Pico-phytoplankton contribute half of global marine carbon export

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Abstract

Earth System Models generally predict increasing upper ocean stratification from 21st century planetary warming, which will cause a decrease in the vertical nutrient flux resulting in declining marine net primary productivity (NPP) and carbon export fluxes. Recent advances in quantifying marine ecosystem carbon to nutrient stoichiometry have identified large latitudinal and biome variability, with low-latitude oligotrophic systems harboring pico-sized phytoplankton exhibiting large phosphorus to carbon cellular plasticity. Climate forced changes in nutrient flux stoichiometry and phytoplankton community composition is thus likely to alter the ocean’s biogeochemical response and feedback with the carbon-climate system. We have added three pico-phytoplankton functional types within the Biogeochemical Elemental Cycling component of the Community Earth System Model while incorporating variable cellular phosphorus to carbon stoichiometry for all represented phytoplankton types. The model simulates Prochlorococcus and Synechococcus populations that dominate the productivity and sinking carbon export of the tropical and subtropical ocean, and pico-eukaryote populations that contribute significantly to productivity and export within the subtropical to mid-latitude transition zone, contributing a combined 50 – 70% of these fluxes. Pico-phytoplankton cellular stoichiometry and resulting sinking export patterns inversely track the distribution of surface phosphate, with the western subtropical regions of each basin supporting the most P-poor stoichiometries. Collectively, pico-phytoplankton contribute ~58% of global NPP and ~46% of global particulate organic carbon export below 100 meters. Subtropical gyre recirculation regions along the poleward flanks of surface western boundary currents are identified as regional hotspots of enhanced carbon export exhibiting C-rich/P-poor stoichiometry, preferentially inhabited by pico-eukaryotes and diatoms.
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**Key Points:**

- Explicitly simulated *Prochlorococcus*, *Synechococcus*, and pico-eukaryotes contribute ~60\% of global net primary productivity
- Pico-phytoplankton contribute half of global marine sinking particle export production
- Carbon export is elevated poleward of western boundary currents, dominated by pico-eukaryotes and diatoms with elevated C:P stoichiometry
Abstract

Earth System Models generally predict increasing upper ocean stratification from 21st century planetary warming, which will cause a decrease in the vertical nutrient flux resulting in declining marine net primary productivity (NPP) and carbon export fluxes. Recent advances in quantifying marine ecosystem carbon to nutrient stoichiometry have identified large latitudinal and biome variability, with low-latitude oligotrophic systems harboring pico-sized phytoplankton exhibiting large phosphorus to carbon cellular plasticity. Climate forced changes in nutrient flux stoichiometry and phytoplankton community composition is thus likely to alter the ocean’s biogeochemical response and feedback with the carbon-climate system. We have added three pico-phytoplankton functional types within the Biogeochemical Elemental Cycling component of the Community Earth System Model while incorporating variable cellular phosphorus to carbon stoichiometry for all represented phytoplankton types. The model simulates Prochlorococcus and Synechococcus populations that dominate the productivity and sinking carbon export of the tropical and subtropical ocean, and pico-eukaryote populations that contribute significantly to productivity and export within the subtropical to mid-latitude transition zone, contributing a combined 50 – 70% of these fluxes. Pico-phytoplankton cellular stoichiometry and resulting sinking export patterns inversely track the distribution of surface phosphate, with the western subtropical regions of each basin supporting the most P-poor stoichiometries. Collectively, pico-phytoplankton contribute ~58% of global NPP and ~46% of global particulate organic carbon export below 100 meters. Subtropical gyre recirculation regions along the poleward flanks of surface western boundary currents are identified as regional hotspots of enhanced carbon export exhibiting C-rich/P-poor stoichiometry, preferentially inhabited by pico-eukaryotes and diatoms.
1. **Introduction**

Ocean biogeochemical (OBGC) models operating within larger Earth System Models (ESMs) have been developed for the purpose of quantifying the feedbacks between ocean biology, chemistry, and the climate system as well as to provide predictions of ocean biogeochemistry fluxes and tracer concentrations decades to centuries into the future (Le Quéré et al., 2005; Friedlingstein et al., 2006; Bopp et al., 2013; Cabré et al., 2015; Moore et al., 2013; 2018). ESM output from the Coupled Model Intercomparison Project Phase 5 (CMIP5) and Phase 6 (CMIP6) has predicted ocean acidification, deoxygenation, and decreasing primary productivity during the 21st century (Bopp et al., 2013; Cabré et al., 2015, Moore et al., 2013; Kwiatkowski et al., 2020) and beyond (Moore et al., 2018) under a business-as-usual scenario, primarily as a result of changes in ocean circulation and stratification in response to warming ocean temperatures. Enhanced upper ocean stratification leads to a reduction in vertical nutrient flux to the sunlight surface layer, decreasing net primary productivity (NPP) by phytoplankton, with a concomitant reduction in carbon export to the ocean interior (Cabré et al., 2015; Wilson et al., 2022), resulting in a positive feedback with the climate system as ocean carbon uptake is diminished with sustained warming. These declines are nearly universal among CMIP5 models, most of which feature fixed C:N:P:-O$_2$ stoichiometry (i.e. the Redfield ratio; Redfield et al., 1934; 1958) to describe phytoplankton nutrient uptake, production of organic matter, and its subsequent remineralization (Bopp et al., 2013). Further, declines in key carbon cycle features persist in the latest generation CMIP6 models despite an increasingly prevalent switch to variable nutrient to carbon quotas within the marine biogeochemistry modules (Séférian et al., 2020; Wilson et al., 2022).
Common to nearly all OBGC models is the representation of one or more ‘groups’ of phytoplankton that consume inorganic nutrients, resulting in photosynthesis, net primary production, and production of autotrophic biomass. Autotrophy is most often distributed among ≥ 2 phytoplankton functional types (PFTs), that are chosen to represent differing growth strategies, e.g., slower-growing small pico- and nano-phytoplankton vs. fast-growing large diatoms, to capture biogeographic latitudinal trends in nutrient uptake rates, cell size, and export efficiency. Phytoplankton growth experiences top-down control due to grazing by one or more zooplankton groups, also resulting in the production of particulate and dissolved organic detrital pools. Bottom-up controls on phytoplankton growth include multiple nutrient limitations, light limitation, and optimum temperature effects. Generally, nutrient uptake affinity is inversely correlated with cell size with smaller PFTs better able to acquire nutrients at low in situ concentrations. Each PFT acquires nutrients to satisfy their particular cellular N:C, P:C, and/or Fe:C quota, depending on the simulated inorganic nutrient fields. These cellular nutrient quotas are invariant for fixed stoichiometry models (i.e., at Redfield proportions; Moore et al., 2004; Dunne et al., 2010; Ilyina et al., 2013; Aumont et al., 2015) or can be variable, dependent on in situ environmental parameters such as nutrient concentration (Galbraith & Martiny, 2015) or optimal resource allocation theory describing phytoplankton cellular functions (Klausmeier et al., 2004; Dunne, 2013; Kwiatkowski et al., 2018). Many culture and field observations have documented variability in cellular nutrient quotas that vary both with ambient nutrient concentrations (Tanioka et al., 2022; Karl et al., 2001; Rhee, 1978) and/or across PFTs (Martiny et al., 2013a; Baer et al., 2017; Geider & La Roche, 2002; Quigg et al., 2003). Additionally, with the emergence of flow cytometry for the study of marine microbes (Lomas et al., 2011), a large literature of field and culture based studies has accumulated in recent decades describing the
physiology, biogeography, and phylogeny of marine pico-phytoplankton, more specifically the cyanobacterial lineages *Prochlorococcus* and *Synechococcus* and pico-eukaryotic phytoplankton, which now allows for their representation in numerical marine ecosystem models (e.g. Moore et al., 1998; Partensky et al., 1998; DuRand et al., 2001; Martiny et al., 2009; Buitenhuis et al., 2012; Pasulka et al., 2013; Flombaum et al., 2013; 2020; Sohm et al., 2016; Baer et al., 2017).

Given the need to balance model complexity against limiting computational resources, representation of additional pico-phytoplankton groups is perhaps of first order importance with respect to introduction of variable elemental stoichiometry within OBGC models, since pico-phytoplankton are known to exhibit the largest carbon to nutrient cellular plasticity (Quigg et al., 2003; Klausmeier et al., 2004; Martiny et al., 2013a; Baer et al., 2017). In light of these advances, representing the pico-phytoplankton and variable carbon to nutrient content of all phytoplankton in OBGC models has become a pressing goal to better capture the ocean ecosystem response to the changing 21st century climate and its associated feedbacks with the climate system.

Many recent studies have identified and quantified the non-Redfield, variable carbon to nutrient stoichiometry that permeates most ocean biogeochemical fluxes and tracer pools including dissolved nutrients (Weber & Deutsch, 2010), biomass (Tanioka & Matsumoto, 2020), particulate organic matter (Martiny et al., 2013a), dissolved organic matter (Letscher & Moore, 2015), export production (Teng et al., 2014), and organic matter remineralization (DeVries & Deutsch, 2014). The emergent pattern from these studies is one of carbon-rich (nutrient-poor) stoichiometry that describes biogeochemical functioning in the lower latitude, nutrient-poor biomes of the subtropics, while the nutrient-replete upwelling and subpolar biomes exhibit stoichiometries at or slightly below Redfield carbon to nutrient proportions. Two mechanisms
have been suggested to contribute to the observed biogeography of marine ecosystem stoichiometry including phosphorus and/or nitrogen plasticity by the extant phytoplankton community (e.g. Galbraith & Martiny, 2015; Van Mooy et al., 2009) as well as shifts in phytoplankton community composition across biomes with relatively nutrient-deficient cyanobacteria dominating the subtropics and more nutrient-rich diatoms and other larger phytoplankton dominating elsewhere (e.g. Arrigo et al., 1999; Weber & Deutsch, 2010; Sharoni & Halevy, 2020). Cell-sorted field populations of cyanobacteria (*Prochlorococcus* and *Synechococcus*) and pico-eukaryotes from the Sargasso Sea, eastern Pacific, and Indian Ocean suggest both mechanisms may be at play as cyanobacteria exhibited lower cellular P content as compared to eukaryotes, while all groups exhibited cellular P plasticity in response to latitudinal gradients in inorganic phosphorus availability, albeit at varying degrees across taxonomic groups (Martiny et al., 2013a; Baer et al., 2017; Lomas et al., 2021).

Does representation of variable carbon to nutrient stoichiometry within OBGC models potentially buffer the well-documented positive ocean carbon cycle feedback to future climate warming? A number of ESM’s that have included representation of variable carbon to phosphorus stoichiometry within the ocean ecosystem component generally predict a more resilient response for future marine NPP and carbon export fluxes, with modest ~0 – <5% declines by year 2100 (Bopp et al., 2013; Cabré et al., 2015; Tanioka & Matsumoto; 2017; Kwiatkowski et al., 2018; Buchanan et al., 2018; Matsumoto et al., 2020; Kwon et al., 2022).

Inclusion of variable phosphorus to carbon stoichiometry within the marine ecosystem component permits flexibility in phytoplankton phosphorus quotas. Such frugal use of nutrients allows phytoplankton to maintain similar growth rates under decreasing P flux as well as a shift towards increasing biomass C:P that partially compensates for warming induced changes to
ocean circulation and stratification. Galbraith and Martiny (2015) demonstrated with a simple box model that changes to the preformed phosphate concentration on the order of 100 nM within low latitude surface waters can produce up to 20 ppm shifts in atmospheric CO$_2$ concentrations, assuming P is the primary growth limiting nutrient. An OBGC modeling study incorporating representation of variable marine elemental stoichiometry found that adding this sensitivity buffered against a predicted decrease in global carbon export production on the order of ~3% for the 21st century (Tanioka & Matsumoto, 2017). Similarly, results from the recent inclusion of this process within the Pelagic Interactions Scheme for Carbon and Ecosystem Studies Quota (PISCES-QUOTA) model, suggest its effects on 21st century ocean carbon uptake are modest with up to a 3.5% underestimation as compared to fixed stoichiometry ESMs (Kwiatkowski et al., 2018). However, larger regional changes of up to 50% reductions or stimulations of marine NPP and carbon export were predicted which could have significant local to regional impacts on marine ecosystems and fisheries. The above studies have thus far considered adaptive nutrient quotas, while other phytoplankton adaptive strategies including altered physiological rates, trait-based evolution, and changes in grazing pressure could all potentially influence phytoplankton fitness and resilience with important controls on marine NPP and carbon export trends (Martiny et al., 2021).

As the next generation of OBGC models mature to better represent variable carbon to nutrient stoichiometry with the goal to study how this variability impacts the marine NPP and carbon export fluxes, an important missing element to this question is the role for changes in marine phytoplankton biodiversity to modulate the response. Most of the current generation of OBGC models represent the marine phytoplankton community with 1 – 3 PFTs, with the notable exception of the ECCO-Darwin OBGC model which simulates 35 up to 350 PFTs (e.g.,
Dutkiewicz et al., 2020; Juranek et al., 2020). This higher biodiversity model has been shown to more accurately represent the marine phytoplankton diversity regionally to globally, as gauged by such metrics as phytoplankton size distribution, biomass, size-fractionated NPP rates, and species richness. How does marine phytoplankton diversity and associated carbon to nutrient stoichiometric variability impact the regional to global patterns of carbon export? Based on current understanding, enhanced thermal stratification of the water column depresses the vertical nutrient input, favoring a shift to smaller sized pico-phytoplankton dominated by cyanobacteria possessing higher plasticity in cellular nutrient quotas. Thus, in a future warmer ocean with expanding stratification, the marine biodiversity response might be hypothesized to favor marine cyanobacteria such as Prochlorococcus and Synechococcus with elevated C:P cellular stoichiometry. While the shift to smaller phytoplankton may depress marine NPP rates, export of a portion of the resultant biomass with high C:P may partially buffer against reductions in marine carbon export and mitigate the expected positive feedback to warming by maintaining the strength of the ocean’s biological carbon pump. In support of this hypothesis, observations from the Bermuda Atlantic Time-series station in the Sargasso Sea document a recent decline in marine NPP due to enhanced stratification, without a concomitant decline in carbon export, explained by a shift in the phytoplankton community composition in favor of cyanobacteria with higher cellular C:P stoichiometry (Lomas et al., 2022).

This paper describes the addition of three new pico-phytoplankton functional types, Prochlorococcus, Synechococcus, and pico-eukaryotes, to the OBGC component of the Community Earth System Model, with the goal to better characterize the role for marine phytoplankton biodiversity and associated variable cellular carbon to phosphorus stoichiometry in controlling marine carbon export patterns. The implementation builds from recent advances in
this OBGC model to incorporate variable C:P stoichiometry within all pertinent tracers of the marine ecosystem (Letscher et al., 2015; Wang et al., 2019) including phytoplankton, dissolved organic matter, and sinking particulate organic matter, to now include representation of six PFTs: *Prochlorococcus*, *Synechococcus*, pico-eukaryotes, nano-phytoplankton (including implicit coccolithophores), diatoms, and diazotrophs.

2. Model and Validation Data

The model is the Biogeochemical Elemental Cycling (BEC) model, which is the biogeochemistry and ecosystem component operating within the ocean circulation model (POP2) of the Community Earth System Model (CESM) v.1.2.2 (http://www.cesm.ucar.edu/models/cesm1.2/). The model resolution employed here is nominally 3° x 3° horizontally with 60 vertical layers (gx3v7 grid) that increase in thickness with depth in the water column. The ocean circulation is the output of the Parallel Ocean Program 2 (POP2) ocean circulation model using the final thirty-year forcing from years 1980 – 2009 of the CORE-II forcing (Smith et al., 2010) and includes parameter value changes to the isopycnal diffusivity ($\kappa_{\text{isop}}$) to partially account for the transport of unresolved equatorial jets (Getzlaff & Dietze, 2013).

