

Trophic interactions with heterotrophic bacteria limit the range of *Prochlorococcus*

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Edited by Pablo Marquet, Ecology, Pontificia Universidad Catolica de Chile, Santiago, Chile; received July 8, 2021; accepted November 8, 2021

Prochlorococcus is both the smallest and numerically most abundant photosynthesizing organism on the planet. While thriving in the warm oligotrophic gyres, *Prochlorococcus* concentrations drop rapidly in higher-latitude regions. Transect data from the North Pacific show the collapse occurring at a wide range of temperatures and latitudes (temperature is often hypothesized to cause this shift), suggesting an ecological mechanism may be at play. An often used size-based theory of phytoplankton community structure that has been incorporated into computational models correctly predicts the dominance of *Prochlorococcus* in the gyres, and the relative dominance of larger cells at high latitudes. However, both theory and computational models fail to explain the poleward collapse. When heterotrophic bacteria and predators that prey nonspecifically on both *Prochlorococcus* and bacteria are included in the theoretical framework, the collapse of *Prochlorococcus* occurs with increasing nutrient supplies. The poleward collapse of *Prochlorococcus* populations then naturally emerges when this mechanism of “shared predation” is implemented in a complex global ecosystem model. Additionally, the theory correctly predicts trends in both the abundance and mean size of the heterotrophic bacteria. These results suggest that ecological controls need to be considered to understand the biogeography of *Prochlorococcus* and predict its changes under future ocean conditions. Indirect interactions within a microbial network can be essential in setting community structure.

Prochlorococcus | biogeography | trophic interactions

P*rochlorococcus* is the world’s smallest photosynthetic organism and is found ubiquitously in the global ocean equatorward of about 45° latitude (1). In regions where it occurs, *Prochlorococcus* tends to numerically dominate the photosynthetic community. Understanding what sets its biogeography is critical because of its important impact on global primary production and biogeochemical cycling. A striking feature of *Prochlorococcus* biogeography is its absence from higher latitudes (2) and regions of high chlorophyll concentrations (3), and, in particular, the rapidness of decline (Fig. 1). This collapse has become even more clear with new high spatial resolution SeaFlow data (4) (Fig. 1B). This “collapse” at high latitudes is often ascribed to temperature sensitivity: *Prochlorococcus* fails to grow below ~8 °C in the laboratory (5). Indeed, global distributions of *Prochlorococcus* appear to be well captured by simple parametric models including only temperature and light (photosynthetically available radiation) (2). This has led some authors (2, 6) to conclude that the biogeography of *Prochlorococcus* will expand poleward under climate change–induced ocean warming.

We explore this temperature-driven hypothesis of the collapse of *Prochlorococcus* by examining observations from several cruises in the northeast Pacific Ocean (Fig. 1). We plot an estimate for the latitudinal location of the collapse of *Prochlorococcus* populations for each of four cruises (7, 8), as well as the temperature at that location. Three things are clear: the decrease in cell abundance can happen over a short distance; the location of the collapse moves by at least 6° latitude over the year; and the collapse occurs at different temperatures. This final point is the

most critical. The temperature of the cell abundance transition for *Prochlorococcus* appears to span ~7° of temperature, some of which are within *Prochlorococcus*’ optimal temperature range (9). This suggests that an additional factor contributes to the collapse of *Prochlorococcus* in this region.

We pose that an ecological interaction might be important, leveraging the knowledge that *Prochlorococcus* and similarly sized phytoplankton and heterotrophic bacteria share common predators (10–14). We hypothesize that this indirect interaction leads to the decline of *Prochlorococcus* as heterotrophic bacterial populations increase in regions of higher nutrient supply. Indeed, although temperature effects could not be excluded, heterotrophic bacteria have been invoked in this fashion as a possible explanation for observed decreases in *Prochlorococcus* indicator pigments at high chlorophyll concentrations (3). In ecological theory, this mechanism is known as “apparent competition” (15), defined as the “negative indirect interactions between victim species that arise because they share a natural enemy” (16). Here we refer to the mechanism as “shared predation” since the term “competition” is strongly associated with nutrient limitation and “bottom-up” controls in plankton ecology.

We provide a size-based theory for how shared predation works in this system. In this model, a critical size emerges that separates the size spectrum into heterotroph-dominated and

Significance

Prochlorococcus is the smallest and most abundant photosynthetic organism on Earth and is thought to be confined to low-latitude regions by its requirement for warm waters. Latitudinal transects in the North Pacific, however, demonstrate that the poleward decrease of this species occurs across a wide range of temperatures. An additional mechanism is likely required. We use theory, computational models, and additional observational data to suggest that the poleward decrease is caused by an ecological interaction: a shared predator which consumes both *Prochlorococcus* and similar-sized heterotrophic bacteria. Understanding the fate of this organism requires a knowledge of the interconnected ecosystem of other organisms, where both direct and indirect interactions control community structure.

Author contributions: C.L.F. and S.D. designed research; C.L.F. and S.D. performed research; E.Z. contributed new reagents/analytic tools; C.L.F., S.D., and F.R. analyzed data; and C.L.F., S.D., F.R., D.C., E.V.A., and M.J.F. wrote the paper.

The authors declare no competing interest.

This article is a PNAS Direct Submission.

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This article contains supporting information online at <https://www.pnas.org/lookup/suppl/doi:10.1073/pnas.2110993118/-DCSupplemental>.

Published January 4, 2022.

