz} - \beta^{diaz}_{grz}, 0)$$
(A105)

$$J_{sp}^{grz} = J_{sp}^{g,max} T_{func} \quad \frac{P_{sp}^{\prime 2}}{P_{sp}^{\prime 2} + (\beta_z^{grz})^2} C_{zoo}$$
(A106)

$$J_{diat}^{grz} = J_{diat}^{g,max} T_{func} \quad \frac{P_{diat}^{\prime 2}}{P_{diat}^{\prime 2} + 0.81(\beta_z^{grz})^2} C_{zoo}$$
(A107)

$$J_{diaz}^{grz} = J_{diaz}^{g,max} T_{func} \quad \frac{P_{diaz}'^2}{P_{diaz}'^2 + (\beta_z^{grz})^2} C_{zoo}$$
(A108)

$$J_{sp}^{g,zoo} = \alpha_{sp}^{grz,zoo} J_{sp}^{grz}$$
(A109)

$$J_{diat}^{g,zoo} = \alpha_{diat}^{grz,zoo} J_{diat}^{grz}$$
(A110)

$$J_{diaz}^{g,zoo} = \alpha_{diaz}^{grz,zoo} J_{diaz}^{grz}$$
(A111)

$$J_{sp}^{g,poc} = \max(Q_{caco3:C}^{sp,max} Q_{caco3:C}^{sp}, \min[0.18 P_{sp}', \alpha_{sp}^{grz,poc})] J_{sp}^{grz}$$
(A112)

$$J_{diat}^{g,poc} = \alpha_{diat}^{grz,poc} J_{diat}^{grz}$$
(A113)

$$J_{diaz}^{g,poc} = \alpha_{diaz}^{grz,poc} J_{diaz}^{grz}$$
(A114)

$$J_{sp}^{g,doc} = \alpha_{sp}^{grz,doc} J_{sp}^{grz} \tag{A115}$$

$$J_{diat}^{g,doc} = \alpha_{diat}^{grz,doc} J_{diat}^{grz}$$
(A116)

$$J_{diaz}^{g,doc} = \alpha_{diaz}^{grz,doc} J_{diaz}^{grz}$$
(A117)

$$J_{sp}^{g,dic} = \alpha_{sp}^{grz,dic} J_{sp}^{grz} \tag{A118}$$

$$J_{diat}^{g,dic} = \alpha_{diat}^{grz,dic} J_{diat}^{grz}$$
(A119)

$$J_{diaz}^{g,dic} = \alpha_{diaz}^{grz,dic} J_{diaz}^{grz}$$
(A120)

Losses

$$J_{sp}^{l} = \lambda_{sp}^{mort} P_{sp}^{\prime} \tag{A121}$$

$$J_{diat}^{l} = \lambda_{diat}^{mort} P_{diat}^{\prime} \tag{A122}$$

$$J_{diaz}^{l} = \lambda_{diaz}^{mort} P_{diaz}^{\prime} \tag{A123}$$

$$J_{sp}^{l,poc} = Q_{caco3:C}^{sp} J_{sp}^{l}$$
(A124)

$$J_{diat}^{l,poc} = \alpha_{diat}^{l,poc} J_{diat}^{l}$$
(A125)

$$J_{diaz}^{l,poc} = \alpha_{diaz}^{l,poc} J_{diaz}^{l}$$
(A126)

$$J_{sp}^{l,doc} = (1 - f_{labile}) (J_{sp}^{l} - J_{sp}^{l,poc}) J_{sp}^{l}$$
(A127)

$$J_{diat}^{l,doc} = (1 - f_{labile}) (J_{diat}^{l} - J_{diat}^{l,poc}) J_{diat}^{l}$$
(A128)

$$J_{diaz}^{l,doc} = (1 - f_{labile}) (J_{diaz}^{l} - J_{diaz}^{l,poc}) J_{diaz}^{l}$$
(A129)

$$J_{sp}^{l,dic} = f_{labile} (J_{sp}^{l} - J_{sp}^{l,poc}) J_{sp}^{l}$$
(A130)

$$J_{diat}^{l,dic} = f_{labile} (J_{sp}^l - J_{diat}^{l,poc}) J_{diat}^l$$
(A131)

$$J_{diaz}^{l,dic} = f_{labile} (J_{sp}^l - J_{diaz}^{l,poc}) J_{diaz}^l$$
(A132)

Aggregation

$$J_{sp}^{agg} = \min[\tau_{sp}^{agg,max} P_{sp}', \lambda_{sp}^{mort2} P_{sp}'^2]$$
(A133)

$$J_{sp}^{agg} = \max[\tau_{diat}^{agg,min} P'_{diat}, \min[\tau_{diat}^{agg,max} P'_{diat}, \lambda_{diat}^{mort2} P'^{2}_{diat}]]$$
(A134)

$$J_{diaz}^{agg} = 0.0 \tag{A135}$$

N-cycle rates

$$J^{ammox} = \begin{cases} \tau^{ammox} N_{nh4}, & \text{where } \text{PAR}_B(k) < \text{PAR}_{max}^{nitrif} \text{ and } O_2 > O_2^{min} \\ 0, & \text{elsewhere} \end{cases}$$
(A136)

$$J^{nit} = \begin{cases} \tau^{nit} N_{no2}, & \text{where } \text{PAR}_B(k) < \text{PAR}_{max}^{nitrif} \text{ and } O_2 > O_2^{min} \\ 0, & \text{elsewhere} \end{cases}$$
(A137)

$$J_{o2}^{cons} = (J_{poc}^{remin} + J_{DOC,dic} + J_{zoo,dic}^{l} + J_{sp,dic}^{l} + J_{sp,dic}^{grz} + J_{diat,dic}^{l} + J_{diat,dic}^{grz} + J_{diat,dic}^{grz} + J_{diaz,dic}^{grz} + J_{diaz,dic}^{poc} + J_{diaz,dic}^{poc} + J_{remin}^{poc})/Q_{C:O} + 1.5 J^{ammox} + 0.5 J^{nit}$$
(A138)

$$J_{no3,no2}^{denit} = \begin{cases} Q_{N:C}^{denit} J_{poc}^{remin}, & \text{where } O_2 \le O_2^{min} \\ 0, & \text{elsewhere} \end{cases}$$
(A139)

$$J_{no2,n2o}^{denit} = f_{n2o}^{denit} J_{no3,no2}^{denit}$$
(A140)

$$J_{n2o,n2}^{denit} = \tau_{n2o}^{cons} N_{n2o}$$
(A141)

$$J_{sed}^{denit} = \begin{cases} a_0^{bohl} + a_1^{bohl} (a_2^{bohl})^{O_2 - N_{no3}} \frac{\Phi_{poc}^{out}}{\Delta z}, & \text{where } z = z_{bottom} \\ 0, & \text{elsewhere} \end{cases}$$
(A142)

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