The BEC tracks the cycling of key biogeochemical tracers (e.g., C, N, P, Fe, and O$_2$) amongst dissolved inorganic, dissolved organic, sinking particulate, and four plankton biomass pools (Moore et al., 2004). Net primary production in the model is partitioned amongst three PFTs–small phytoplankton (with implicit calcifiers), diatoms, and diazotrophs, which differ in their relative competitiveness for limiting nutrients (N, P, Si, and Fe). In addition to bottom-up nutrient and light control, phytoplankton mortality is controlled top-down by grazing from one zooplankton group. Version 1.2.2 of the BEC contains significant improvements to the cycling of
marine dissolved organic matter (Letscher et al., 2015). More recent model development for
BEC v1.2.2 has incorporated variable phosphorus quotas for the three phytoplankton types
(Wang et al., 2019) that depend on *in situ* simulated inorganic phosphate concentration following
the linear relationship in Galbraith & Martiny (2015) but with an imposed minimum (110:1) and
maximum (210:1) C:P value for all PFTs. This current work describes the addition of three new
pico-phytoplankton groups with variable carbon to nutrient (P and Fe) stoichiometry to the BEC
model, building from the code base for the variable 3 PFT phosphorus quota model of Wang et
al., (2019). The ‘small phytoplankton’ group from the 3 PFT BEC model which jointly
represented pico- and nano-sized phytoplankton has been reparameterized to more closely
represent larger nano-phytoplankton. The newly named nano-phytoplankton along with the
diatom and diazotroph groups are updated to represent variable C:P and C:Fe stoichiometry. The
model retains one zooplankton group with Redfieldian stoichiometry. The marine ecosystem in
the BEC model is thus represented by 6 phytoplankton and 1 zooplankton groups, or 6p1z.

Output of the pico-phytoplankton BEC model is validated for the simulation of
chlorophyll a and NPP against the Sea-Viewing Wide Field-of-View Sensor (SeaWiFS) satellite
mission climatology products (1997 – 2010) related to chlorophyll a concentration (NASA
GSFC, 2018) and a model of carbon-based net primary productivity estimated from SeaWiFS
data products (Behrenfeld et al., 2005; Westberry et al., 2008). Simulated nutrient concentrations
from the pico-phytoplankton BEC model are compared against the World Ocean Atlas 2013
climatologies of nitrate, phosphate, and silicate (Garcia et al., 2013) and a global compilation of
dissolved Fe concentrations (Tagliabue et al., 2012). Simulated pico-phytoplankton biomass are
validated against the data assimilative niche modeling of Flombaum et al. (2013; 2020) that
predicts field observed *Prochlorococcus, Synechococcus* from inputs of temperature and
irradiance (Flombaum et al., 2013) and pico-eukaryote biomass from inputs of temperature, irradiance, and dissolved nitrate (Flombaum et al., 2020). Simulated pico-phytoplankton C:P stoichiometry and biomass are compared against observations of populations from the North Atlantic, eastern Pacific, and eastern Indian Oceans of Prochlorococcus, Synechococcus, and pico-eukaryotes measured for C and P content (Baer et al., 2017; Lomas et al., 2021). Simulated total phytoplankton biomass C:P stoichiometry is compared against a global compilation of bulk particulate organic matter C, N, and P content (Martiny et al., 2014).

3. Methods

The addition of three pico-phytoplankton groups with variable carbon to nutrient stoichiometry introduces 12 additional tracers to the BEC model code: biomass carbon (C), chlorophyll (Chl), iron (Fe), and phosphorus (P) for each of the 3 pico-phytoplankton groups, termed ‘Prochlorococcus’ (pro), ‘Synechococcus’ (syn), and ‘Pico-eukaryotes’ (peuk). The ‘small phytoplankton’ group from BEC v1.2.2 and earlier versions has been renamed ‘nano-phytoplankton’ (nano) and retains the implicit calcifiers. Nano-phytoplankton as well as the diatoms (diat) and diazotroph (diaz) phytoplankton groups carry the same four biomass tracers (C, Chl, Fe, P) as the newly added pico-phytoplankton, with the addition of biomass silicon for diatoms. Variable biomass nitrogen (N) content is not considered at this time and phytoplankton biomass N:C is fixed for all six phytoplankton groups at a value of 0.1333 (C:N = 7.5) near the mean of open ocean suspended particles (Martiny et al., 2013b) and nutrient replete versus deplete cells across a range of phyla (Geider & La Roche, 2002).

Growth and mortality of each phytoplankton type is modeled with more than two dozen parameter values specific to each group including half saturation constants for nutrient uptake,
photosynthesis vs irradiance curves, Fe and P quotas, temperature thresholds for growth, contributions to detrital pools, and zooplankton grazing rates among others. Table 1 lists these parameter values for each phytoplankton group including the relevant literature sources. Where literature sources are lacking to inform the value for a particular parameter, the principle of allometric scaling is employed to assign values corresponding with cell size with the order of increasing cell size following: pro/syn/peuk < nano/diaz < diat.

**Table 1.** Description of phytoplankton parameters and values from the PICO simulation of the BEC model.

Phytoplankton iron uptake half saturation constants are prescribed with cyanobacteria having the lowest values (pro = 30 pM; syn+diaz = 40 pM), followed by pico-eukaryotes (43 pM), nano-eukaryotes (55 pM), and the large celled diatoms (60 pM) following the culture studies of Shire and Kustka, (2015) and Timmermans et al., (2005) and the field study of Hutchins et al., (2002) (Table 1). Phosphate half saturation constants are selected based on the
field study from the Sargasso Sea of Lomas et al., (2014), the culture study of Garcia et al., (2015), and the Amazon River plume ecosystem modeling work of Stukel et al., (2014) with kPO4 values increasing from a value of 8 nM for pro, 10 nM for syn, 15 nM for peuk, 25 nM for diaz, 40 nM for nano, and 70 nM for diatoms. Semilabile dissolved organic phosphorus (DOP) is bioavailable for phytoplankton to utilize as a P source when phosphate levels are low in the BEC model (Letscher et al., 2015) with the half saturation constants for its uptake selected based on several field studies from the Sargasso Sea and eastern North Pacific (Dyhrman & Ruttenberg, 2006; Sohm & Capone, 2006; Casey et al., 2009; Orchard et al., 2010). Following these studies, eukaryotes and diazotrophs are assigned the lowest half saturation values for DOP uptake ranging from 100 – 250 nM, with higher kDOP values ranging from 800 – 900 nM for the two cyanobacteria groups. Half saturation constants for ammonium are assigned following allometric scaling in the range 5 – 35 nM. Nitrate uptake half saturation constants are similarly assigned following allometric scaling in the range 150 – 350 nM, with exceptions for syn and peuk which are assigned the lowest values of 180 and 150 nM respectively, owing to their demonstrated increased affinity for nitrate relative to pro in the Sargasso Sea (e.g., Glover et al., 1988; Fawcett et al., 2013) and observed correlations between peuk abundance and in situ nitrate (Flombaum et al., 2020). Diazotroph kNH4 and kNO3 are set to the highest value across the 6 PFTs to reflect its reliance on internally fixed N for a significant but not exclusive fraction of its N quota (Knapp, 2012).

Phytoplankton iron quotas in the BEC model allow for flexibility in phytoplankton biomass Fe:C based on in situ dissolved iron concentrations, varying in the range of 3 µmol:mol to 180 µmol:mol (Table 1). Following the field study of Boyd et al., (2015), eukaryotes are assigned the lowest minimum iron quotas followed by diazotrophs and non-N-fixing
cyanobacteria (Prochlorococcus and Synechococcus). The diazotroph group is assigned the
highest maximum iron quotas based on the high iron requirement of the nitrogen-fixing enzyme
nitrogenase (Berman-Frank et al., 2007). Phytoplankton iron quotas are reduced linearly when in
situ dissolved Fe concentrations fall below a prescribed threshold (FeOpt), set to 1.75 nM for
diatoms, 1.6 nM for nano, 1.5 nM for diaz, 1.0 nM for peuk, and 0.8 nM for pro and syn
(Wisemann et al., under review).

Variable phosphorus quotas are implemented here for the six phytoplankton group BEC
model following the approach introduced in Wang et al. (2019) and similar to the variable Fe:C
approach described above. Variable phytoplankton biomass P:C is modeled as a function of in
situ dissolved phosphate concentration using the linear relation detailed
in Galbraith & Martiny
(2015) diagnosed from a global compilation of suspended particulate organic phosphorus and
carbon concentrations. Each PFT P:C quota (gQp) is computed from:

\[
gQp = \left(\frac{gQp_0 - gQp_{\text{min}}}{P_{\text{Opt}}}ight) \times PO4 + gQp_{\text{min}} \quad \text{PO4 < POpt} \quad \text{(Eq. 1)}
\]

\[
gQp = gQp_0 \quad \text{PO4 \geq POpt}
\]

where PO4 is the in situ dissolved inorganic phosphate concentration (µM) and gQp_0 and
gQp_min describe the minimum and maximum P:C quota (see Table 1). Equation 1 linearly
decreases the phytoplankton P:C quota (increases its C:P ratio) below a prescribed threshold
(POpt; range 0.5 – 1.0 µM across PFTs), from its initial maximum value, currently prescribed at
gQp_0 = 0.01111; C:P = 90:1 for the eukaryotes (peuk+nano+diat), gQp_0 = 0.01042; C:P =
96:1 for the cyanobacteria, and gQp_0 = 0.006667; C:P = 150:1 for diazotrophs, towards a
minimum P:C quota that varies for each PFT (Table 1). This formulation is similar in approach
to that presented in Galbraith & Martiny (2015). Here, field observations of surface ocean
particulate organic carbon and phosphorus (of which autotrophic biomass is a significant
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contributor) were found to fit a linear function of *in situ* phosphate concentration. The formulation presented here prescribes differing P:C versus PO₄ slopes and y-intercepts for each PFT set by the choice of gQp_min. Field-based estimates of cell-sorted biomass C:P and N:P for *Prochlorococcus*, *Synechococcus*, and pico-eukaryotes from the Sargasso Sea (Martiny et al., 2013a) and elsewhere (Baer et al., 2017; Lomas et al., 2021) are used to assign values of gQp_min with a maximum C:P of 300:1 for diazotrophs, a C:P of 270:1 for *Prochlorococcus* and 192:1 for *Synechococcus*, and a maximum C:P of 120:1 for peuk, nano, and diat (retained from the variable C:P BEC model; Wang et al., 2019).

Based on the phytoplankton niche modeling of Flombaum et al. (2013; 2020), each phytoplankton group in the BEC is assigned a temperature threshold below which photosynthesis rates and thus growth rates are depressed, with *Prochlorococcus* and diazotroph niches favoring warmer temperatures above 13°C and 18°C, respectively. *Synechococcus* and pico-eukaryote growth decreases as temperatures fall below 4°C and -1.5°C, respectively, while nano-phytoplankton and diatom growth rates are not depressed as a function of temperature (temp_thres value set to -10°C in Table 1).

Physiological response to light parameters describing the pico-phytoplankton are assigned based on the culture study of Stawiarski et al. (2017) in which two strains of *Prochlorococcus* (HL RCC 296, LL RCC 162), one *Synechococcus* strain (RCC 30), and four pico-eukaryote species *Triparma eleuthera* (RCC 212), *Micromonas pusilla* (RCC 1677), *Picochlorum* sp. (TCC 289), and *Nannochloropsis granulata* (RCC 438) from the Roscoff culture collection (Vaulot et al., 2004) were studied. Following this study, cyanobacteria are assigned slower maximum growth rates compared with eukaryotes, in the range 1.9 – 4 d⁻¹ (based on a 24-hour light period). Stawiarski et al. (2017) found that light affinity is inversely related to
the maximum growth rate. *Prochlorococcus* and *Synechococcus* are assigned the highest light
affinities, as encoded within the ‘alphaPI’ model parameter which describes the initial slope of
the photosynthesis-irradiance curve, with decreasing affinities for the faster growing eukaryotes
(Table 1). The maximum Chl:N, the ‘thetaN_max’ model parameter, is assigned inversely
proportional to maximum growth rate with the exception of *Synechococcus* which exhibits lower
Chl:C in line with the results of Stawiarski et al. (2017).

Grazing on each of the six phytoplankton groups in the updated BEC model is carried out
by one adaptive zooplankton group. Grazing rates are largely unconstrained by field data. In the
BEC model, maximum grazing rates, ‘z_umax_0’ parameter, are adjusted to achieve satisfactory
biomass distributions and C:P stoichiometry for each phytoplankton group as validated against
phytoplankton niche modeling and flow-sorted biomass stoichiometry distributions. Variable
fractions of phytoplankton biomass are routed to zooplankton losses to detritus (e.g., fecal
pellets) and particulate organic matter pools resulting from zooplankton grazing (Table 1). Each
PFT’s contribution to the sinking export flux following grazing varies from a minimum of 3%
for diazotrophs to a maximum of 40% for diatoms (‘graze_poc’; Table 1) informed in part by
pigment biomarker analyses of the sediment trap flux in the Sargasso Sea (Lomas & Moran,
2011). Zooplankton contribute varying proportions of grazed phytoplankton biomass to detritus
(POC) based on the PFT grazed, ranging from a minimum of 20% for nano-phytoplankton and
diatoms to a maximum of 36% when grazing diazotrophs (‘f_zoo_detr’; Table 1). Grazing of
phytoplankton biomass also contributes 6% to the dissolved organic matter pool following the
inverse model diagnosed rate from our previous work (Letscher et al., 2015). In the BEC model,
combinations of two or more phytoplankton types can be grouped together in ‘graze’ classes,
for which zooplankton grazing rates are computed jointly based on the total amount of biomass
in each simulated in situ ‘grazee’ class. Experimentation was carried out to decide which
combinations of phytoplankton groups in each ‘grazee’ class achieved distributions of
phytoplankton biomass, NPP, and dissolved nutrients that best matched observations. A model
configuration in which all six PFTs were ungrouped, i.e., parameter grazee_ind was set equal to
auto_ind, the unique index of each of the six PFTs, was chosen.

BEC simulations were carried out for sixty model years from initial conditions, allowing
the upper ocean ecosystem to achieve tracer equilibrium. After selection of the set of parameter
values that best matched observational constraints, a longer 300 model year simulation was
performed, repeating the 30-year (1980 – 2009) CORE-II forcing for a total of ten repetitions.
The annually averaged output from the final twenty years of this simulation were averaged and is
presented in all results to follow. Reported flux variability represents ±1 S.D. of the mean from
the twenty years of output. The BEC simulation containing the additional pico-phytoplankton
groups for a total of 6 PFTs is termed PICO. In Section 4, this simulation is compared against
two similarly forced previous versions of the BEC model: a variable C:P stoichiometry version
with 3 PFTs herein termed VAR-3 (Wang et al., 2019) and a 3 PFT version with Redfield
stoichiometry governing phytoplankton dynamics herein termed RED-3 (Letscher et al., 2015).

4. Results

Chlorophyll a & NPP—Sea surface chlorophyll a (chl a) concentrations [mg m\(^{-3}\)] from the
PICO simulation and the SeaWiFS satellite climatology are shown in Figure 1a-b. BEC
simulated chl a captures the overall pattern of lowest concentrations within the subtropical ocean
gyres, highest concentrations within coastal and subpolar zones, and intermediate concentrations
in upwelling regions and subtropical to mid-latitude transitions zones (Fig. 1). The PICO
simulation exhibits an overall global negative mean bias in chl a (~6%; Table 2) relative to the SeaWiFS climatology, reducing by a factor of ~3 compared with the 3 PFT variable C:P stoichiometry BEC model (VAR-3) and an earlier version containing 3 PFTs and Redfield stoichiometry (RED-3) (Table 2).