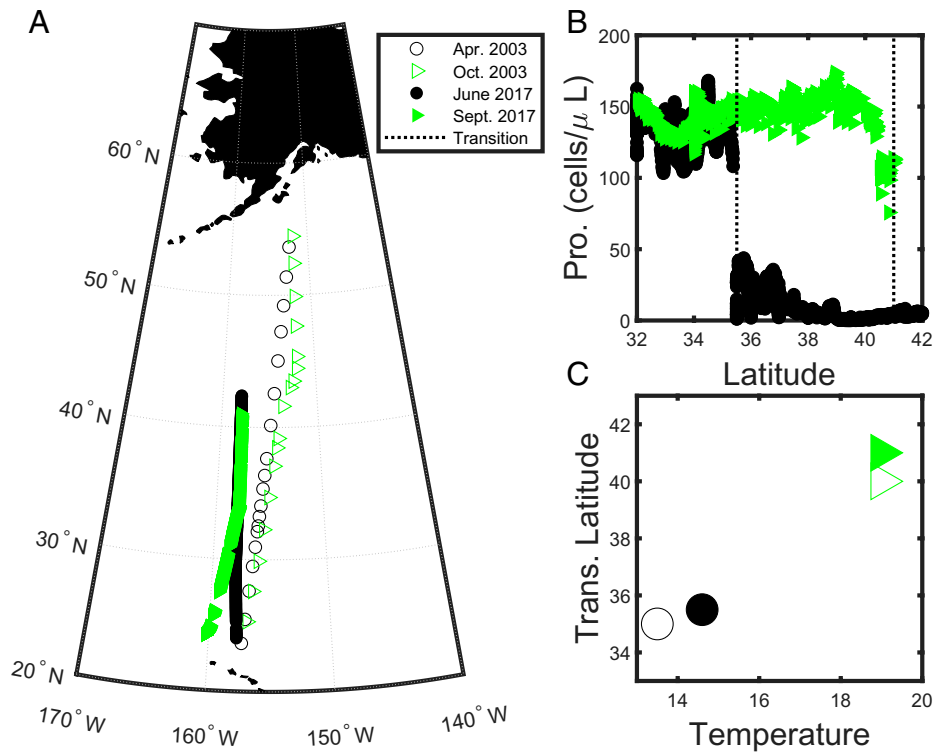


Fig. 1. Cruise transect data: The transition from relatively stable to very few *Prochlorococcus* occurs at different latitudes over the seasons, and at different temperatures. (A) Cruise tracks for four latitudinal cruises across the northeast Pacific Ocean from April 2003, October 2003, April 2016, and September 2017. Black circles represent spring, and green triangles represent fall. Filled symbols are measurements from cruises associated with the Simons Collaboration on Ocean Processes and Ecology (SCOPE)-Gradients (7) cruise campaign, and open symbols are associated with cruises as part of the Comprehensive Oligotrophic Ocean Knowledge-Biogeochemical Observations Oahu-Kodiak (COOK-BOOK) program (8). (B) Mixed layer abundances of *Prochlorococcus* (cells per microliter) along the transect of the two cruises in June 2017 and September 2017. The dashed vertical lines indicate the latitude of the abrupt drop in cell numbers. (C) The corresponding latitude (y axis) and temperature (x axis) where the transition occurs.

photoautotroph-dominated sections. We show that this critical size increases as a function of resource supply, demonstrating how *Prochlorococcus*-like cells could be excluded in favor of heterotrophic bacteria in regions with sufficiently high resource supply. This theory therefore contains both top-down (e.g., grazers) and bottom-up (e.g., rate of supply of limiting nutrient) components. After showing how this theory works in simplified models, we implement it in a fully coupled global ecosystem model. The transition naturally emerges, solving outstanding issues with our ability to simulate and interpret *Prochlorococcus* globally. Finally, we return to transect data, showing that the trends in *Prochlorococcus* and heterotrophic bacterial abundance as well as their size agree with our predictions.

Theoretical Framework for Plankton Community Size Structure

Previous theoretical frameworks to explore the controls on size structuring of phytoplankton assemblages have highlighted the important interaction between predator and prey, and the supply rate of the limiting resource (3, 17–20). To illustrate these ideas, we first consider the following set of equations (schematically shown with black arrows in Fig. 2A):

$$\frac{dR_p}{dt} = - \sum_j \mu_{pj} P_j + S_{R_p} \quad [1]$$

$$\frac{dP_j}{dt} = +\mu_{pj} P_j - g_j P_j Z_j - m_p P_j \quad [2]$$

$$\frac{dZ_j}{dt} = +\gamma g_j P_j Z_j - m_z Z_j \quad [3]$$

Here R_p is the limiting resource for the phytoplankton (e.g., nutrient such as nitrate), P_j is the biomass of the phytoplankton j , and Z_j is the zooplankton that grazes on the j th phytoplankton. The resource is supplied to the system at rate S_{R_p} . Plankton growth (μ_{pj}) is considered a function of a maximum growth rate, μ_{pj}^{max} , a half-saturation constant κ_{pj} , and the limiting resource in a Monod kinetics,

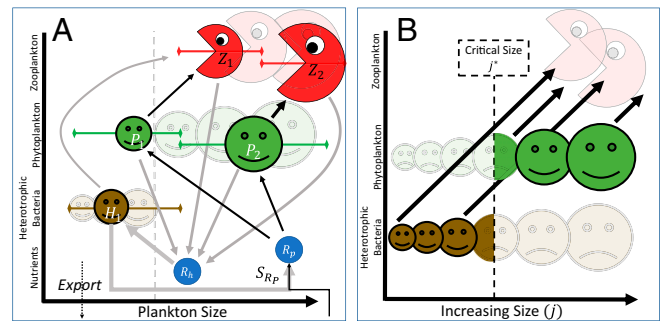


Fig. 2. Ecosystem schematics. (A) Heterotrophic bacteria, H_i , phytoplankton, P_j , and zooplankton, Z_j , along with their required resources R_i ($i = p, h$) are shown with arrows representing the elemental flow. The earlier size-based PZ model (Eqs. 1–3) formulation with two sizes is represented by the black arrows. The addition of a heterotroph, H_1 and organic matter resource R_h shown with gray arrows (i.e., the system represented by Eqs. 6–10), allows for the exclusion of P_1 at large enough resource supplies (S_{R_p}). (B) The mechanism for exclusion can be described in terms of a critical size class, j^* , separating a heterotroph-dominated from an autotroph-dominated section of the size spectrum. This critical size increases as a function of inorganic resource supply.

$$\mu_{pj} = \frac{\mu_{pj}^{max} R_p}{R_p + \kappa_{pj}}$$

Zooplankton graze with a per capita rate g_j , and assimilate a fraction γ of the prey biomass. Both plankton have a linear loss rate (m_p, m_z). Here we consider a simple food chain with a range of phytoplankton sizes ($j = 1$ is the smallest) that are preyed on by a size range of zooplankton ($j = 1$ is the smallest). We assume that the smallest phytoplankton has the highest affinity, $\mu_{p1}^{max}/\kappa_{p1}$, for the resource, the second smallest has the next highest affinity, and so on, as suggested empirically (21) (see *Materials and Methods*).