<table>
<thead>
<tr>
<th></th>
<th>PICO</th>
<th>VAR-3</th>
<th>RED-3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chl a</td>
<td>0.087</td>
<td>-6%</td>
<td>0.218</td>
</tr>
<tr>
<td>CbPM NPP</td>
<td>0.241</td>
<td>+3%</td>
<td>0.108</td>
</tr>
<tr>
<td>[NO₃⁻]</td>
<td>0.938</td>
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<tr>
<td>[PO₄³⁻]</td>
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<tr>
<td>[SiO₄⁴⁻]</td>
<td>0.935</td>
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</tr>
<tr>
<td>[dFe]</td>
<td>0.398</td>
<td>0.18 nM</td>
<td>0.332</td>
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**Table 2.** Comparison metrics for PICO, VAR-3, and RED-3 simulated chlorophyll a and NPP versus satellite (SeaWiFS) climatology estimates using the CbPM model. Metrics for dissolved nutrients from the same three simulations are compared versus the World Ocean Atlas 2013 (NO₃⁻, PO₄³⁻, SiO₄⁴⁻) and Tagliabue et al. (2012) Fe data compilation.

Vertically integrated annual net primary productivity (NPP) [g C m⁻² yr⁻¹] from the PICO simulation is provided along with the Carbon-based Productivity Model estimate of NPP from the SeaWiFS climatology (Westberry et al., 2008) (Figure 1c-d). The globally integrated annual NPP from PICO is 51.4 ± 0.75 Pg C yr⁻¹, within the range of satellite-based estimates – 60-67 Pg C yr⁻¹ (Behrenfeld et al., 2005), 52 Pg C yr⁻¹ (Westberry et al., 2008; Silsbe et al., 2016), 58 ± 7 Pg C yr⁻¹ (Buitenhuis et al., 2013). Highest rates of annual NPP are simulated in the equatorial Pacific, Arabian Sea, and the South Pacific and North Atlantic eastern boundary upwelling zones, reaching rates of ~400 – 600 g C m⁻² yr⁻¹ in the PICO simulation, which is approximately
25 – 75% greater than those estimated by the CbPM. The lowest rates of NPP simulated by PICO are found within the Arctic Ocean, which are approximately one-quarter of those estimated by CbPM. The PICO simulation also simulates higher rates of NPP as compared to CbPM within the mid-latitudes of both hemispheres, especially near the western boundary of each ocean basin. Overall, the PICO simulation exhibits a small positive mean bias of +3% in estimated global annual NPP rates when comparing binned 3ºx3º pixels with the CbPM SeaWiFS climatology (Table 2).

**Figure 1.** PICO simulated 20-yr averaged sea surface chlorophyll a [mg m⁻³] (a), climatological average sea surface chlorophyll a from the SeaWiFS satellite during 1997-2010 (b), PICO simulated 20-yr averaged NPP [g C m⁻² yr⁻¹] (c), and from the CbPM model using SeaWiFS satellite data from the same era (d) (Westberry et al., 2008).

**PFT contributions to spatial NPP patterns** – The fractional contribution of each of the six PFTs to simulated NPP rates within the PICO simulation is shown in Figure 2. The majority of
NPP within the five subtropical gyre regions is contributed by *Prochlorococcus* (fraction of NPP ~ 0.50 – 0.90), with a general western intensification (Fig. 2a). *Synechococcus* NPP is similarly elevated within the subtropics, albeit at lower fractional contributions in the range ~0.06 – 0.12 (Fig. 2b). *Prochlorococcus* and *Synechococcus* are mostly absent from the polar oceans due to their imposed temperature threshold of 13°C and 4°C for positive growth rates (Flombaum et al., 2013; Table 1). Pico-eukaryotes dominate NPP (fraction of NPP > 0.80) in many of the eastern boundary and mid-latitude regions of the world ocean in PICO (Fig. 2c). Elsewhere, pico-eukaryote contribution to simulated NPP is much reduced, in the range ~0.02 – 0.10. Nano-phytoplankton contribute variably to NPP rates within a range of 0.02 – 0.20 throughout the world ocean in PICO (Fig. 2d). Lowest simulated contributions are found in the eastern and equatorial upwelling regions; highest contributions are located in the polar oceans. Diatoms dominate simulated NPP rates (fraction > 0.80) within the subpolar and polar oceans as well as the equatorial Pacific (Fig. 2e), with lowest fractional contributions to NPP (~0.01 – 0.05) located within the subtropical gyres. Diazotrophs contribute < 0.07 to simulated NPP rates and are absent from the subpolar and polar oceans, due to the imposed minimum temperature of 18°C for positive growth rates (Table 1). Highest diazotroph contributions to NPP rates are found within the western tropical to subtropical North and South Pacific Oceans (~0.04 – 0.07) (Fig. 2f), similar to other OBGC model simulations that incorporate variable nutrient stoichiometry (e.g., Wang et al., 2019).
Figure 2. Fraction of vertically integrated, annual marine NPP attributable to each PFT from the PICO simulation. (a) *Prochlorococcus*, (b) *Synechococcus*, (c) pico-eukaryotes, (d) nano-phytoplankton, (e) diatoms, (f) diazotrophs. Note change in range in panels a,c,e from panels b,d, and panel f.

**Nutrients** – Upper ocean (0 – 500 m) simulated dissolved nutrient concentrations from PICO have been compared against the World Ocean Atlas (WOA) 2013 climatology (Garcia et al., 2013) for nitrate (NO$_3^-$), phosphate (PO$_4^{3-}$), and silicate (SiO$_4^{4-}$) and a global compilation of dissolved Fe (dFe) measurements (Tagliabue et al., 2012). Table 2 provides the r-correlation parameter and mean bias (µM) from a regression of log-transformed model-observation pairs. Dissolved NO$_3^-$, PO$_4^{3-}$, and SiO$_4^{4-}$ are equally well correlated with WOA nutrients, exhibiting r-correlation coefficients of ~0.94 and mean biases of +0.01 µM NO$_3^-$, +0.02 µM PO$_4^{3-}$, and +0.01 µM SiO$_4^{4-}$. PICO simulated dFe is the most weakly correlated nutrient with observational data, r = 0.40, with a mean bias of +0.18 nM. The PICO upper ocean dissolved nutrient mean biases are similar to those of the other variable C:P stoichiometry simulation, VAR-3, and the Redfield simulation, RED-3, with a notable improvement to the mean bias of SiO$_4^{4-}$ from VAR-3 (Table 2).
Biomass – PICO simulated biomass ($\mu$g C L$^{-1}$) is compared against the data assimilative niche model output for Prochlorococcus and Synechococcus (Flombaum et al., 2013) and pico-eukaryotes (Flombaum et al., 2020) binned by 3°x3° pixels. Simulated pico-phytoplankton biomass correlation coefficients with the niche model output range from 0.485 (pico-eukaryotes) to 0.637 (Prochlorococcus) (Table 3). However, significant negative biases are present within PICO simulated Prochlorococcus biomass ($-56\%$), Synechococcus biomass ($-85\%$), and pico-eukaryotes biomass ($-76\%$) (Table 3). PICO pico-phytoplankton biomass has also been compared against a globally compiled field-based dataset of flow-sorted populations (Baer et al., 2017; Lomas et al., 2021), yielding metrics of: Prochlorococcus biomass ($+76\%$, $r = 0.199$), Synechococcus biomass ($-72\%$, $r = 0.245$), and pico-eukaryotes biomass ($-72\%$, $r = 0.142$) (Table 3).

<table>
<thead>
<tr>
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<th>r-correlation</th>
<th>mean bias</th>
<th>n</th>
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<tbody>
<tr>
<td>Pro C niche</td>
<td>0.637</td>
<td>-56%</td>
<td>24,447</td>
</tr>
<tr>
<td>Syn C niche</td>
<td>0.459</td>
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<tr>
<td>Peuk C niche</td>
<td>0.485</td>
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<td>39,244</td>
</tr>
<tr>
<td>Pro C:P</td>
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</tr>
<tr>
<td>Syn C:P</td>
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<tr>
<td>Peuk C:P</td>
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</tr>
<tr>
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<td>+20%</td>
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</tr>
<tr>
<td>Pro C</td>
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<td>+76%</td>
<td>2188</td>
</tr>
<tr>
<td>Syn C</td>
<td>0.245</td>
<td>-72%</td>
<td>4120</td>
</tr>
<tr>
<td>Peuk C</td>
<td>0.142</td>
<td>-72%</td>
<td>4158</td>
</tr>
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Table 3. Comparison metrics for PICO simulated pico-phytoplankton biomass versus the data assimilative niche models of Flombaum et al. (2013; 2020) (Pro C niche, Syn C niche, Peuk C niche); versus pico-phytoplankton biomass C:P stoichiometry (Baer et al., 2017; Lomas et al., 2021) (Pro C:P, Syn C:P, Peuk C:P) and biomass C (Pro C, Syn C, Peuk C); and versus bulk particulate organic matter C:P stoichiometry (Martiny et al., 2014) (POC:POP).
Phytoplankton Stoichiometry – Euphotic zone-averaged PFT C:P stoichiometry from the PICO simulation is shown in Figure 3. Prochlorococcus (Fig. 3a) and diazotrophs (Fig. 3f) exhibit the greatest range in cellular C:P stoichiometry, with the minimum stoichiometry found within the eastern equatorial Pacific Ocean, C:P = ~90 for pro; C:P = ~150 for diaz, increasing to maximum cellular C:P = ~250 – 300 within the North and South Atlantic subtropical gyres. Elevated C:P stoichiometry for Prochlorococcus and diazotrophs is also found within the other three subtropical gyre regions, with typical values in the range ~180 – 300. Synechococcus (Fig. 3b), pico-eukaryotes (Fig. 3c), nano-phytoplankton (Fig. 3d), and diatoms (Fig. 3e) all exhibit similar geographical variability in cellular C:P stoichiometry to Prochlorococcus and diazotrophs, with elevated values typical of the five subtropical ocean gyre regions, syn C:P = ~150 – 210; peuk + nano + diat C:P = ~110 – 125. Phytoplankton C:P stoichiometry is ~25 – 80% more elevated within the western half of each subtropical gyre relative to the eastern half.

Figure 3. Euphotic zone averaged cellular C:P stoichiometry for each PFT from the PICO simulation. (a) Prochlorococcus, (b) Synechococcus, (c) pico-eukaryotes, (d) nano-phytoplankton, (e) diatoms, (f) diazotrophs.
PICO simulated *Prochlorococcus*, *Synechococcus*, and pico-eukaryotes cellular C:P stoichiometry has been compared against a subset of the cell-sorted population dataset for which biomass P was measured (Baer et al., 2017; Lomas et al., 2021). PICO simulated *Prochlorococcus* C:P exhibits the smallest mean bias of ~6% below the average field-based estimate of C:P = 221 (Table 3). PICO simulated *Synechococcus* C:P has a negative mean bias of ~17% below the field estimate of 187. PICO simulated pico-eukaryotes exhibited a positive mean bias compared to field estimates at +11% above the field estimate of 103. PICO simulated total autotrophic biomass C:P stoichiometry is compared with a global compilation of surface ocean suspended organic matter C:P stoichiometry (Martiny et al., 2014), exhibiting a positive mean bias of +20% above the field-based estimate of POC : POP = 171 : 1 (Table 3).

*Sinking carbon export* – The sinking particulate organic carbon flux through the 100 meter depth horizon ($\text{POC}_{\text{exp,100m}}$) from the PICO simulation is plotted in Figure 4a. $\text{POC}_{\text{exp,100m}}$ exhibits a similar geographical distribution as simulated NPP (Fig. 1c), with maximum rates of ~90 – 120 g C m$^{-2}$ yr$^{-1}$ found within the eastern equatorial Pacific, Peruvian, and Arabian Sea upwelling regions as well as the SW Atlantic Ocean and minimum rates, < 20 g C m$^{-2}$ yr$^{-1}$, within the five subtropical ocean gyres and their bordering tropical regions, the Southern Ocean, and the Arctic Ocean. Other regions exhibiting elevated $\text{POC}_{\text{exp,100m}}$ include the eastern boundary upwelling zones and the subtropical to mid-latitude transition zones with rates in the range ~40 – 70 g C m$^{-2}$ yr$^{-1}$. The pico-phytoplankton contribution to $\text{POC}_{\text{exp,100m}}$ is greatest across the tropics, subtropics, and mid-latitudes (Fig. 4b), with a fractional contribution in the range ~0.50 – 0.70. In both hemispheres, pico-phytoplankton make their largest contribution to sinking POC export in the mid-latitudes (~40 – 50 °N and °S), approaching 65 – 70% of $\text{POC}_{\text{exp,100m}}$, with minima
(-0.10 – 0.30) estimated for the subpolar, polar, and equatorial Pacific Oceans. The globally integrated $POC_{exp,100m}$ flux below 100 m from PICO is $7.0 \pm 0.1$ Pg C yr$^{-1}$; within the range of observational-, satellite-, and model-based estimates of global marine export production, e.g., 4 – 14 Pg C yr$^{-1}$ (summarized in Emerson, 2014; Britten & Primeau, 2016; Siegel et al., 2014; Henson et al., 2011). It should be noted when comparing the PICO carbon export flux to other estimates from the literature that DOC export below 100 m contributes an additional ~2.1 Pg C yr$^{-1}$ of carbon export production in the BEC model (Letscher et al., 2015; Letscher & Moore, 2015) bringing estimated total organic carbon export below 100 m to ~9.1 Pg C yr$^{-1}$.

**Figure 4.** PICO simulated 20-yr averaged sinking particulate organic carbon export below 100 m $POC_{exp,100m}$ [g C m$^{-2}$ yr$^{-1}$] (a), the fraction of total $POC_{exp,100m}$ attributable to pico-phytoplankton (pro + syn + peuk) (b), the C:P stoichiometry of sinking organic particles C:P$_{exp}$ at 100 m (c), and the zonal mean C:P$_{exp}$ at 100 m (black line), C:P of cyanobacteria (pro + syn + diaz; red line), C:P of eukaryotic phytoplankton biomass (peuk + nano + diat; blue line), and C:P of total phytoplankton (green line) within the upper 100 m (d).
The C:P stoichiometry of the sinking organic particulate flux at 100 m (C:P_{exp}) from the PICO simulation is shown in Fig. 4c. Carbon rich/P-poor stoichiometry for C:P_{exp} is found in the ocean tropics and subtropics, being a function of both the large contribution of pico-phytoplankton productivity (Fig. 2) in these regions as well as the elevated C:P stoichiometry of the cyanobacteria (pro + syn + diaz) PFTs inhabiting these regions (Fig. 4d). Sinking particulate export C:P stoichiometry exhibits global maxima within the western subtropical gyres of each basin, reaching values of ~140 – 150 (Fig. 4c). C:P export stoichiometry is elevated elsewhere within the subtropics, ~120 – 140, near Redfieldian in the mid-latitudes and the Arctic Ocean, and depressed within the eastern equatorial Pacific, subpolar North Pacific, and the Southern Ocean, exhibiting values in the range ~90 – 105. The zonal mean C:P_{exp} is near the minimum imposed value of 90 for eukaryotic phytoplankton groups south of 50 °S and near ~100 north of 50 °N where the phytoplankton community is dominated by diatoms and other eukaryote groups (Fig. 2). Maximal zonal mean C:P_{exp} is found at subtropical latitudes exhibiting values of ~128 at 23 °S and a slightly higher value of ~133 at 26 °N in the northern hemisphere (Fig. 4d).

Cyanobacteria phytoplankton (pro + syn + diaz), eukaryotic phytoplankton (peuk + nano + diat), and the total phytoplankton community C:P stoichiometry all exhibit similar latitudinal patterns with high latitude minima and subtropical maxima (Fig. 4d). Cyanobacteria phytoplankton C:P varies between ~96 – 223, eukaryotic phytoplankton C:P varies between ~90 – 117, with total phytoplankton C:P varying between ~90 – 202.

5. Discussion
Pico-phytoplankton were found to make their largest contribution to marine NPP within the tropical and subtropical ocean, reaching contributions of ~50–90% within the latitudinal range 45°N–45°S (Fig. 2). In general, *Prochlorococcus* dominates NPP within the subtropical gyre regions, transitioning to dominance by pico-eukaryotes both equatorward and poleward of ~30° latitude to ~45°. *Synechococcus* follows a similar spatial pattern as *Prochlorococcus* but never dominates the phytoplankton community, with fractional contributions peaking at ~10–12% in the subtropical gyres. Diazotrophs, which are parameterized to represent the cyanobacteria lineages of *Trichodesmium* and *Crocosphaera* in the BEC model, make a smaller yet important additional contribution in these same low-latitude regions. The pico-phytoplankton and diazotroph groups within the PICO simulation were assigned the largest cellular C:P plasticity and higher minimum temperature thresholds for growth (Table 1) than the larger nano-phytoplankton and diatoms, with the pico-phytoplankton niche being realized within the nutrient-poor, high temperature waters of the low latitudes. The three pico-phytoplankton PFTs in PICO contributed to globally integrated NPP at a percentage of 16.6 ± 0.4% for *Prochlorococcus*, 3.9 ± 0.5% for *Synechococcus*, and 37.1 ± 0.6% for pico-eukaryotes, for a total pico-phytoplankton contribution of 57.6 ± 0.8% to marine NPP (Table 4). These estimates for global NPP attributable to *Prochlorococcus* and *Synechococcus* within PICO are approximately double and one-quarter, respectively, compared to an independent estimate based on data assimilative modeling of niche partitioning, (pro = 8.5%, syn = 16.7%, Flombaum et al., 2013), however both model estimates converge on a combined pro + syn contribution to marine NPP of ~20–25%.

Nano-phytoplankton contribute 3.6 ± 0.7% of global NPP in PICO (likely an underestimate, e.g., Brewin et al., 2010; Juranek et al., 2020), concentrated in the Southern Ocean, the subpolar and Arctic Ocean, as well as the eastern tropical Pacific Ocean (Fig. 2d). Diatoms are the largest
contributor to global marine NPP in PICO, contributing 37.5 ± 0.5%, concentrated in the subpolar and polar oceans as well as the equatorial Pacific upwelling (Table 4; Figure 2d). This rate of diatom contribution to global NPP is consistent with the ~38-43% contribution estimated from marine biogenic silica budgets (Nelson et al., 1995). Diazotrophs contribute 1.2 ± 0.04% to marine NPP in PICO (Table 4), concentrated in the same tropical and subtropical regions as pico-phytoplankton, especially the western tropical and subtropical Pacific Ocean (Fig. 2e).

<table>
<thead>
<tr>
<th>PFT</th>
<th>% of NPP</th>
<th>% of POC\text{exp}</th>
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<tbody>
<tr>
<td>\textit{Prochlorococcus}</td>
<td>16.6</td>
<td>14.9</td>
</tr>
<tr>
<td>\textit{Synechococcus}</td>
<td>3.9</td>
<td>12.6</td>
</tr>
<tr>
<td>Pico-eukaryotes</td>
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<tr>
<td>Nano-phytoplankton</td>
<td>3.6</td>
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</tr>
<tr>
<td>Diatoms</td>
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</tr>
<tr>
<td>Diazotrophs</td>
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</tr>
<tr>
<td>(\Sigma) Pico-phytoplankton</td>
<td>57.6</td>
<td>45.6</td>
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</tbody>
</table>

Table 4. Percentages of globally integrated marine NPP and POC\text{exp,100m} attributable to each PFT.

Ecosystem processing of the newly fixed organic matter by each phytoplankton PFT contributes to the detrital particulate organic matter pools, which constitute the source terms to the sinking export fluxes of POC\text{exp} and particulate organic phosphorus (POP\text{exp}) in the BEC model. These processes include losses to the detrital pool upon grazing by zooplankton, zooplankton contributions to the detrital pool as a result of grazing (e.g., fecal pellet production), and aggregation and sinking of phytoplankton biomass upon cell mortality within the BEC model (Moore et al., 2004). The parameter values governing these processes were prescribed in a size-scaling manner across the six PFTs in PICO (Table 1), with the pico-phytoplankton
contributing the smallest to the particulate sinking fluxes, increasing with diazotrophs and nano-plankton, to the largest contribution from diatoms. The fractions of pico-phytoplankton grazing and zooplankton losses that accumulate within the particulate fluxes were prescribed in the range $0.07 - 0.33$ (Table 1), at the upper end of estimates based on pigment biomarker analyses of the sediment trap flux collected from the Sargasso Sea (Lomas & Moran, 2011). The higher estimates for pico-phytoplankton detrital contributions were needed to export enough N and P below the euphotic zone to prevent spurious accumulation of $\text{NO}_3^-$ and $\text{PO}_4^{3-}$ concentrations in the upper layers of the subtropics. The nano-phytoplankton and diatom contributions to NPP are $<5\%$ each over much of the subtropical regions in PICO (Fig. 2), which are biased low compared to field observations (e.g., Nelson & Brzezinski, 1997; Brzezinski et al., 2011; Juranek et al., 2020) and may potentially explain the low N and P export efficiency, partially satisfied by prescribed elevated pico-phytoplankton detrital contributions in PICO. PFT-specific parameters linked to the detrital contributions are held constant in PICO, however future formulations of the BEC model could include parameterizations that are dependent on in situ temperature, PFT growth rate, grazing rate, or nutrient limitation status among others.

Within PICO, pico-phytoplankton contribute nearly half ($45.6 \pm 0.3\%$) of the globally integrated $\text{POC}_{\text{exp,100m}}$ flux through 100 meters when accounting for both direct (e.g., aggregation) and indirect (e.g., zooplankton processing of pico-produced organic matter) contributions. Approximately 40% of the pico-phytoplankton total is contributed by pico-eukaryotes ($18.1 \pm 0.2\%$), an additional third contributed by Prochlorococcus ($14.9 \pm 0.1\%$), and a quarter contributed by Synechococcus ($12.6 \pm 0.1\%$) (Table 4). Pico-phytoplankton dominate the phytoplankton contribution to carbon export across large areas of the tropics, subtropics, and mid-latitudes (Fig. 4b), which is an emergent result from the added phytoplankton biodiversity,
representation of variable carbon to phosphorus cellular stoichiometry, and ecosystem processing
of the detrital pool mediated by zooplankton grazing. Perhaps counterintuitively, pico-phytoplankton contribution to carbon export peaks in importance (~0.65 – 0.7), not where their greatest environmental niche is realized, i.e., the high light, low nutrient waters of the oligotrophic subtropical gyres, but at mid-latitudes, ~35 – 45°, driven largely by regional maxima in C:P stoichiometry (Fig. 4c).

The estimate of pico-phytoplankton contribution to sinking carbon export in PICO (~46%) is approximately 80% their combined contributions to global NPP (~58%), in partial agreement with inversions of PFT-specific observational data from the equatorial Pacific and Arabian Sea suggesting that pico-phytoplankton contribute to carbon export at rates comparable to their contribution to NPP (Richardson & Jackson, 2007). Group specific C export fraction to NPP fraction ratios, computed from Table 4, range from a minimum of ~0.5 to ~3.2, within the range of the available group-specific estimates diagnosed from a range of field data, ~0.1 to ~4.5 (Richardson & Jackson, 2011; Stukel & Landry, 2010; Lomas & Moran, 2011; Fawcett et al., 2011; Stukel et al., 2013), albeit at the higher end especially for Pro and Syn which are 0.9 and 3.2 in PICO and ~0.1 and ~0.1 to ~0.4 from the literature (Lomas & Moran, 2011; Fawcett et al., 2011; Stukel et al., 2013), respectively. A notable outlier is the C export fraction to NPP fraction ratio for diaz which is ~11 in PICO. Dissolved organic matter (DOM) contributes ~20-25% of total organic carbon export production in the CESM (Letscher et al., 2015). We are unable to provide an estimate of each PFT contribution to the DOM export flux, as DOM export is spatially and temporally offset from its region of production, and the model does not track DOM sourced from individual PFTs. Each PFT contributes a constant fraction of its newly produced
organic matter to the DOM pool; therefore, PFT contributions to DOM export should generally
track the proportions of PFT contributions to NPP (e.g., Table 4).

The regions located immediately along the poleward and western edges of the surface western boundary currents of the subtropical gyre regions of each ocean basin exhibit elevated POC$_{exp,100m}$ fluxes on the order of ~50 – 90 g C m$^{-2}$ yr$^{-1}$ (Fig. 4a), approximately ~2-3 fold higher as compared to the adjacent regions. These so-called ‘gyre recirculation regions’ of the surface western boundary currents are characterized by strong eddying flow and enhanced lateral nutrient concentration gradients that stand out as important regions for the delivery of a lateral nutrient flux that dominates nutrient resupply to the subtropical gyres (Williams & Follows, 1998; Oschlies, 2002; Jenkins & Doney, 2003; Torres-Valdés et al., 2009; Palter et al., 2013; Letscher et al., 2016; Yamamoto et al., 2019). The resolution of the BEC model employed in PICO is non-eddy resolving, however it is able to simulate somewhat elevated rates of NPP (Fig. 1a) and associated POC$_{exp,100m}$ (Fig 4a) along these subtropical to subpolar gyre flanks at the western side. These gyre recirculation region biogeochemical features are driven by a larger contribution of pico-eukaryotes and diatoms to simulated NPP (Fig. 2). These regions are also where PFT cellular C:P stoichiometry transitions from C-rich/P-poor stoichiometry within the subtropical gyres, to more Redfield-like C:P stoichiometry poleward (Fig. 3; Fig. 4c,d). Thus, the magnitude of POC$_{exp,100m}$ within the subtropical gyre recirculation regions identified as ‘POC$_{exp}$ hotspots’ by PICO may be sensitive to small changes to the nutrient flux stoichiometry which drive cellular C:P stoichiometry. Further field-based study of the in situ phytoplankton community composition and associated biomass C:P stoichiometry within these regions could help confirm their regional ‘POC$_{exp}$ hotspot’ status.
The PICO simulation builds from recent advances in a BEC model formulation that added variable C:P stoichiometry within the three traditionally simulated PFTs (small phytoplankton, diatoms, diazotrophs; Wang et al., 2019), by adding three pico-phytoplankton PFTs with variable C:P stoichiometry. As compared to the variable C:P, three PFT version of the BEC model (VAR-3; Wang et al., 2019), the PICO simulation exhibits ~1% higher globally integrated NPP (51.4 Pg C yr\(^{-1}\) vs. 50.8 Pg C yr\(^{-1}\)) and ~18% lower PO\(_{\text{exp},100\text{m}}\) (7.0 Pg C yr\(^{-1}\) vs. 8.5 Pg C yr\(^{-1}\)). Thus, given that both the three and six PFT, variable C:P formulations of the BEC model reproduce similarly-well, the surface to deep ocean spatial gradients in dissolved nutrient concentrations (Table 2), the pico-phytoplankton enabled (6p1z) BEC model predicts similar rates of NPP, with lower rates of PO\(_{\text{exp},100\text{m}}\), and lower C:PO\(_{\text{exp}}\) per unit of phosphorus utilized within the euphotic zone as compared with the three PFT BEC model version (VAR-3; Wang et al., 2019). The geographical distribution of PO\(_{\text{exp},100\text{m}}\) is similar between the PICO and VAR-3 simulations, however VAR-3 simulated PO\(_{\text{exp},100\text{m}}\) is approximately double to triple that estimated by PICO within the subtropics to mid-latitudes (not shown), contributing to its overall ~18% higher estimate of global PO\(_{\text{exp},100\text{m}}\). The lower PO\(_{\text{exp},100\text{m}}\) estimate by the PICO model is likely attributable to the lower prescribed values of zooplankton grazing routed to the particulate organic pool, ‘graze_poc’ parameter, for the pico PFTs (graze_poc = 0.07 – 0.13; Table 1) versus the small phytoplankton PFT in Wang et al. (2019) (graze_poc = 0.3). Maximum C:PO\(_{\text{exp}}\) is estimated within the ocean’s subtropical gyres (Fig. 4c) in both simulations, with PICO predicting slightly lower maximum C:PO\(_{\text{exp}}\) within the Indo-Pacific subtropics, ~140 – 145 vs. ~180 in VAR-3 (not shown). PICO simulated C:PO\(_{\text{exp}}\) generally agrees with the spatial pattern estimated from an inversion of dissolved inorganic carbon and phosphate data (Teng et al., 2014), with notable differences in C:PO\(_{\text{exp}}\) magnitude. PICO minimum PFT C:P is set at 90 while
the Teng et al. (2014) inversion estimates minimum C:P_{exp} of ~60 – 100 for the equatorial and subpolar regions. Maximal C:P_{exp} in the subtropical North Atlantic is ~155 in PICO compared to ~355 by Teng et al. (2014). These comparisons demonstrate that estimates of the globally important fluxes of marine NPP and carbon export including its C:P stoichiometry vary on the order of ~10 – 50% between OBGC model formulations, highlighting the continued need for observational datasets to further constrain and validate model simulations of ocean biogeochemistry.

We are aware of two other ESMs to which addition of variable phosphorus to carbon ratios within the OBGC model has been performed as well as an additional global ocean biogeochemistry model. While nearly all ESMs participating in the CMIP5 suite predict declining marine NPP during the 21st century (Bopp et al., 2013), the GFDL-ESM2 which runs the variable C:P enabled TOPAZ2 OBGC model (Dunne, 2013), was the only ESM to predict a resilient rate of marine NPP throughout this century. Presumably the mechanism at play is one in which declining nutrient fluxes to the surface ocean with increasing upper ocean stratification are countered by a switch in phytoplankton community composition towards smaller, more nutrient plastic types and/or an increase in phytoplankton C:P that allows for maintenance of carbon based NPP rates at depressed phosphorus supply fluxes. The PICO simulation presented here supports this interpretation, exhibiting a marginally ~1% higher overall rate of marine NPP than its analogous three PFT, Redfield C:P sister simulation (RED-3; Letscher & Moore, 2015). Results from a variable C:P OBGC model that added representation of pico-phytoplankton have been published (the PISCES-QUOTA model; Kwiatkowski et al., 2018). The model predicts similar 21st century declines in marine NPP (~8%) and POC_{exp} (~14%) as the fixed (Redfield) stoichiometry ESMs that participated in CMIP5 (Bopp et al., 2013), while noting an increase in
carbon export efficiency per unit phosphorus, which increases by 4.5%. Both the TOPAZ2
OBGC model and PISCES-QUOTA parameterize variable C:P in the phytoplankton following
the optimal resource cell allocation model of Klausmeier et al., (2004), so it is surprising that the
two models yield divergent predictions for 21st century marine NPP (no decline in GFDL-ESM2
using TOPAZ2, decreasing in PISCES-QUOTA), that may be due to differences in model
physics and the simulated nutrient supply to the surface. The global ocean biogeochemistry
modeling study of Tanioka & Matsumoto (2017) in which variable C:P stoichiometry was
represented in a two phytoplankton marine ecosystem model, found that predicted declines in
global export production could be mitigated by ~3% relative to a Redfield stoichiometry model
over the 21st century that the authors attributed to combined mechanisms of flexible cellular
nutrient to carbon ratios and regional shifts in phytoplankton community composition.

6. Future Directions

A next step with the CESM-BEC model is to perform a future climate forced simulation
using PICO parameterizations to investigate changes in 21st century marine NPP and export in
the context of marine phytoplankton diversity and cellular nutrient plasticity. We also wish to
point out that during development of the PICO simulation it was found that the model solution
was very sensitive to the choice of the grazing rates for each of the six PFTs by the single
zooplankton group. This single zooplankton group formulation was not well suited to act as the
top-down control on the growth of six PFTs, with the model exhibiting behavior in which it was
difficult to obtain mixed communities of phytoplankton everywhere in the ocean. Future
improvement of the BEC model, now carried out within the Marine Biogeochemistry Library
(MARBL) in CESM v2, should include the addition of explicit nanophytoplankton groups (e.g.,
coccolithophores) and additional zooplankton groups representing multiple size classes and
grazing formulations that will likely result in a more robust simulation of plankton growth,
grazing, and export dynamics. Currently, a version with four zooplankton and eight
phytoplankton functional types is in development (8p4z).