Although highly simplified, these equations provide a clear illustration of the mechanisms at play, especially when we consider steady state (see *Materials and Methods*, Eqs. M1–M3). In particular, the nonzero steady-state solution for resource and phytoplankton are (where * indicates the steady state solution)

$$R_{pj}^* = \frac{\kappa_j(m_p + g_j Z_j^*)}{\mu_{pj}^{max} - (m_p + g_j Z_j^*)} \quad [4]$$

$$P_j^* = \frac{m_z}{\gamma g_j} \quad [5]$$

Following the terminology of resource competition theory (22, 23), the phytoplankton type with the lowest subsistence resource concentration R_{pj}^* (Eq. 4) is the fittest. In the absence of grazing, the smallest phytoplankton with highest affinity has the lowest R^* and will outcompete all others (Fig. 3A). This occurs when resource supplies are very low. As one transitions to regions of higher and higher nutrient supplies, the grazer biomass increases (when only one zooplankton exists, $Z_1^* = \frac{\gamma S_{Rp}}{m_z} - \frac{m_p}{g_1}$; see *Materials and Methods*, Eq. M4), and, subsequently, R_{p1}^* increases. It eventually reaches the value of R_{p2}^* (in the absence of grazing), and the two smallest phytoplankton can coexist (second vertical dashed line in Fig. 3A). The smallest phytoplankton biomass, P_1 , is capped as given in Eq. 5. Additional resource supply goes into the biomass of the next largest phytoplankton (P_2) and its paired zooplankton (Z_2) and so on for additional size classes. This inclusion of larger and larger phytoplankton transitioning from oligotrophic to increasingly eutrophic (higher R^*) waters is a well-known phenomenon observed in the world's oceans (24, 25).

However, in the simplified theory, P_1 persists with biomass $m_z/\gamma g_1$ (Eq. 5) no matter the resource supply. Thus, there is no mechanism in this theory to explain the collapse of *Prochlorococcus* as shown in Fig. 1.

Shared Predation between Prochlorococcus and Heterotrophic Bacteria Causes the Collapse. We introduce into this framework heterotrophic bacteria, H_j , which are consumed along with similar size phytoplankton, P_j , by zooplankton, Z_j (Fig. 2A; gray lines indicate the additional flows of matter),

$$\frac{dR_p}{dt} = - \sum \mu_{pj} P_j + S_{Rp} \quad [6]$$

$$\frac{dP_j}{dt} = +\mu_{pj} P_j - g_j P_j Z_j - m_p P_j \quad [7]$$

$$\frac{dH_j}{dt} = +\mu_{hj} H_j - g_j H_j Z_j - m_h H_j \quad [8]$$

$$\frac{dZ_j}{dt} = +\gamma g_j (P_j + H_j) Z_j - m_z Z_j \quad [9]$$

$$\frac{dR_h}{dt} = - \sum \delta \mu_{hj} H_j + S_{Rh} \quad [10]$$

Heterotrophic bacteria are assumed to consume detrital matter, R_h ,

$$\mu_{hj} = \frac{\mu_{hj}^{max} R_h}{R_h + \kappa_{hj}}$$

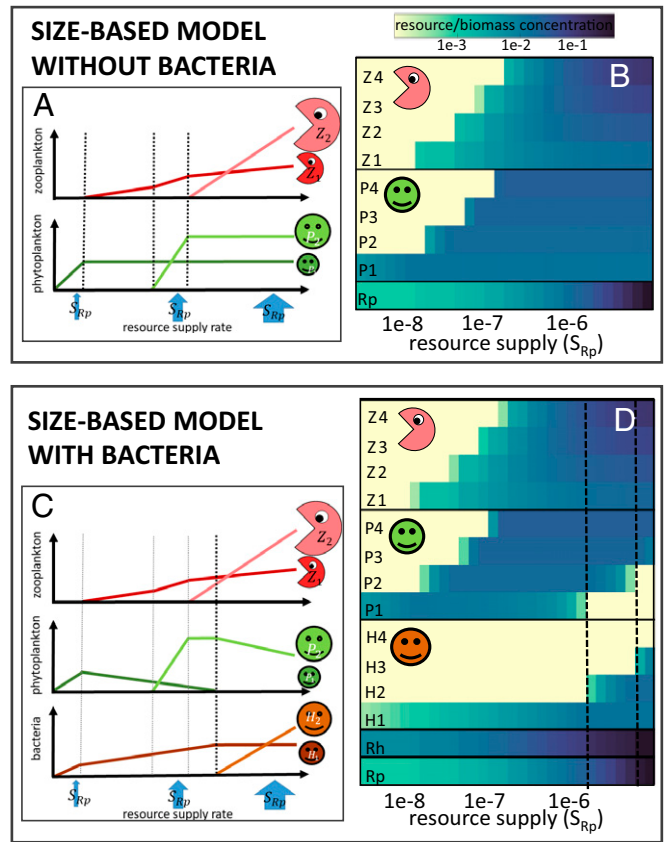


Fig. 3. (A and C) Schematic of the theoretical steady-state results for different resource rates (S_R) for (A) simple R_p, P_j, Z_j size structured model (Eqs. 1–3) and (C) the more complex model including size classes of heterotrophic bacteria H_j (Eqs. 6–10). (B and D) Solutions for zero-dimensional numerical model for different resource supply rates for (B) Eqs. 1–3 and (D) Eqs. 6–10. Solutions are steady state for low resource supply rates, and are averages over several cycles of predator–prey oscillations that occur for higher resource supply rates in the simplified equations.