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thanked for providing output of pico-phytoplankton biomass from predictive niche models. The
authors claim no financial conflicts of interest.

Open Research

The modified BEC model source code used to generate the PICO simulation can be found on
GitHub here: https://github.com/rletscher/GBC_PICO. The 20-year annually averaged output file
for PICO presented in this work is available by request from the corresponding author.

References

& Lizotte, M. P. (1999). Phytoplankton community structure and the drawdown of nutrients and


plasticity on bulk particulate stoichiometry across ocean basins. Communications Earth & Environment, 2(1), 1-10.


Moore, J. K., Lindsay, K., Doney, S. C., Long, M. C., & Misumi, K. (2013). Marine ecosystem dynamics and biogeochemical cycling in the Community Earth System Model [CESM1 (BGC)]: Comparison of the 1990s with the 2090s under the RCP4.5 and RCP8.5 scenarios. *Journal of Climate, 26*(23), 9291-9312.


Pico-phytoplankton contribute half of global marine carbon export

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**Key Points:**

- Explicitly simulated *Prochlorococcus*, *Synechococcus*, and pico-eukaryotes contribute ~60% of global net primary productivity
- Pico-phytoplankton contribute half of global marine sinking particle export production
- Carbon export is elevated poleward of western boundary currents, dominated by pico-eukaryotes and diatoms with elevated C:P stoichiometry
Abstract

Earth System Models generally predict increasing upper ocean stratification from 21st century planetary warming, which will cause a decrease in the vertical nutrient flux resulting in declining marine net primary productivity (NPP) and carbon export fluxes. Recent advances in quantifying marine ecosystem carbon to nutrient stoichiometry have identified large latitudinal and biome variability, with low-latitude oligotrophic systems harboring pico-sized phytoplankton exhibiting large phosphorus to carbon cellular plasticity. Climate forced changes in nutrient flux stoichiometry and phytoplankton community composition is thus likely to alter the ocean’s biogeochemical response and feedback with the carbon-climate system. We have added three pico-phytoplankton functional types within the Biogeochemical Elemental Cycling component of the Community Earth System Model while incorporating variable cellular phosphorus to carbon stoichiometry for all represented phytoplankton types. The model simulates Prochlorococcus and Synechococcus populations that dominate the productivity and sinking carbon export of the tropical and subtropical ocean, and pico-eukaryote populations that contribute significantly to productivity and export within the subtropical to mid-latitude transition zone, contributing a combined 50 – 70% of these fluxes. Pico-phytoplankton cellular stoichiometry and resulting sinking export patterns inversely track the distribution of surface phosphate, with the western subtropical regions of each basin supporting the most P-poor stoichiometries. Collectively, pico-phytoplankton contribute ~58% of global NPP and ~46% of global particulate organic carbon export below 100 meters. Subtropical gyre recirculation regions along the poleward flanks of surface western boundary currents are identified as regional hotspots of enhanced carbon export exhibiting C-rich/P-poor stoichiometry, preferentially inhabited by pico-eukaryotes and diatoms.
1. Introduction

Ocean biogeochemical (OBGC) models operating within larger Earth System Models (ESMs) have been developed for the purpose of quantifying the feedbacks between ocean biology, chemistry, and the climate system as well as to provide predictions of ocean biogeochemistry fluxes and tracer concentrations decades to centuries into the future (Le Quéré et al., 2005; Friedlingstein et al., 2006; Bopp et al., 2013; Cabré et al., 2015; Moore et al., 2013; 2018). ESM output from the Coupled Model Intercomparison Project Phase 5 (CMIP5) and Phase 6 (CMIP6) has predicted ocean acidification, deoxygenation, and decreasing primary productivity during the 21st century (Bopp et al., 2013; Cabré et al., 2015, Moore et al., 2013; Kwiatkowski et al., 2020) and beyond (Moore et al., 2018) under a business-as-usual scenario, primarily as a result of changes in ocean circulation and stratification in response to warming ocean temperatures. Enhanced upper ocean stratification leads to a reduction in vertical nutrient flux to the sunlight surface layer, decreasing net primary productivity (NPP) by phytoplankton, with a concomitant reduction in carbon export to the ocean interior (Cabré et al., 2015; Wilson et al., 2022), resulting in a positive feedback with the climate system as ocean carbon uptake is diminished with sustained warming. These declines are nearly universal among CMIP5 models, most of which feature fixed C:N:P:-O$_2$ stoichiometry (i.e. the Redfield ratio; Redfield et al., 1934; 1958) to describe phytoplankton nutrient uptake, production of organic matter, and its subsequent remineralization (Bopp et al., 2013). Further, declines in key carbon cycle features persist in the latest generation CMIP6 models despite an increasingly prevalent switch to variable nutrient to carbon quotas within the marine biogeochemistry modules (Séférian et al., 2020; Wilson et al., 2022).
Common to nearly all OBGC models is the representation of one or more ‘groups’ of phytoplankton that consume inorganic nutrients, resulting in photosynthesis, net primary production, and production of autotrophic biomass. Autotrophy is most often distributed among ≥ 2 phytoplankton functional types (PFTs), that are chosen to represent differing growth strategies, e.g., slower-growing small pico- and nano-phytoplankton vs. fast-growing large diatoms, to capture biogeographic latitudinal trends in nutrient uptake rates, cell size, and export efficiency. Phytoplankton growth experiences top-down control due to grazing by one or more zooplankton groups, also resulting in the production of particulate and dissolved organic detrital pools. Bottom-up controls on phytoplankton growth include multiple nutrient limitations, light limitation, and optimum temperature effects. Generally, nutrient uptake affinity is inversely correlated with cell size with smaller PFTs better able to acquire nutrients at low in situ concentrations. Each PFT acquires nutrients to satisfy their particular cellular N:C, P:C, and/or Fe:C quota, depending on the simulated inorganic nutrient fields. These cellular nutrient quotas are invariant for fixed stoichiometry models (i.e., at Redfield proportions; Moore et al., 2004; Dunne et al., 2010; Ilyina et al., 2013; Aumont et al., 2015) or can be variable, dependent on in situ environmental parameters such as nutrient concentration (Galbraith & Martiny, 2015) or optimal resource allocation theory describing phytoplankton cellular functions (Klausmeier et al., 2004; Dunne, 2013; Kwiatkowski et al., 2018). Many culture and field observations have documented variability in cellular nutrient quotas that vary both with ambient nutrient concentrations (Tanioka et al., 2022; Karl et al., 2001; Rhee, 1978) and/or across PFTs (Martiny et al., 2013a; Baer et al., 2017; Geider & La Roche, 2002; Quigg et al., 2003). Additionally, with the emergence of flow cytometry for the study of marine microbes (Lomas et al., 2011), a large literature of field and culture based studies has accumulated in recent decades describing the
physiology, biogeography, and phylogeny of marine pico-phytoplankton, more specifically the
cyanobacterial lineages *Prochlorococcus* and *Synechococcus* and pico-eukaryotic phytoplankton,
which now allows for their representation in numerical marine ecosystem models (e.g. Moore et
al., 1998; Partensky et al., 1998; DuRand et al., 2001; Martiny et al., 2009; Buitenhuis et al.,
2012; Pasulka et al., 2013; Flombaum et al., 2013; 2020; Sohm et al., 2016; Baer et al., 2017).
Given the need to balance model complexity against limiting computational resources,
representation of additional pico-phytoplankton groups is perhaps of first order importance with
respect to introduction of variable elemental stoichiometry within OBGC models, since pico-
phytoplankton are known to exhibit the largest carbon to nutrient cellular plasticity (Quigg et al.,
2003; Klausmeier et al., 2004; Martiny et al., 2013a; Baer et al., 2017). In light of these
advances, representing the pico-phytoplankton and variable carbon to nutrient content of all
phytoplankton in OBGC models has become a pressing goal to better capture the ocean
ecosystem response to the changing 21st century climate and its associated feedbacks with the
climate system.

Many recent studies have identified and quantified the non-Redfield, variable carbon to
nutrient stoichiometry that permeates most ocean biogeochemical fluxes and tracer pools
including dissolved nutrients (Weber & Deutsch, 2010), biomass (Tanioka & Matsumoto, 2020),
particulate organic matter (Martiny et al., 2013a), dissolved organic matter (Letscher & Moore,
2015), export production (Teng et al., 2014), and organic matter remineralization (DeVries &
Deutsch, 2014). The emergent pattern from these studies is one of carbon-rich (nutrient-poor)
stoichiometry that describes biogeochemical functioning in the lower latitude, nutrient-poor
biomes of the subtropics, while the nutrient-replete upwelling and subpolar biomes exhibit
stoichiometries at or slightly below Redfield carbon to nutrient proportions. Two mechanisms
have been suggested to contribute to the observed biogeography of marine ecosystem stoichiometry including phosphorus and/or nitrogen plasticity by the extant phytoplankton community (e.g. Galbraith & Martiny, 2015; Van Mooy et al., 2009) as well as shifts in phytoplankton community composition across biomes with relatively nutrient-deficient cyanobacteria dominating the subtropics and more nutrient-rich diatoms and other larger phytoplankton dominating elsewhere (e.g. Arrigo et al., 1999; Weber & Deutsch, 2010; Sharoni & Halevy, 2020). Cell-sorted field populations of cyanobacteria (Prochlorococcus and Synechococcus) and pico-eukaryotes from the Sargasso Sea, eastern Pacific, and Indian Ocean suggest both mechanisms may be at play as cyanobacteria exhibited lower cellular P content as compared to eukaryotes, while all groups exhibited cellular P plasticity in response to latitudinal gradients in inorganic phosphorus availability, albeit at varying degrees across taxonomic groups (Martiny et al., 2013a; Baer et al., 2017; Lomas et al., 2021).

Does representation of variable carbon to nutrient stoichiometry within OBGC models potentially buffer the well-documented positive ocean carbon cycle feedback to future climate warming? A number of ESM’s that have included representation of variable carbon to phosphorus stoichiometry within the ocean ecosystem component generally predict a more resilient response for future marine NPP and carbon export fluxes, with modest ~0 – <5% declines by year 2100 (Bopp et al., 2013; Cabré et al., 2015; Tanioka & Matsumoto; 2017; Kwiatkowski et al., 2018; Buchanan et al., 2018; Matsumoto et al., 2020; Kwon et al., 2022). Inclusion of variable phosphorus to carbon stoichiometry within the marine ecosystem component permits flexibility in phytoplankton phosphorus quotas. Such frugal use of nutrients allows phytoplankton to maintain similar growth rates under decreasing P flux as well as a shift towards increasing biomass C:P that partially compensates for warming induced changes to
ocean circulation and stratification. Galbraith and Martiny (2015) demonstrated with a simple box model that changes to the preformed phosphate concentration on the order of 100 nM within low latitude surface waters can produce up to 20 ppm shifts in atmospheric CO$_2$ concentrations, assuming P is the primary growth limiting nutrient. An OBGC modeling study incorporating representation of variable marine elemental stoichiometry found that adding this sensitivity buffered against a predicted decrease in global carbon export production on the order of ~3% for the 21$^{\text{st}}$ century (Tanioka & Matsumoto, 2017). Similarly, results from the recent inclusion of this process within the Pelagic Interactions Scheme for Carbon and Ecosystem Studies Quota (PISCES-QUOTA) model, suggest its effects on 21$^{\text{st}}$ century ocean carbon uptake are modest with up to a 3.5% underestimation as compared to fixed stoichiometry ESMs (Kwiatkowski et al., 2018). However, larger regional changes of up to 50% reductions or stimulations of marine NPP and carbon export were predicted which could have significant local to regional impacts on marine ecosystems and fisheries. The above studies have thus far considered adaptive nutrient quotas, while other phytoplankton adaptive strategies including altered physiological rates, trait-based evolution, and changes in grazing pressure could all potentially influence phytoplankton fitness and resilience with important controls on marine NPP and carbon export trends (Martiny et al., 2021).

As the next generation of OBGC models mature to better represent variable carbon to nutrient stoichiometry with the goal to study how this variability impacts the marine NPP and carbon export fluxes, an important missing element to this question is the role for changes in marine phytoplankton biodiversity to modulate the response. Most of the current generation of OBGC models represent the marine phytoplankton community with 1 – 3 PFTs, with the notable exception of the ECCO-Darwin OBGC model which simulates 35 up to 350 PFTs (e.g.,
Dutkiewicz et al., 2020; Juranek et al., 2020). This higher biodiversity model has been shown to more accurately represent the marine phytoplankton diversity regionally to globally, as gauged by such metrics as phytoplankton size distribution, biomass, size-fractionated NPP rates, and species richness. How does marine phytoplankton diversity and associated carbon to nutrient stoichiometric variability impact the regional to global patterns of carbon export? Based on current understanding, enhanced thermal stratification of the water column depresses the vertical nutrient input, favoring a shift to smaller sized pico-phytoplankton dominated by cyanobacteria possessing higher plasticity in cellular nutrient quotas. Thus, in a future warmer ocean with expanding stratification, the marine biodiversity response might be hypothesized to favor marine cyanobacteria such as Prochlorococcus and Synechococcus with elevated C:P cellular stoichiometry. While the shift to smaller phytoplankton may depress marine NPP rates, export of a portion of the resultant biomass with high C:P may partially buffer against reductions in marine carbon export and mitigate the expected positive feedback to warming by maintaining the strength of the ocean’s biological carbon pump. In support of this hypothesis, observations from the Bermuda Atlantic Time-series station in the Sargasso Sea document a recent decline in marine NPP due to enhanced stratification, without a concomitant decline in carbon export, explained by a shift in the phytoplankton community composition in favor of cyanobacteria with higher cellular C:P stoichiometry (Lomas et al., 2022).

This paper describes the addition of three new pico-phytoplankton functional types, Prochlorococcus, Synechococcus, and pico-eukaryotes, to the OBGC component of the Community Earth System Model, with the goal to better characterize the role for marine phytoplankton biodiversity and associated variable cellular carbon to phosphorus stoichiometry in controlling marine carbon export patterns. The implementation builds from recent advances in
this OBGC model to incorporate variable C:P stoichiometry within all pertinent tracers of the marine ecosystem (Letscher et al., 2015; Wang et al., 2019) including phytoplankton, dissolved organic matter, and sinking particulate organic matter, to now include representation of six PFTs: *Prochlorococcus*, *Synechococcus*, pico-eukaryotes, nano-phytoplankton (including implicit coccolithophores), diatoms, and diazotrophs.

2. Model and Validation Data

The model is the Biogeochemical Elemental Cycling (BEC) model, which is the biogeochemistry and ecosystem component operating within the ocean circulation model (POP2) of the Community Earth System Model (CESM) v.1.2.2 (http://www.cesm.ucar.edu/models/cesm1.2/). The model resolution employed here is nominally 3º x 3º horizontally with 60 vertical layers (g3v7 grid) that increase in thickness with depth in the water column. The ocean circulation is the output of the Parallel Ocean Program 2 (POP2) ocean circulation model using the final thirty-year forcing from years 1980 – 2009 of the CORE-II forcing (Smith et al., 2010) and includes parameter value changes to the isopycnal diffusivity (κiso) to partially account for the transport of unresolved equatorial jets (Getzlaff & Dietze, 2013).