The constant μ_{hj}^{max} is the maximum growth rate and κ_{hj} is the detritus half-saturation rate for bacteria j . Only a small fraction of the detrital matter taken up by heterotrophic bacteria goes into biomass growth (26), and the excess matter ($\delta - 1$, where $\delta > 1$) is remineralized to the inorganic resource, R_p . Detrital matter is supplied by the ecosystem loss terms $S_{R_h} = f(m_p P_j, m_z Z_j, m_h H_j)$, although it could also include some of the nonconsumed grazed plankton. For ease of illustration, we do not express this term explicitly, but do note that S_{R_h} increases with increased total biomass.

The steady-state subsistence solutions (see *Materials and Methods*, Eqs. M5–M9) for inorganic resource are the same as in the simpler model (Eq. 4), but the analogous equation to Eq. 5 has a significant difference,

$$R_{pj}^* = \frac{\kappa_{pj}(m_p + g_j Z_j^*)}{\mu_{pj}^{max} - (m_p + g_j Z_j^*)} \quad [11]$$

$$P_j^* = \frac{m_z}{\gamma g_j} - H_j^* \quad [12]$$

The predator-mediated, “apparent” competition manifests in Eq. 12 where P_j^* declines as H_j^* increases (and vice versa). By having a shared predator, similarly sized phytoplankton and heterotrophic bacteria negatively impact each other.

The steady-state solution for detrital matter has a similar form to the inorganic resource,

$$R_{hj}^* = \frac{\kappa_{hj}(m_h + g_j Z_j^*)}{\mu_{hj}^{max} - (m_h + g_j Z_j^*)}. \quad [13]$$

If we assume [as suggested empirically (27, 28), and see *Materials and Methods*] that smaller bacteria have a higher affinity for dissolved organic matter than larger bacteria, these equations suggest an increase in size classes of bacteria across a gradient of increasing supply of resource, S_{R_p} (schematically shown in Fig. 3C). Z_j concentrations increase with higher inorganic and organic resource supply (in the case of a single size class $Z_1^* = \frac{\gamma}{m_z}(S_{R_p} + \frac{S_{R_h}^*}{\delta}) - \frac{m_p}{g_1}$; see *Materials and Methods*, Eq. M9). Thus, with increasing resource supply, R_h^* increases until another size class of bacteria can be supported. Instead of the bacteria being capped at a constant value, it is the sum of the similar-sized phytoplankton and bacteria (j) that is held at $m_z/\gamma g_j$ (Fig. 3B and Eq. 12). As detrital matter accumulates in regions with higher inorganic resource supply, heterotrophic bacterial biomass increases, to the detriment of the similar-sized phytoplankton. As H_j approaches $m_z/\gamma g_j$, the phytoplankton are excluded (black dashed vertical line in Fig. 3C). This provides an ecological mechanism for the demise of a size class with increased resource supply.

A Size-Based Ecological Transition from Heterotrophs to Autotrophs.

The steady-state solutions for Eqs. 7 and 8 provide two constraints on the zooplankton biomass (*Materials and Methods*, Eq. M7). To match these constraints simultaneously (if we assume that the death rate constants for H_j and P_j equate, $m_h = m_p$), coexistence in steady state occurs only when the growth rates of the similar size bacteria and phytoplankton are equal,

$$\frac{\mu_{pj}^{max} R_p^*}{R_p^* + \kappa_{pj}} = \frac{\mu_{hj}^{max} R_h^*}{R_h^* + \kappa_{hj}}. \quad [14]$$

This suggests that a tight coupling between accumulation of detrital matter and supply of inorganic resources is necessary for the coexistence of the two prey. For how many sizes classes, (j), is this condition met?

We have assumed here that the growth rate constants μ^{max} and κ are functions of the organism type (subscripts p or h) and increasing size, j . There is strong theoretical and empirical evidence to suggest that the size-dependent growth rate constants μ^{max} and κ for phytoplankton can be expressed as power law functions of size (21, 29), and, for simplicity, here we assume the growth of heterotrophic bacteria scales similarly (see *Materials and Methods*). Given these scalings, we can write μ^{max} and κ as separable functions; a function of size ($f_\mu(j)$, $f_\kappa(j)$, which are independent of type) multiplied by type-dependent constants (Ψ_c^μ or Ψ_c^κ , $c \Rightarrow p$ or h). We expand Eq. 14 with these functions,

$$\frac{\Psi_p^\mu f_\mu(j) R_p^*}{\Psi_p^\kappa f_\kappa(j) + R_p^*} = \frac{\Psi_h^\mu f_\mu(j) R_h^*}{\Psi_h^\kappa f_\kappa(j) + R_h^*}, \quad [15]$$

which can be solved for the size-dependent portion of the half-saturation function,

$$f_\kappa(j) = \frac{(\Psi_h^\mu - \Psi_p^\mu) R_p^* R_h^*}{\Psi_p^\mu \Psi_h^\kappa R_p^* - \Psi_h^\mu \Psi_p^\kappa R_h^*}. \quad [16]$$

The half-saturation constant is a monotonically increasing function of size (see *Materials and Methods*), and the right-hand side of Eq. 16 is independent of size. Thus, there is, at most, a single size class (i.e., single j , which we will refer to as j^*) which contains both plankton types for any fixed combination of R_p^* and R_h^* (set by the resource supply, S_{R_p}). As the derivative d/dR_p^* of the right-hand side of Eq. 16 is negative under our conditions, size classes of phytoplankton will be systematically eliminated with increasing nutrient supplies.

Numerical Simulations Support Theory

The theory we lay out above suggests that smaller phytoplankton will be eliminated with higher resource supply due to shared predation with similar-sized heterotrophic bacteria. The theory also suggests that only one sized pair of bacteria and phytoplankton can coexist at any resource supply rate. The number of size classes of the eliminated phytoplankton increases with higher resource supply. Here we demonstrate these findings with two sets of numerical simulations.

The first model setup encodes the exact equations as laid out in *Theoretical Framework for Plankton Community Size Structure* (see *Materials and Methods*; parameters are provided in *SI Appendix, Table S1*), and we solve the solutions for parcels of water (we will call this the zero-dimensional model setup) with different resource supply rates (S_{R_p}). First, we show the solutions for Eqs. 1–3 for the size-based model including only phytoplankton and zooplankton (Fig. 3B) represented by the black arrows in Fig. 2A for four size classes. Nutrient affinity decreases with size, although note that we employ a unimodal distribution for maximum growth rates (30), such that the smallest type has a low rate as observed for *Prochlorococcus* (9) (see *Materials and Methods*). The anticipated increase in the number of coexisting phytoplankton size classes with increasing resource supply is evident. The resource supply at which any size class of phytoplankton j can coexist is higher than the resource supply rate where the smaller-size class grazer ($j - 1$) is supported, indicating that grazer control of a smaller phytoplankton size class allows the next larger phytoplankton size class to survive. Once the j th grazer can survive, the prey j has its biomass capped. These results support the finding of previous size-based models (e.g., refs. 17 and 19).

In Fig. 3D, we show the results for the system extended to include heterotrophic bacteria and detrital matter (Eqs. 6–10, additional gray arrows in Fig. 2A). Again, we see the increase of coexisting phytoplankton size classes and their grazers with increasing resource supply (S_{R_p}). Additionally, several size classes of heterotrophic bacteria can coexist at larger resource supply rates. Increasing resource supply leads to increased detrital supply, which, in turn, leads to increased bacterial biomass in the j th size class at the expense of the j th phytoplankton. At a critical resource supply rate (vertical black dashed lines in Fig. 3D), when the heterotrophic bacteria reach its maximum biomass ($m_z/\gamma g_j$), the j th phytoplankton is eliminated. Additionally, we see that only one size class of phytoplankton and bacteria can coexist. Moving along the axis of resource supply rate, we see each new size class $j + 1$ of bacteria emerge only after the smaller phytoplankton j no longer coexists.

This computational examination of Eqs. 6–10 provides evidence that indirect competition due to a shared predator can cause the collapse of a phytoplankton size class. However, this setup is highly simplistic, and we now put these ideas in the context of the *Prochlorococcus* collapse seen in Fig. 1. First, we note that the smallest heterotrophic bacteria (e.g., SAR11) are smaller than *Prochlorococcus*. For instance, SAR11 have been found to have cell diameters between about 0.3 and 0.45 μm (28), while *Prochlorococcus* has been observed with diameters varying between about 0.45 and 0.8 μm (5, 9, 31, 32). Additionally, the carbon cell quota of SAR11 has been shown to be ~ 10 times less than for *Prochlorococcus* (33). We repeat the experiment seen in Fig. 3D with the smallest size class only consisting of bacteria, and find, qualitatively, the same results (*SI Appendix, Fig. S1A*). Nevertheless, the model described by the set of Eqs. 6–10 is extremely simplified, neglecting the many additional complexities of the real ocean, including influences of seasonality, light, temperature, multiple limiting nutrients, more complex food web dynamics (e.g., inclusion of viruses), and constant mixing and transport by the three-dimensional fluid flow. We now consider a second set of numerical experiments using a more complex three-dimensional global biophysical model.

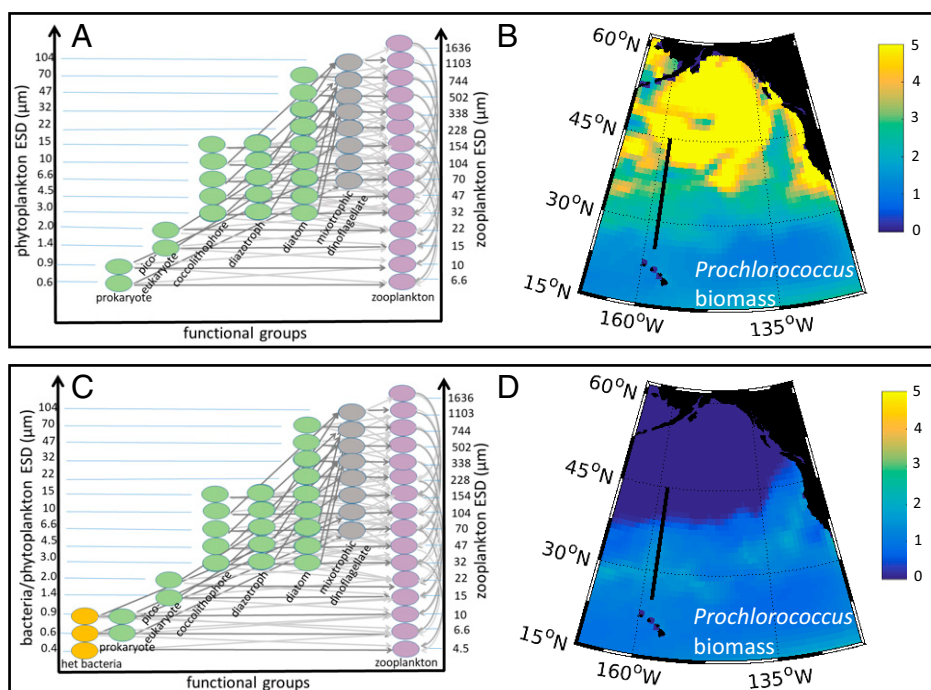


Fig. 4. Global biophysical model. (A and C) Schematic of the ecosystem structure with y axis representing the size of organisms and x axis representing their functional type. In A, remineralization is parameterized as a rate, while, in C, three heterotrophic bacteria size classes are explicitly included in the food web. (B and D) Modeled surface biomass (milligrams C per cubic meter) of the model *Prochlorococcus* analogs for the case with no explicit bacteria (B) and the case with bacteria (D).