The BEC tracks the cycling of key biogeochemical tracers (e.g., C, N, P, Fe, and O₂) amongst dissolved inorganic, dissolved organic, sinking particulate, and four plankton biomass pools (Moore et al., 2004). Net primary production in the model is partitioned amongst three PFTs—small phytoplankton (with implicit calcifiers), diatoms, and diazotrophs, which differ in their relative competitiveness for limiting nutrients (N, P, Si, and Fe). In addition to bottom-up nutrient and light control, phytoplankton mortality is controlled top-down by grazing from one zooplankton group. Version 1.2.2 of the BEC contains significant improvements to the cycling of
marine dissolved organic matter (Letscher et al., 2015). More recent model development for
BEC v1.2.2 has incorporated variable phosphorus quotas for the three phytoplankton types
(Wang et al., 2019) that depend on in situ simulated inorganic phosphate concentration following
the linear relationship in Galbraith & Martiny (2015) but with an imposed minimum (110:1) and
maximum (210:1) C:P value for all PFTs. This current work describes the addition of three new
pico-phytoplankton groups with variable carbon to nutrient (P and Fe) stoichiometry to the BEC
model, building from the code base for the variable 3 PFT phosphorus quota model of Wang et
al., (2019). The ‘small phytoplankton’ group from the 3 PFT BEC model which jointly
represented pico- and nano-sized phytoplankton has been reparameterized to more closely
represent larger nano-phytoplankton. The newly named nano-phytoplankton along with the
diatom and diazotroph groups are updated to represent variable C:P and C:Fe stoichiometry. The
model retains one zooplankton group with Redfieldian stoichiometry. The marine ecosystem in
the BEC model is thus represented by 6 phytoplankton and 1 zooplankton groups, or 6p1z.

Output of the pico-phytoplankton BEC model is validated for the simulation of
chlorophyll a and NPP against the Sea-Viewing Wide Field-of-View Sensor (SeaWiFS) satellite
mission climatology products (1997 – 2010) related to chlorophyll a concentration (NASA
GSFC, 2018) and a model of carbon-based net primary productivity estimated from SeaWiFS
data products (Behrenfeld et al., 2005; Westberry et al., 2008). Simulated nutrient concentrations
from the pico-phytoplankton BEC model are compared against the World Ocean Atlas 2013
climatologies of nitrate, phosphate, and silicate (Garcia et al., 2013) and a global compilation of
dissolved Fe concentrations (Tagliabue et al., 2012). Simulated pico-phytoplankton biomass are
validated against the data assimilative niche modeling of Flombaum et al. (2013; 2020) that
predicts field observed Prochlorococcus, Synechococcus from inputs of temperature and
irradiance (Flombaum et al., 2013) and pico-eukaryote biomass from inputs of temperature, irradiance, and dissolved nitrate (Flombaum et al., 2020). Simulated pico-phytoplankton C:P stoichiometry and biomass are compared against observations of populations from the North Atlantic, eastern Pacific, and eastern Indian Oceans of Prochlorococcus, Synechococcus, and pico-eukaryotes measured for C and P content (Baer et al., 2017; Lomas et al., 2021). Simulated total phytoplankton biomass C:P stoichiometry is compared against a global compilation of bulk particulate organic matter C, N, and P content (Martiny et al., 2014).

3. Methods

The addition of three pico-phytoplankton groups with variable carbon to nutrient stoichiometry introduces 12 additional tracers to the BEC model code: biomass carbon (C), chlorophyll (Chl), iron (Fe), and phosphorus (P) for each of the 3 pico-phytoplankton groups, termed ‘Prochlorococcus’ (pro), ‘Synechococcus’ (syn), and ‘Pico-eukaryotes’ (peuk). The ‘small phytoplankton’ group from BEC v1.2.2 and earlier versions has been renamed ‘nano-phytoplankton’ (nano) and retains the implicit calcifiers. Nano-phytoplankton as well as the diatoms (diat) and diazotroph (diaz) phytoplankton groups carry the same four biomass tracers (C, Chl, Fe, P) as the newly added pico-phytoplankton, with the addition of biomass silicon for diatoms. Variable biomass nitrogen (N) content is not considered at this time and phytoplankton biomass N:C is fixed for all six phytoplankton groups at a value of 0.1333 (C:N = 7.5) near the mean of open ocean suspended particles (Martiny et al., 2013b) and nutrient replete versus deplete cells across a range of phyla (Geider & La Roche, 2002).

Growth and mortality of each phytoplankton type is modeled with more than two dozen parameter values specific to each group including half saturation constants for nutrient uptake,
photosynthesis vs irradiance curves, Fe and P quotas, temperature thresholds for growth,
contributions to detrital pools, and zooplankton grazing rates among others. Table 1 lists these
parameter values for each phytoplankton group including the relevant literature sources. Where
literature sources are lacking to inform the value for a particular parameter, the principle of
allometric scaling is employed to assign values corresponding with cell size with the order of
increasing cell size following: pro/syn/peuk < nano/dia < diat.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Units</th>
<th>pro</th>
<th>syn</th>
<th>peak</th>
<th>nano</th>
<th>diat</th>
<th>diaz</th>
<th>Reference</th>
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<td>KFe</td>
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<td>0.0404-3</td>
<td>0.0572-3</td>
<td>0.060-3</td>
<td>0.060-3</td>
<td>0.060-3</td>
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</tr>
<tr>
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<td>umol/mL</td>
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<td>0.015</td>
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<td>0.025</td>
<td>Lomas et al., 2014; Stukel et al., 2014; Garcia et al., 2015</td>
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<tr>
<td>KdOP</td>
<td>half sat. const. organic P</td>
<td>umol/mL</td>
<td>0.8</td>
<td>0.9</td>
<td>0.15</td>
<td>0.2</td>
<td>0.1</td>
<td>0.25</td>
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<tr>
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Table 1. Description of phytoplankton parameters and values from the PICO simulation of the
BEC model.

Phytoplankton iron uptake half saturation constants are prescribed with cyanobacteria
having the lowest values (pro = 30 pM; syn+dia = 40 pM), followed by pico-eukaryotes (43
pM), nano-eukaryotes (55 pM), and the large celled diatoms (60 pM) following the culture
studies of Shiré and Kustka, (2015) and Timmermans et al., (2005) and the field study of
Hutchins et al., (2002) (Table 1). Phosphate half saturation constants are selected based on the
field study from the Sargasso Sea of Lomas et al., (2014), the culture study of Garcia et al., (2015), and the Amazon River plume ecosystem modeling work of Stukel et al., (2014) with kPO4 values increasing from a value of 8 nM for pro, 10 nM for syn, 15 nM for peuk, 25 nM for diaz, 40 nM for nano, and 70 nM for diatoms. Semilabile dissolved organic phosphorus (DOP) is bioavailable for phytoplankton to utilize as a P source when phosphate levels are low in the BEC model (Letcher et al., 2015) with the half saturation constants for its uptake selected based on several field studies from the Sargasso Sea and eastern North Pacific (Dyhrman & Ruttenberg, 2006; Sohm & Capone, 2006; Casey et al., 2009; Orchard et al., 2010). Following these studies, eukaryotes and diazotrophs are assigned the lowest half saturation values for DOP uptake ranging from 100 – 250 nM, with higher kDOP values ranging from 800 – 900 nM for the two cyanobacteria groups. Half saturation constants for ammonium are assigned following allometric scaling in the range 5 – 35 nM. Nitrate uptake half saturation constants are similarly assigned following allometric scaling in the range 150 – 350 nM, with exceptions for syn and peuk which are assigned the lowest values of 180 and 150 nM respectively, owing to their demonstrated increased affinity for nitrate relative to pro in the Sargasso Sea (e.g., Glover et al., 1988; Fawcett et al., 2013) and observed correlations between peuk abundance and in situ nitrate (Flombaum et al., 2020). Diazotroph kNH4 and kNO3 are set to the highest value across the 6 PFTs to reflect its reliance on internally fixed N for a significant but not exclusive fraction of its N quota (Knapp, 2012).

Phytoplankton iron quotas in the BEC model allow for flexibility in phytoplankton biomass Fe:C based on in situ dissolved iron concentrations, varying in the range of 3 µmol:mol to 180 µmol:mol (Table 1). Following the field study of Boyd et al., (2015), eukaryotes are assigned the lowest minimum iron quotas followed by diazotrophs and non-N-fixing
cyanobacteria (*Prochlorococcus* and *Synechococcus*). The diazotroph group is assigned the highest maximum iron quotas based on the high iron requirement of the nitrogen-fixing enzyme nitrogenase (Berman-Frank et al., 2007). Phytoplankton iron quotas are reduced linearly when *in situ* dissolved Fe concentrations fall below a prescribed threshold (FeOpt), set to 1.75 nM for diatoms, 1.6 nM for nano, 1.5 nM for diaz, 1.0 nM for peuk, and 0.8 nM for pro and syn (Wisemann et al., under review).

Variable phosphorus quotas are implemented here for the six phytoplankton group BEC model following the approach introduced in Wang et al. (2019) and similar to the variable Fe:C approach described above. Variable phytoplankton biomass P:C is modeled as a function of *in situ* dissolved phosphate concentration using the linear relation detailed in Galbraith & Martiny (2015) diagnosed from a global compilation of suspended particulate organic phosphorus and carbon concentrations. Each PFT P:C quota (gQp) is computed from:

\[
gQp = \begin{cases} 
(gQp_0 - gQp_{min}) \times \frac{PO4}{POpt} + gQp_{min} & \text{PO4 < POpt} \\
gQp_0 & \text{PO4 \geq POpt}
\end{cases} \tag{Eq. 1}
\]

where PO4 is the *in situ* dissolved inorganic phosphate concentration (µM) and gQp_0 and gQp_min describe the minimum and maximum P:C quota (see Table 1). Equation 1 linearly decreases the phytoplankton P:C quota (increases its C:P ratio) below a prescribed threshold (POpt; range 0.5 – 1.0 µM across PFTs), from its initial maximum value, currently prescribed at gQp_0 = 0.01111; C:P = 90:1 for the eukaryotes (peuk+nano+diat), gQp_0 = 0.01042; C:P = 96:1 for the cyanobacteria, and gQp_0 = 0.006667; C:P = 150:1 for diazotrophs, towards a minimum P:C quota that varies for each PFT (Table 1). This formulation is similar in approach to that presented in Galbraith & Martiny (2015). Here, field observations of surface ocean particulate organic carbon and phosphorus (of which autotrophic biomass is a significant
contributor) were found to fit a linear function of *in situ* phosphate concentration. The formulation presented here prescribes differing P:C versus PO₄ slopes and y-intercepts for each PFT set by the choice of gQp_min. Field-based estimates of cell-sorted biomass C:P and N:P for *Prochlorococcus*, *Synechococcus*, and pico-eukaryotes from the Sargasso Sea (Martiny et al., 2013a) and elsewhere (Baer et al., 2017; Lomas et al., 2021) are used to assign values of gQp_min with a maximum C:P of 300:1 for diazotrophs, a C:P of 270:1 for *Prochlorococcus* and 192:1 for *Synechococcus*, and a maximum C:P of 120:1 for peuk, nano, and diat (retained from the variable C:P BEC model; Wang et al., 2019).

Based on the phytoplankton niche modeling of Flombaum et al. (2013; 2020), each phytoplankton group in the BEC is assigned a temperature threshold below which photosynthesis rates and thus growth rates are depressed, with *Prochlorococcus* and diazotroph niches favoring warmer temperatures above 13°C and 18°C, respectively. *Synechococcus* and pico-eukaryote growth decreases as temperatures fall below 4°C and -1.5°C, respectively, while nano-phytoplankton and diatom growth rates are not depressed as a function of temperature (temp_thres value set to -10°C in Table 1).

Physiological response to light parameters describing the pico-phytoplankton are assigned based on the culture study of Stawiarzski et al. (2017) in which two strains of *Prochlorococcus* (HL RCC 296, LL RCC 162), one *Synechococcus* strain (RCC 30), and four pico-eukaryote species *Triparma eleuthera* (RCC 212), *Micromonas pusilla* (RCC 1677), *Picochlorum* sp. (TCC 289), and *Nannochloropsis granulata* (RCC 438) from the Roscoff culture collection (Vaulot et al., 2004) were studied. Following this study, cyanobacteria are assigned slower maximum growth rates compared with eukaryotes, in the range 1.9 – 4 d⁻¹ (based on a 24-hour light period). Stawiarzski et al. (2017) found that light affinity is inversely related to
the maximum growth rate. *Prochlorococcus* and *Synechococcus* are assigned the highest light affinities, as encoded within the ‘alphaPI’ model parameter which describes the initial slope of the photosynthesis-irradiance curve, with decreasing affinities for the faster growing eukaryotes (Table 1). The maximum Chl:N, the ‘thetaN_max’ model parameter, is assigned inversely proportional to maximum growth rate with the exception of *Synechococcus* which exhibits lower Chl:C in line with the results of Stawiarski et al. (2017).

Grazing on each of the six phytoplankton groups in the updated BEC model is carried out by one adaptive zooplankton group. Grazing rates are largely unconstrained by field data. In the BEC model, maximum grazing rates, ‘z_umax_0’ parameter, are adjusted to achieve satisfactory biomass distributions and C:P stoichiometry for each phytoplankton group as validated against phytoplankton niche modeling and flow-sorted biomass stoichiometry distributions. Variable fractions of phytoplankton biomass are routed to zooplankton losses to detritus (e.g., fecal pellets) and particulate organic matter pools resulting from zooplankton grazing (Table 1). Each PFT’s contribution to the sinking export flux following grazing varies from a minimum of 3% for diazotrophs to a maximum of 40% for diatoms (‘graze_poc’; Table 1) informed in part by pigment biomarker analyses of the sediment trap flux in the Sargasso Sea (Lomas & Moran, 2011). Zooplankton contribute varying proportions of grazed phytoplankton biomass to detritus (POC) based on the PFT grazed, ranging from a minimum of 20% for nano-phytoplankton and diatoms to a maximum of 36% when grazing diazotrophs (‘f_zoo_detr’; Table 1). Grazing of phytoplankton biomass also contributes 6% to the dissolved organic matter pool following the inverse model diagnosed rate from our previous work (Letscher et al., 2015). In the BEC model, combinations of two or more phytoplankton types can be grouped together in ‘grazee’ classes, for which zooplankton grazing rates are computed jointly based on the total amount of biomass
in each simulated \textit{in situ} ‘grazee’ class. Experimentation was carried out to decide which combinations of phytoplankton groups in each ‘grazee’ class achieved distributions of phytoplankton biomass, NPP, and dissolved nutrients that best matched observations. A model configuration in which all six PFTs were ungrouped, i.e., parameter grazee\_ind was set equal to auto\_ind, the unique index of each of the six PFTs, was chosen.

BEC simulations were carried out for sixty model years from initial conditions, allowing the upper ocean ecosystem to achieve tracer equilibrium. After selection of the set of parameter values that best matched observational constraints, a longer 300 model year simulation was performed, repeating the 30-year (1980 – 2009) CORE-II forcing for a total of ten repetitions. The annually averaged output from the final twenty years of this simulation were averaged and is presented in all results to follow. Reported flux variability represents ±1 S.D. of the mean from the twenty years of output. The BEC simulation containing the additional pico-phytoplankton groups for a total of 6 PFTs is termed PICO. In Section 4, this simulation is compared against two similarly forced previous versions of the BEC model: a variable C:P stoichiometry version with 3 PFTs herein termed VAR-3 (Wang et al., 2019) and a 3 PFT version with Redfield stoichiometry governing phytoplankton dynamics herein termed RED-3 (Letscher et al., 2015).

4. Results

\textit{Chlorophyll a & NPP}—Sea surface chlorophyll a (chl a) concentrations [mg m$^{-3}$] from the PICO simulation and the SeaWiFS satellite climatology are shown in Figure 1a-b. BEC simulated chl a captures the overall pattern of lowest concentrations within the subtropical ocean gyres, highest concentrations within coastal and subpolar zones, and intermediate concentrations in upwelling regions and subtropical to mid-latitude transitions zones (Fig. 1). The PICO
simulation exhibits an overall global negative mean bias in chl a (−6%; Table 2) relative to the SeaWiFS climatology, reducing by a factor of ~3 compared with the 3 PFT variable C:P stoichiometry BEC model (VAR-3) and an earlier version containing 3 PFTs and Redfield stoichiometry (RED-3) (Table 2).