We use a modified version of the size-based numerical ecosystem model used in ref. 34. The model includes 31 phytoplankton types (Fig. 4A and C) covering 15 size classes from 0.6 μm to 104 μm equivalent spherical diameter (ESD). The phytoplankton are additionally split into functional groups including diatoms, coccolithophores, prokaryotes, picoeukaryotes, diatoms, and mixotrophs. The smallest phytoplankton (with ESD of 0.6 μm) is an analog of *Prochlorococcus*. Phytoplankton growth (and sinking) parameters are allometrically defined similarly to the simple zero-dimensional model shown in Fig. 3, but with functional group specifics (see ref. 35, *Materials and Methods*, and *SI Appendix*, Table S1). Growth rates are also a function of multiple potential limiting nutrients (dissolved inorganic nitrogen, phosphate, iron, and silicic acid), spectral light, photoacclimation, and temperature. The model includes a size range of zooplankton that graze on plankton 5 to 15 times smaller than themselves, but, preferentially, 10 times smaller (36–38). We additionally include three heterotrophic bacteria, ranging from smaller than to the same size as the two smallest phytoplankton, following ref. 26. This complex ecosystem, along with inorganic, dissolved, and particulate material, is advected and mixed within a three-dimensional global ocean model (see *Materials and Methods*).

We run two experiments: one where we explicitly include the heterotrophic bacteria and another where we exclude the bacteria and approximate remineralization as a rate-specific function of the detrital matter's concentration. Both models capture similar (and reasonable) patterns of total chlorophyll *a* concentrations (Chl), size distribution of phytoplankton, and functional groups (*SI Appendix*, Figs. S2 and S3). The patterns of high and low total Chl are realistic, including the sharp increase in Chl and size classes in the transition from subtropical to subpolar conditions in the North Pacific covered by the shipboard transects (Fig. 1 and *SI Appendix*, Fig. S2). In the simulation with explicit bacteria, the smallest bacteria type is ubiquitous, with an increase in larger types also occurring in the transition zone (*SI Appendix*, Fig. S4).

In the model without explicit bacteria, *Prochlorococcus* analogs continue to exist in poleward regions (Fig. 4B), in strong contrast to the version with explicit bacteria (Fig. 4D). With the explicit bacteria, *Prochlorococcus* biomass is relatively uniform in the subtropical regions, with a sharp decrease in the midlatitudes similar to that suggested by the observations. The collapse in the population occurs slightly poleward from where the total phytoplankton biomass increases dramatically, coincident with the grow-in of the larger heterotrophic bacteria (*SI Appendix*, Fig. S4). Over this region (several degrees of latitude, shifting between 30°N and 40°N seasonally), the environment shifts from oligotrophic conditions with relatively uniform nutrient supplies (i.e., S_{R_p} = constant; *SI Appendix*, Fig. S5) via remineralization to a region of seasonally large fluxes of nutrients from upwelled waters at higher latitudes (see, e.g., ref. 39). This increase in resource supply rates (at least seasonally) increases dramatically over a very short distance (i.e., the x axis, S_{R_p} , of Fig. 3D is collapsed over just a few degrees of latitude). This sharp increase in resource supply rate promotes the sharp increase in total phytoplankton biomass (i.e., more size classes), increases supply of organic matter, and thus allows the coexistence of the larger bacteria size class (*SI Appendix*, Fig. S4). Shared predation with these larger heterotrophic bacteria driven by the 6.6- μm grazer leads to the collapse of the *Prochlorococcus* analog. The simulation without explicit bacteria shows no such collapse (Fig. 4B). These results suggest that, even in a much more realistic environment (multiple limitations on growth, lack of steady state) and far more complex food web dynamics, the insight from the theoretical framework of shared predation can explain the collapse of *Prochlorococcus*.

Data Comparison and Summary

We have developed a theory that combines size-based predator-prey interactions (19, 40), including shared predation [known as “apparent competition” in classical ecological theory (15)], to