### Table 2. Comparison metrics for PICO, VAR-3, and RED-3 simulated chlorophyll a and NPP versus satellite (SeaWiFS) climatology estimates using the CbPM model. Metrics for dissolved nutrients from the same three simulations are compared versus the World Ocean Atlas 2013 (NO$_3^-$, PO$_4^{3-}$, SiO$_4^{4-}$) and Tagliabue et al. (2012) Fe data compilation.

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<td>[dFe]</td>
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Vertically integrated annual net primary productivity (NPP) [g C m$^{-2}$ yr$^{-1}$] from the PICO simulation is provided along with the Carbon-based Productivity Model estimate of NPP from the SeaWiFS climatology (Westberry et al., 2008) (Figure 1c-d). The globally integrated annual NPP from PICO is $51.4 \pm 0.75$ Pg C yr$^{-1}$, within the range of satellite-based estimates – 60-67 Pg C yr$^{-1}$ (Behrenfeld et al., 2005), 52 Pg C yr$^{-1}$ (Westberry et al., 2008; Silsbe et al., 2016), 58 ± 7 Pg C yr$^{-1}$ (Buitenhuis et al., 2013). Highest rates of annual NPP are simulated in the equatorial Pacific, Arabian Sea, and the South Pacific and North Atlantic eastern boundary upwelling zones, reaching rates of ~400 – 600 g C m$^{-2}$ yr$^{-1}$ in the PICO simulation, which is approximately
25 – 75% greater than those estimated by the CbPM. The lowest rates of NPP simulated by PICO are found within the Arctic Ocean, which are approximately one-quarter of those estimated by CbPM. The PICO simulation also simulates higher rates of NPP as compared to CbPM within the mid-latitudes of both hemispheres, especially near the western boundary of each ocean basin. Overall, the PICO simulation exhibits a small positive mean bias of +3% in estimated global annual NPP rates when comparing binned 3ºx3º pixels with the CbPM SeaWiFS climatology (Table 2).

**Figure 1.** PICO simulated 20-yr averaged sea surface chlorophyll a [mg m⁻³] (a), climatological average sea surface chlorophyll a from the SeaWiFS satellite during 1997-2010 (b), PICO simulated 20-yr averaged NPP [g C m⁻² yr⁻¹] (c), and from the CbPM model using SeaWiFS satellite data from the same era (d) (Westberry et al., 2008).

PFT contributions to spatial NPP patterns – The fractional contribution of each of the six PFTs to simulated NPP rates within the PICO simulation is shown in Figure 2. The majority of
NPP within the five subtropical gyre regions is contributed by *Prochlorococcus* (fraction of NPP \( \sim 0.50 \text{–} 0.90 \)), with a general western intensification (Fig. 2a). *Synechococcus* NPP is similarly elevated within the subtropics, albeit at lower fractional contributions in the range \( \sim 0.06 \text{–} 0.12 \) (Fig. 2b). *Prochlorococcus* and *Synechococcus* are mostly absent from the polar oceans due to their imposed temperature threshold of 13°C and 4°C for positive growth rates (Flombaum et al., 2013; Table 1). Pico-eukaryotes dominate NPP (fraction of NPP > 0.80) in many of the eastern boundary and mid-latitude regions of the world ocean in PICO (Fig. 2c). Elsewhere, pico-eukaryote contribution to simulated NPP is much reduced, in the range \( \sim 0.02 \text{–} 0.10 \). Nano-phytoplankton contribute variably to NPP rates within a range of 0.02 – 0.20 throughout the world ocean in PICO (Fig. 2d). Lowest simulated contributions are found in the eastern and equatorial upwelling regions; highest contributions are located in the polar oceans. Diatoms dominate simulated NPP rates (fraction > 0.80) within the subpolar and polar oceans as well as the equatorial Pacific (Fig. 2e), with lowest fractional contributions to NPP (\( \sim 0.01 \text{–} 0.05 \)) located within the subtropical gyres. Diazotrophs contribute < 0.07 to simulated NPP rates and are absent from the subpolar and polar oceans, due to the imposed minimum temperature of 18°C for positive growth rates (Table 1). Highest diazotroph contributions to NPP rates are found within the western tropical to subtropical North and South Pacific Oceans (\( \sim 0.04 \text{–} 0.07 \)) (Fig. 2f), similar to other OBGC model simulations that incorporate variable nutrient stoichiometry (e.g., Wang et al., 2019).
Figure 2. Fraction of vertically integrated, annual marine NPP attributable to each PFT from the PICO simulation. (a) Prochlorococcus, (b) Synechococcus, (c) pico-eukaryotes, (d) nano-phytoplankton, (e) diatoms, (f) diazotrophs. Note change in range in panels a,c,e from panels b,d, and panel f.

Nutrients – Upper ocean (0 – 500 m) simulated dissolved nutrient concentrations from PICO have been compared against the World Ocean Atlas (WOA) 2013 climatology (Garcia et al., 2013) for nitrate (NO$_3^-$), phosphate (PO$_4^{3-}$), and silicate (SiO$_4^{4-}$) and a global compilation of dissolved Fe (dFe) measurements (Tagliabue et al., 2012). Table 2 provides the r-correlation parameter and mean bias (µM) from a regression of log-transformed model-observation pairs. Dissolved NO$_3^-$, PO$_4^{3-}$, and SiO$_4^{4-}$ are equally well correlated with WOA nutrients, exhibiting r-correlation coefficients of ~0.94 and mean biases of +0.01 µM NO$_3^-$, +0.02 µM PO$_4^{3-}$, and +0.01 µM SiO$_4^{4-}$. PICO simulated dFe is the most weakly correlated nutrient with observational data, r = 0.40, with a mean bias of +0.18 nM. The PICO upper ocean dissolved nutrient mean biases are similar to those of the other variable C:P stoichiometry simulation, VAR-3, and the Redfield simulation, RED-3, with a notable improvement to the mean bias of SiO$_4^{4-}$ from VAR-3 (Table 2).
Biomass – PICO simulated biomass (µg C L⁻¹) is compared against the data assimilative niche model output for Prochlorococcus and Synechococcus (Flombaum et al., 2013) and pico-eukaryotes (Flombaum et al., 2020) binned by 3°x3° pixels. Simulated pico-phytoplankton biomass correlation coefficients with the niche model output range from 0.485 (pico-eukaryotes) to 0.637 (Prochlorococcus) (Table 3). However, significant negative biases are present within PICO simulated Prochlorococcus biomass (–56%), Synechococcus biomass (–85%), and pico-eukaryotes biomass (–76%) (Table 3). PICO pico-phytoplankton biomass has also been compared against a globally compiled field-based dataset of flow-sorted populations (Baer et al., 2017; Lomas et al., 2021), yielding metrics of: Prochlorococcus biomass (+76%, r = 0.199), Synechococcus biomass (–72%, r = 0.245), and pico-eukaryotes biomass (–72%, r = 0.142) (Table 3).

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<th>r-correlation</th>
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<td>Peuk C</td>
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<td>-72%</td>
<td>4158</td>
</tr>
</tbody>
</table>

Table 3. Comparison metrics for PICO simulated pico-phytoplankton biomass versus the data assimilative niche models of Flombaum et al. (2013; 2020) (Pro C niche, Syn C niche, Peuk C niche); versus pico-phytoplankton biomass C:P stoichiometry (Baer et al., 2017; Lomas et al., 2021) (Pro C:P, Syn C:P, Peuk C:P) and biomass C (Pro C, Syn C, Peuk C); and versus bulk particulate organic matter C:P stoichiometry (Martiny et al., 2014) (POC:POP).
Phytoplankton Stoichiometry – Euphotic zone-averaged PFT C:P stoichiometry from the PICO simulation is shown in Figure 3. *Prochlorococcus* (Fig. 3a) and diazotrophs (Fig. 3f) exhibit the greatest range in cellular C:P stoichiometry, with the minimum stoichiometry found within the eastern equatorial Pacific Ocean, C:P = ~90 for pro; C:P = ~150 for diaz, increasing to maximum cellular C:P = ~250 – 300 within the North and South Atlantic subtropical gyres. Elevated C:P stoichiometry for *Prochlorococcus* and diazotrophs is also found within the other three subtropical gyre regions, with typical values in the range ~180 – 300. *Synechococcus* (Fig. 3b), pico-eukaryotes (Fig. 3c), nano-phytoplankton (Fig. 3d), and diatoms (Fig. 3e) all exhibit similar geographical variability in cellular C:P stoichiometry to *Prochlorococcus* and diazotrophs, with elevated values typical of the five subtropical ocean gyre regions, syn C:P = ~150 – 210; peuk + nano + diat C:P = ~110 – 125. Phytoplankton C:P stoichiometry is ~25 – 80% more elevated within the western half of each subtropical gyre relative to the eastern half.

**Figure 3.** Euphotic zone averaged cellular C:P stoichiometry for each PFT from the PICO simulation. (a) *Prochlorococcus*, (b) *Synechococcus*, (c) pico-eukaryotes, (d) nano-phytoplankton, (e) diatoms, (f) diazotrophs.
PICO simulated *Prochlorococcus*, *Synechococcus*, and pico-eukaryotes cellular C:P stoichiometry has been compared against a subset of the cell-sorted population dataset for which biomass P was measured (Baer et al., 2017; Lomas et al., 2021). PICO simulated *Prochlorococcus* C:P exhibits the smallest mean bias of −6% below the average field-based estimate of C:P = 221 (Table 3). PICO simulated *Synechococcus* C:P has a negative mean bias of −17% below the field estimate of 187. PICO simulated pico-eukaryotes exhibited a positive mean bias compared to field estimates at +11% above the field estimate of 103. PICO simulated total autotrophic biomass C:P stoichiometry is compared with a global compilation of surface ocean suspended organic matter C:P stoichiometry (Martiny et al., 2014), exhibiting a positive mean bias of +20% above the field-based estimate of POC : POP = 171 : 1 (Table 3).

*Sinking carbon export* – The sinking particulate organic carbon flux through the 100 meter depth horizon (POC\textsubscript{exp,100m}) from the PICO simulation is plotted in Figure 4a. POC\textsubscript{exp,100m} exhibits a similar geographical distribution as simulated NPP (Fig. 1c), with maximum rates of ~90 – 120 g C m\(^{-2}\) yr\(^{-1}\) found within the eastern equatorial Pacific, Peruvian, and Arabian Sea upwelling regions as well as the SW Atlantic Ocean and minimum rates, < 20 g C m\(^{-2}\) yr\(^{-1}\), within the five subtropical ocean gyres and their bordering tropical regions, the Southern Ocean, and the Arctic Ocean. Other regions exhibiting elevated POC\textsubscript{exp,100m} include the eastern boundary upwelling zones and the subtropical to mid-latitude transition zones with rates in the range ~40 – 70 g C m\(^{-2}\) yr\(^{-1}\). The pico-phytoplankton contribution to POC\textsubscript{exp,100m} is greatest across the tropics, subtropics, and mid-latitudes (Fig. 4b), with a fractional contribution in the range ~0.50 – 0.70. In both hemispheres, pico-phytoplankton make their largest contribution to sinking POC export in the mid-latitudes (~40 – 50 °N and °S), approaching 65 – 70% of POC\textsubscript{exp,100m}, with minima
(~0.10 – 0.30) estimated for the subpolar, polar, and equatorial Pacific Oceans. The globally integrated POC\textsubscript{exp,100m} flux below 100 m from PICO is 7.0 ± 0.1 Pg C yr\textsuperscript{-1}; within the range of observational-, satellite-, and model-based estimates of global marine export production, e.g., 4 – 14 Pg C yr\textsuperscript{-1} (summarized in Emerson, 2014; Britten & Primeau, 2016; Siegel et al., 2014; Henson et al., 2011). It should be noted when comparing the PICO carbon export flux to other estimates from the literature that DOC export below 100 m contributes an additional ~2.1 Pg C yr\textsuperscript{-1} of carbon export production in the BEC model (Letscher et al., 2015; Letscher & Moore, 2015) bringing estimated total organic carbon export below 100 m to ~9.1 Pg C yr\textsuperscript{-1}.

Figure 4. PICO simulated 20-yr averaged sinking particulate organic carbon export below 100 m POC\textsubscript{exp,100m} [g C m\textsuperscript{-2} yr\textsuperscript{-1}] (a), the fraction of total POC\textsubscript{exp,100m} attributable to pico-phytoplankton (pro + syn + peuk) (b), the C:P stoichiometry of sinking organic particles C:P\textsubscript{exp} at 100 m (c), and the zonal mean C:P\textsubscript{exp} at 100 m (black line), C:P of cyanobacteria (pro + syn + diaz; red line), C:P of eukaryotic phytoplankton biomass (peuk + nano + diat; blue line), and C:P of total phytoplankton (green line) within the upper 100 m (d).
The C:P stoichiometry of the sinking organic particulate flux at 100 m (C:P_{exp}) from the PICO simulation is shown in Fig. 4c. Carbon rich/P-poor stoichiometry for C:P_{exp} is found in the ocean tropics and subtropics, being a function of both the large contribution of pico-phytoplankton productivity (Fig. 2) in these regions as well as the elevated C:P stoichiometry of the cyanobacteria (pro + syn + diaz) PFTs inhabiting these regions (Fig. 4d). Sinking particulate export C:P stoichiometry exhibits global maxima within the western subtropical gyres of each basin, reaching values of ~140 – 150 (Fig. 4c). C:P export stoichiometry is elevated elsewhere within the subtropics, ~120 – 140, near Redfieldian in the mid-latitudes and the Arctic Ocean, and depressed within the eastern equatorial Pacific, subpolar North Pacific, and the Southern Ocean, exhibiting values in the range ~90 – 105. The zonal mean C:P_{exp} is near the minimum imposed value of 90 for eukaryotic phytoplankton groups south of 50 °S and near ~100 north of 50 °N where the phytoplankton community is dominated by diatoms and other eukaryote groups (Fig. 2). Maximal zonal mean C:P_{exp} is found at subtropical latitudes exhibiting values of ~128 at 23 °S and a slightly higher value of ~133 at 26 °N in the northern hemisphere (Fig. 4d). Cyanobacteria phytoplankton (pro + syn + diaz), eukaryotic phytoplankton (peuk + nano + diat), and the total phytoplankton community C:P stoichiometry all exhibit similar latitudinal patterns with high latitude minima and subtropical maxima (Fig. 4d). Cyanobacteria phytoplankton C:P varies between ~96 – 223, eukaryotic phytoplankton C:P varies between ~90 – 117, with total phytoplankton C:P varying between ~90 – 202.

5. Discussion
Pico-phytoplankton were found to make their largest contribution to marine NPP within the tropical and subtropical ocean, reaching contributions of ~50 – 90% within the latitudinal range 45ºN – 45ºS (Fig. 2). In general, *Prochlorococcus* dominates NPP within the subtropical gyre regions, transitioning to dominance by pico-eukaryotes both equatorward and poleward of ~30º latitude to ~45º. *Synechococcus* follows a similar spatial pattern as *Prochlorococcus* but never dominates the phytoplankton community, with fractional contributions peaking at ~10 – 12% in the subtropical gyres. Diazotrophs, which are parameterized to represent the cyanobacteria lineages of *Trichodesmium* and *Crocosphaera* in the BEC model, make a smaller yet important additional contribution in these same low-latitude regions. The pico-phytoplankton and diazotroph groups within the PICO simulation were assigned the largest cellular C:P plasticity and higher minimum temperature thresholds for growth (Table 1) than the larger nano-phytoplankton and diatoms, with the pico-phytoplankton niche being realized within the nutrient-poor, high temperature waters of the low latitudes. The three pico-phytoplankton PFTs in PICO contributed to globally integrated NPP at a percentage of 16.6 ± 0.4% for *Prochlorococcus*, 3.9 ± 0.5% for *Synechococcus*, and 37.1 ± 0.6% for pico-eukaryotes, for a total pico-phytoplankton contribution of 57.6 ± 0.8% to marine NPP (Table 4). These estimates for global NPP attributable to *Prochlorococcus* and *Synechococcus* within PICO are approximately double and one-quarter, respectively, compared to an independent estimate based on data assimilative modeling of niche partitioning, (pro = 8.5%, syn = 16.7%, Flombaum et al., 2013), however both model estimates converge on a combined pro + syn contribution to marine NPP of ~20 – 25%. Nano-phytoplankton contribute 3.6 ± 0.7% of global NPP in PICO (likely an underestimate, e.g., Brewin et al., 2010; Juranek et al., 2020), concentrated in the Southern Ocean, the subpolar and Arctic Ocean, as well as the eastern tropical Pacific Ocean (Fig. 2d). Diatoms are the largest
contributor to global marine NPP in PICO, contributing 37.5 ± 0.5%, concentrated in the
subpolar and polar oceans as well as the equatorial Pacific upwelling (Table 4; Figure 2d). This
rate of diatom contribution to global NPP is consistent with the ~38-43% contribution estimated
from marine biogenic silica budgets (Nelson et al., 1995). Diazotrophs contribute 1.2 ± 0.04% to
marine NPP in PICO (Table 4), concentrated in the same tropical and subtropical regions as
pico-phytoplankton, especially the western tropical and subtropical Pacific Ocean (Fig. 2e).

<table>
<thead>
<tr>
<th>PFT</th>
<th>% of NPP</th>
<th>% of POC_{exp}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prochlorococcus</td>
<td>16.6</td>
<td>14.9</td>
</tr>
<tr>
<td>Synechococcus</td>
<td>3.9</td>
<td>12.6</td>
</tr>
<tr>
<td>Pico-eukaryotes</td>
<td>37.1</td>
<td>18.1</td>
</tr>
<tr>
<td>Nano-phytoplankton</td>
<td>3.6</td>
<td>9.9</td>
</tr>
<tr>
<td>Diatoms</td>
<td>37.5</td>
<td>31.2</td>
</tr>
<tr>
<td>Diazotrophs</td>
<td>1.2</td>
<td>13.4</td>
</tr>
<tr>
<td>Σ Pico-phytoplankton</td>
<td>57.6</td>
<td>45.6</td>
</tr>
</tbody>
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Table 4. Percentages of globally integrated marine NPP and POC_{exp,100m} attributable to each
PFT.

Ecosystem processing of the newly fixed organic matter by each phytoplankton PFT
contributes to the detrital particulate organic matter pools, which constitute the source terms to
the sinking export fluxes of POC_{exp} and particulate organic phosphorus (POC_{exp}) in the BEC
model. These processes include losses to the detrital pool upon grazing by zooplankton,
zooplankton contributions to the detrital pool as a result of grazing (e.g., fecal pellet production),
and aggregation and sinking of phytoplankton biomass upon cell mortality within the BEC
model (Moore et al., 2004). The parameter values governing these processes were prescribed in a
size-scaling manner across the six PFTs in PICO (Table 1), with the pico-phytoplankton
contributing the smallest to the particulate sinking fluxes, increasing with diazotrophs and nano-
plankton, to the largest contribution from diatoms. The fractions of pico-phytoplankton grazing
and zooplankton losses that accumulate within the particulate fluxes were prescribed in the range
0.07 – 0.33 (Table 1), at the upper end of estimates based on pigment biomarker analyses of the
sediment trap flux collected from the Sargasso Sea (Lomas & Moran, 2011). The higher
estimates for pico-phytoplankton detrital contributions were needed to export enough N and P
below the euphotic zone to prevent spurious accumulation of NO$_3^-$ and PO$_4^{3-}$ concentrations in
the upper layers of the subtropics. The nano-phytoplankton and diatom contributions to NPP are
<5% each over much of the subtropical regions in PICO (Fig. 2), which are biased low compared
to field observations (e.g., Nelson & Brzezinski, 1997; Brzezinski et al., 2011; Juranek et al.,
2020) and may potentially explain the low N and P export efficiency, partially satisfied by
prescribed elevated pico-phytoplankton detrital contributions in PICO. PFT-specific parameters
linked to the detrital contributions are held constant in PICO, however future formulations of the
BEC model could include parameterizations that are dependent on in situ temperature, PFT
growth rate, grazing rate, or nutrient limitation status among others.

Within PICO, pico-phytoplankton contribute nearly half (45.6 ± 0.3%) of the globally
integrated POC$_{exp,100m}$ flux through 100 meters when accounting for both direct (e.g.,
aggregation) and indirect (e.g., zooplankton processing of pico-produced organic matter)
contributions. Approximately 40% of the pico-phytoplankton total is contributed by pico-
eukaryotes (18.1 ± 0.2%), an additional third contributed by Prochlorococcus (14.9 ± 0.1%), and
a quarter contributed by Synechococcus (12.6 ± 0.1%) (Table 4). Pico-phytoplankton dominate
the phytoplankton contribution to carbon export across large areas of the tropics, subtropics, and
mid-latitudes (Fig. 4b), which is an emergent result from the added phytoplankton biodiversity,
representation of variable carbon to phosphorus cellular stoichiometry, and ecosystem processing
of the detrital pool mediated by zooplankton grazing. Perhaps counterintuitively, pico-
phytoplankton contribution to carbon export peaks in importance (~0.65 – 0.7), not where their
greatest environmental niche is realized, i.e., the high light, low nutrient waters of the
oligotrophic subtropical gyres, but at mid-latitudes, ~35 – 45°, driven largely by regional maxima
in C:P stoichiometry (Fig. 4c).

The estimate of pico-phytoplankton contribution to sinking carbon export in PICO
(~46%) is approximately 80% their combined contributions to global NPP (~58%), in partial
agreement with inversions of PFT-specific observational data from the equatorial Pacific and
Arabian Sea suggesting that pico-phytoplankton contribute to carbon export at rates comparable
to their contribution to NPP (Richardson & Jackson, 2007). Group specific C export fraction to
NPP fraction ratios, computed from Table 4, range from a minimum of ~0.5 to ~3.2, within the
range of the available group-specific estimates diagnosed from a range of field data, ~0.1 to ~4.5
(Richardson & Jackson, 2011; Stukel & Landry, 2010; Lomas & Moran, 2011; Fawcett et al.,
2011; Stukel et al., 2013), albeit at the higher end especially for Pro and Syn which are 0.9 and
3.2 in PICO and ~0.1 and ~0.1 to ~0.4 from the literature (Lomas & Moran, 2011; Fawcett et al.,
2011; Stukel et al., 2013), respectively. A notable outlier is the C export fraction to NPP fraction
ratio for diaz which is ~11 in PICO. Dissolved organic matter (DOM) contributes ~20-25% of
total organic carbon export production in the CESM (Letscher et al., 2015). We are unable to
provide an estimate of each PFT contribution to the DOM export flux, as DOM export is
spatially and temporally offset from its region of production, and the model does not track DOM
sourced from individual PFTs. Each PFT contributes a constant fraction of its newly produced
organic matter to the DOM pool; therefore, PFT contributions to DOM export should generally
track the proportions of PFT contributions to NPP (e.g., Table 4).

The regions located immediately along the poleward and western edges of the surface
western boundary currents of the subtropical gyre regions of each ocean basin exhibit elevated
POC_{exp,100m} fluxes on the order of ~50 – 90 g C m^{-2} yr^{-1} (Fig. 4a), approximately ~2-3 fold higher
as compared to the adjacent regions. These so-called ‘gyre recirculation regions’ of the surface
western boundary currents are characterized by strong eddying flow and enhanced lateral
nutrient concentration gradients that stand out as important regions for the delivery of a lateral
nutrient flux that dominates nutrient resupply to the subtropical gyres (Williams & Follows,
1998; Oschlies, 2002; Jenkins & Doney, 2003; Torres-Valdés et al., 2009; Palter et al., 2013;
Letscher et al., 2016; Yamamoto et al., 2019). The resolution of the BEC model employed in
PICO is non-eddy resolving, however it is able to simulate somewhat elevated rates of NPP (Fig.
1a) and associated POC_{exp,100m} (Fig 4a) along these subtropical to subpolar gyre flanks at the
western side. These gyre recirculation region biogeochemical features are driven by a larger
contribution of pico-eukaryotes and diatoms to simulated NPP (Fig. 2). These regions are also
where PFT cellular C:P stoichiometry transitions from C-rich/P-poor stoichiometry within the
subtropical gyres, to more Redfield-like C:P stoichiometry poleward (Fig. 3; Fig. 4c,d). Thus, the
magnitude of POC_{exp,100m} within the subtropical gyre recirculation regions identified as ‘POC_{exp}
hotspots’ by PICO may be sensitive to small changes to the nutrient flux stoichiometry which
drive cellular C:P stoichiometry. Further field-based study of the in situ phytoplankton
community composition and associated biomass C:P stoichiometry within these regions could
help confirm their regional ‘POC_{exp} hotspot’ status.
The PICO simulation builds from recent advances in a BEC model formulation that added variable C:P stoichiometry within the three traditionally simulated PFTs (small phytoplankton, diatoms, diazotrophs; Wang et al., 2019), by adding three pico-phytoplankton PFTs with variable C:P stoichiometry. As compared to the variable C:P, three PFT version of the BEC model (VAR-3; Wang et al., 2019), the PICO simulation exhibits ~1% higher globally integrated NPP (51.4 Pg C yr\(^{-1}\) vs. 50.8 Pg C yr\(^{-1}\)) and ~18% lower POC\(_{\text{exp},100m}\) (7.0 Pg C yr\(^{-1}\) vs. 8.5 Pg C yr\(^{-1}\)). Thus, given that both the three and six PFT, variable C:P formulations of the BEC model reproduce similarly-well, the surface to deep ocean spatial gradients in dissolved nutrient concentrations (Table 2), the pico-phytoplankton enabled (6p1z) BEC model predicts similar rates of NPP, with lower rates of POC\(_{\text{exp},100m}\), and lower C:P\(_{\text{exp}}\) per unit of phosphorus utilized within the euphotic zone as compared with the three PFT BEC model version (VAR-3; Wang et al., 2019). The geographical distribution of POC\(_{\text{exp},100m}\) is similar between the PICO and VAR-3 simulations, however VAR-3 simulated POC\(_{\text{exp},100m}\) is approximately double to triple that estimated by PICO within the subtropics to mid-latitudes (not shown), contributing to its overall ~18% higher estimate of global POC\(_{\text{exp},100m}\). The lower POC\(_{\text{exp},100m}\) estimate by the PICO model is likely attributable to the lower prescribed values of zooplankton grazing routed to the particulate organic pool, ‘graze_poc’ parameter, for the pico PFTs (graze_poc = 0.07 – 0.13; Table 1) versus the small phytoplankton PFT in Wang et al. (2019) (graze_poc = 0.3). Maximum C:P\(_{\text{exp}}\) is estimated within the ocean’s subtropical gyres (Fig. 4c) in both simulations, with PICO predicting slightly lower maximum C:P\(_{\text{exp}}\) within the Indo-Pacific subtropics, ~140 – 145 vs. ~180 in VAR-3 (not shown). PICO simulated C:P\(_{\text{exp}}\) generally agrees with the spatial pattern estimated from an inversion of dissolved inorganic carbon and phosphate data (Teng et al., 2014), with notable differences in C:P\(_{\text{exp}}\) magnitude. PICO minimum PFT C:P is set at 90 while
the Teng et al. (2014) inversion estimates minimum C:P$_\text{exp}$ of ~60 – 100 for the equatorial and subpolar regions. Maximal C:P$_\text{exp}$ in the subtropical North Atlantic is ~155 in PICO compared to ~355 by Teng et al. (2014). These comparisons demonstrate that estimates of the globally important fluxes of marine NPP and carbon export including its C:P stoichiometry vary on the order of ~10 – 50% between OBGC model formulations, highlighting the continued need for observational datasets to further constrain and validate model simulations of ocean biogeochemistry.

We are aware of two other ESMs to which addition of variable phosphorus to carbon ratios within the OBGC model has been performed as well as an additional global ocean biogeochemistry model. While nearly all ESMs participating in the CMIP5 suite predict declining marine NPP during the 21st century (Bopp et al., 2013), the GFDL-ESM2 which runs the variable C:P enabled TOPAZ2 OBGC model (Dunne, 2013), was the only ESM to predict a resilient rate of marine NPP throughout this century. Presumably the mechanism at play is one in which declining nutrient fluxes to the surface ocean with increasing upper ocean stratification are countered by a switch in phytoplankton community composition towards smaller, more nutrient plastic types and/or an increase in phytoplankton C:P that allows for maintenance of carbon based NPP rates at depressed phosphorus supply fluxes. The PICO simulation presented here supports this interpretation, exhibiting a marginally ~1% higher overall rate of marine NPP than its analogous three PFT, Redfield C:P sister simulation (RED-3; Letscher & Moore, 2015).

Results from a variable C:P OBGC model that added representation of pico-phytoplankton have been published (the PISCES-QUOTA model; Kwiatkowski et al., 2018). The model predicts similar 21st century declines in marine NPP (–8%) and POC$_\text{exp}$ (–14%) as the fixed (Redfield) stoichiometry ESMs that participated in CMIP5 (Bopp et al., 2013), while noting an increase in
carbon export efficiency per unit phosphorus, which increases by 4.5%. Both the TOPAZ2 OBGC model and PISCES-QUOTA parameterize variable C:P in the phytoplankton following the optimal resource cell allocation model of Klausmeier et al., (2004), so it is surprising that the two models yield divergent predictions for 21st century marine NPP (no decline in GFDL-ESM2 using TOPAZ2, decreasing in PISCES-QUOTA), that may be due to differences in model physics and the simulated nutrient supply to the surface. The global ocean biogeochemistry modeling study of Tanioka & Matsumoto (2017) in which variable C:P stoichiometry was represented in a two phytoplankton marine ecosystem model, found that predicted declines in global export production could be mitigated by ~3% relative to a Redfield stoichiometry model over the 21st century that the authors attributed to combined mechanisms of flexible cellular nutrient to carbon ratios and regional shifts in phytoplankton community composition.

6. Future Directions

A next step with the CESM-BEC model is to perform a future climate forced simulation using PICO parameterizations to investigate changes in 21st century marine NPP and export in the context of marine phytoplankton diversity and cellular nutrient plasticity. We also wish to point out that during development of the PICO simulation it was found that the model solution was very sensitive to the choice of the grazing rates for each of the six PFTs by the single zooplankton group. This single zooplankton group formulation was not well suited to act as the top-down control on the growth of six PFTs, with the model exhibiting behavior in which it was difficult to obtain mixed communities of phytoplankton everywhere in the ocean. Future improvement of the BEC model, now carried out within the Marine Biogeochemistry Library (MARBL) in CESM v2, should include the addition of explicit nanophytoplankton groups (e.g.,
coccolithophores) and additional zooplankton groups representing multiple size classes and
grazing formulations that will likely result in a more robust simulation of plankton growth,
grazing, and export dynamics. Currently, a version with four zooplankton and eight phytoplankton functional types is in development (8p4z).

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Open Research

The modified BEC model source code used to generate the PICO simulation can be found on GitHub here: https://github.com/rletscher/GBC_PICO. The 20-year annually averaged output file for PICO presented in this work is available by request from the corresponding author.

References


Cabrè, A., Marinov, I., & Leung, S. (2015). Consistent global responses of marine ecosystems to future climate change across the IPCC AR5 earth system models. Climate Dynamics, 45(5-6), 1253-1280.


plasticity on bulk particulate stoichiometry across ocean basins. *Communications Earth & Environment, 2*(1), 1-10.


Moore, J. K., Lindsay, K., Doney, S. C., Long, M. C., & Misumi, K. (2013). Marine ecosystem dynamics and biogeochemical cycling in the Community Earth System Model [CESM1 (BGC)]: Comparison of the 1990s with the 2090s under the RCP4.5 and RCP8.5 scenarios. *Journal of Climate, 26*(23), 9291-9312.