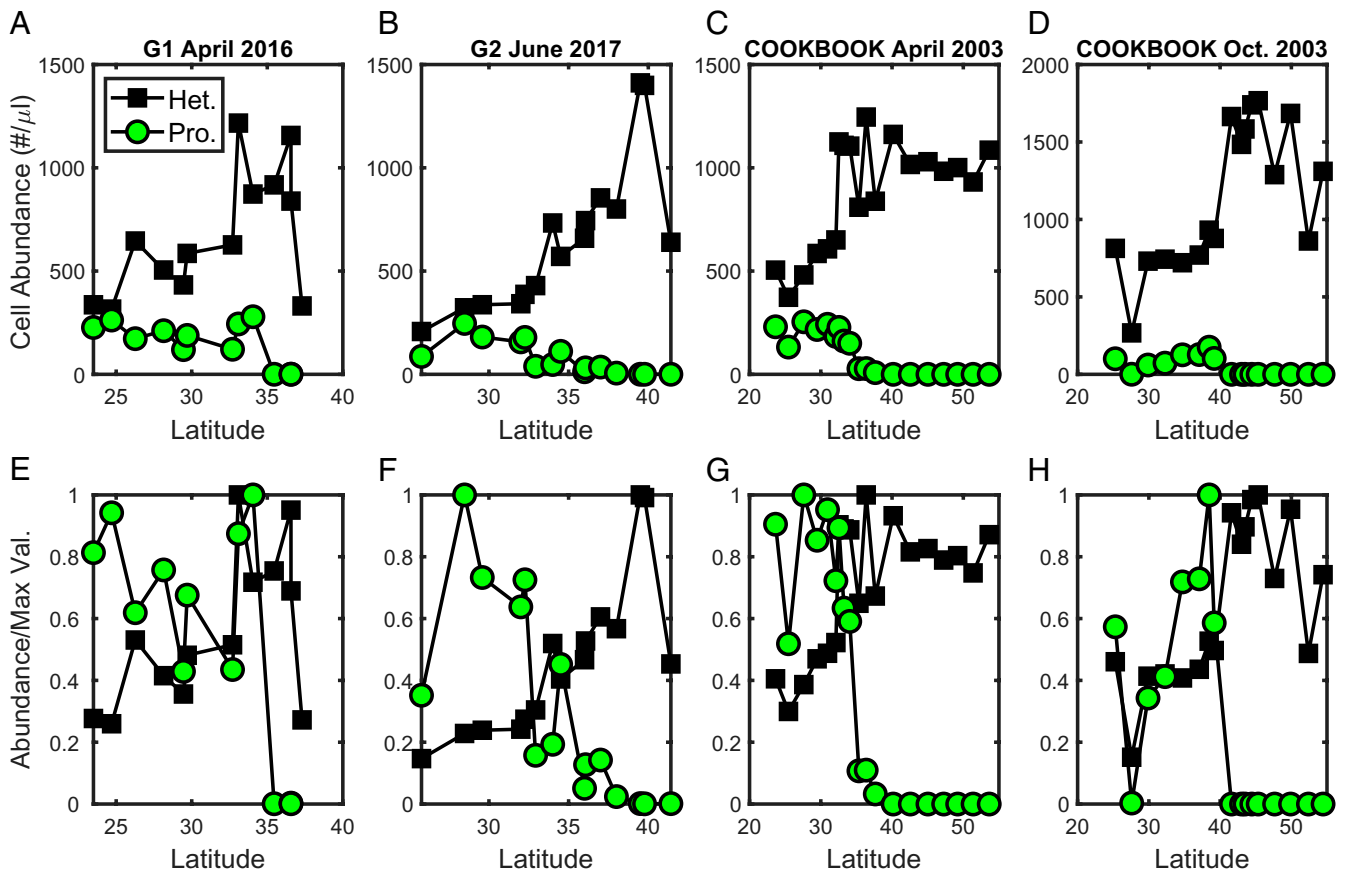


Fig. 5. Observed surface data for *Prochlorococcus* (green circles) and heterotrophic bacteria (black squares). (A–D) Cell abundance (cells per microliter) across four different transects spanning 2003–2016. Cruises in B–D are the same as in Fig. 1. Additional cruise information is available for the SCOPE-Gradients cruises 1 and 2 (A and B; see ref. 7) and the COOK-BOOK cruises (C and D; see ref. 8). (E–H) Cell abundance normalized to the maximum value of that type on that transect (above, A–D).

consider the controls on community structure. The theory, by including both bottom-up (e.g., supply of nutrients) and top-down (e.g., grazing pressure) processes, lays out a set of predictions that can be evaluated with observed data from latitudinal surface transects in the North Pacific. The simplest prediction from apparent competition is that the abundance of heterotrophic bacteria should increase as the abundance of *Prochlorococcus* decreases, clearly seen in the zero-dimensional simulation results (Fig. 3D). However, growth rates in the real ocean will be affected by a number of factors other than a single resource, including temperature and, in the case of phytoplankton, light, while grazing rarely manifests as food chains. The global biophysical models suggest that a more complex marine food web with multiple limitations to growth, and subject to physical advection and mixing (i.e., never in steady state), provides similar results (Fig. 4D and SI Appendix, Fig. S4). We also see that this prediction is consistent with the surface data from four different cruise transects (Fig. 5). These transects are from different times of year and span almost 15 y. We find that the relative cell abundance of heterotrophic bacteria undergoes a strong increase coincident with the strong decrease in *Prochlorococcus*. Across this transect, nutrient supply rates increase poleward (as shown in the model; SI Appendix, Fig. S5). Surface nutrient concentrations also increase (SI Appendix, Fig. S7) (39), as suggested by increasing R^* . The increased nutrient supply leads to higher primary production and higher biomass of larger phytoplankton (Fig. 6A) poleward. As a result, more organic matter is produced, which, in turn, supports more heterotrophic bacteria. The theory of shared predation predicts such a pattern; as both the mutual grazer

and the bacteria population increase, the loss of *Prochlorococcus* increases until it no longer coexists.

The theory also suggests that only a single size class of mutual prey can coexist, and that this critical size, j^* , is a function of resource supply. This result is illustrated in the more abstract zero-dimensional model (Fig. 3D). Allowing for the different

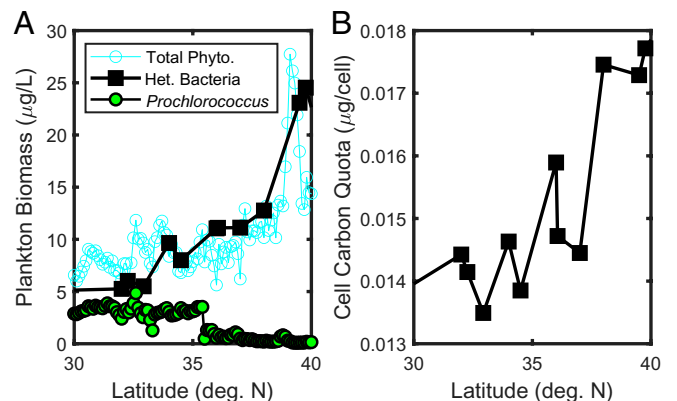


Fig. 6. Observed surface data from the June 2017 north–south transect near 158°W (MGL1704, SCOPE-Gradients 2; also see Fig. 1). (A) Total estimated carbon biomass (micrograms C per liter; see Materials and Methods) for heterotrophic bacteria (black squares), *Prochlorococcus* (green circles), and total phytoplankton (blue open circles). (B) Mean cell mass (micrograms C per cell) of heterotrophic bacteria.

ranges in sizes of phytoplankton and heterotrophic bacteria, our global model results also suggest that it is, in fact, increases in larger bacteria that force the collapse of *Prochlorococcus* (SI Appendix, Fig. S4B). The smallest heterotrophic bacteria are ubiquitous in the model ocean, similar to the wide distribution of SAR11 (28). It is only as the larger bacteria grow, supported in regions of higher plankton biomass and nutrient supply rates, that *Prochlorococcus* is eliminated. We find that this additional component of the theory/model is also consistent with observations from cruise results. Not only does the biomass of bacteria increase just south of the collapse of *Prochlorococcus* (Fig. 6A), but the average cell size of the bacteria also increases significantly moving poleward (Fig. 6B). The cell size increase closely follows the increase in total phytoplankton biomass, and hence the increase in production of organic matter and supply of inorganic resource.

Here we have focused on the decrease of *Prochlorococcus* abundance moving poleward in surface waters of the North Pacific, but similar patterns of declining *Prochlorococcus* cell abundances have also been observed in the Atlantic (5, 41) in both northern and southern gyre transitions. Coincidentally, increasing bacterial cell abundances have been observed across these transitions (41). Thus, we believe that this theoretical understanding is not restricted to the North Pacific and has a larger application to understanding global *Prochlorococcus* distributions.

We do not preclude a role for temperature, as *Prochlorococcus* does indeed fail to grow in cold water (5). *Prochlorococcus* also has a low maximum growth rate relative to eukaryotic phytoplankton (9), consistent with its very small size (30). It will have a growth disadvantaged relative to others, particularly with a combination of higher nutrients and low temperatures (42). However, its ability to avoid predation due to its small size remains, which is why this mechanism fails to cause the collapse in models (as in Fig. 4). Our work suggests that shared predation may be a leading process in determining the spatial distribution of *Prochlorococcus*. In addition to explaining the demise of *Prochlorococcus* at high latitudes, we believe that shared predation with bacteria may also partially explain the demise of *Prochlorococcus* with depth. Similar to the poleward decrease of *Prochlorococcus* growth rate with temperature, the relative growth rates of *Prochlorococcus* and other autotrophs decline significantly with depth as the amount of available light decreases. In this case, the demise could be set by a balance between organic material sinking from above and the decreasing light intensity.

We have focused on shared predation with bacteria. However, shared predation could also be at play with similar-sized phytoplankton or even between phytoplankton and generalist viruses. For instance, some predators can consume (11, 12) and some cyanophages can infect (43–45) both *Prochlorococcus* and *Synechococcus*. There are, indeed, some interesting shifts in *Prochlorococcus* distributions that may be related to indirect competition with *Synechococcus* (SI Appendix, Fig. S6 and Fig. 3D), although we note that the relationship between these two populations is less clear. Nonetheless, expanding the concept of shared predation to other functional types and size classes (e.g., *Synechococcus* and a similar-sized heterotrophic bacteria) could very likely help explain these nuances.

We contend that the observational data are consistent with the theoretical framework of shared predation combined with size class grazing that can explain the decreases of *Prochlorococcus* in higher latitudes. This theory does not require any assumption about how *Prochlorococcus* growth is impacted by temperature. However, we do note that the theory is, by design, extremely simple in order to illustrate key concepts. Indeed, the real world is far more complex, and growth rates are impacted by other factors, such as temperature, nutrients, and light. These complexities are captured in the more complex global biophysical model

(Fig. 4 C and D), but the theoretical ideas of the collapse still hold, showing that the underlying mechanism is robust. However, even the three-dimensional model is simplified with regard to the real ocean. Here we have, for instance, assumed heterotrophic bacteria and phytoplankton do not compete for resources, and the food web remains relatively simple. Nonetheless, the observations in the real world are consistent with the theoretical insight, suggesting that shared predation plays an important role in the dramatic collapse of *Prochlorococcus* populations (Fig. 1). Further observational exploration of these ideas, especially those focused on the seasonal progression of these ecological transitions (39), should help quantify the importance of shared predation and related mechanisms in structuring plankton biogeography.

Understanding what sets the biogeography of plankton and their dynamics has become additionally important under a changing climate. These organisms set the foundation for global fisheries and are the conduits for long-term carbon storage in the deep ocean. Current methods for predicting the range shifts of phytoplankton are often based on assuming that current correlations with temperature are predictive in a future ocean. Ecological mechanisms, such as shared predation, show how small changes in our understanding can lead to large changes in our predictions into the future. If we want to correctly project and understand how the ocean ecosystem will evolve moving forward, it is critical that we build a mechanistic understanding of how the community structure currently functions. The controls of *Prochlorococcus* distributions are important to understand, given this species' numerical dominance in many regions of the ocean. Currently suggested controls based on a thermal growth threshold (5) are insufficient to explain the species' collapse at high latitudes (Fig. 1). We utilize a bacteria-driven mechanism (3), which allows for decreasing *Prochlorococcus* abundance at higher production rates. We provide theoretical, numerical, and observational support for the idea that shared predation with heterotrophic bacteria controls the range of *Prochlorococcus*. In our theory, size-based predation leads to the emergence of a critical size separating heterotrophic bacteria from phytoplankton. This size increases with nutrient supply, eventually eliminating the smallest phytoplankton, *Prochlorococcus*, from the ecosystem. This same mechanism is likely to play a role in the dynamics of the next largest phytoplankton, *Synechococcus*, and the formulation may prove useful for simplifying our understanding of the relationships between other functional groups of plankton.

While the importance of direct interactions such as grazing have been established (18, 19, 40) and included in models, the role of indirect interactions such as shared grazing is often ignored. Yet, as we show here, these interactions may be crucial to understanding *Prochlorococcus* distributions. Thus, the knowledge of the microbial network of interactions, including processes such as shared grazing and other indirect interactions (e.g., refs. 3 and 46–50), is essential for being able to understand and predict community structure and its changes in the future. Equally important are frameworks, like the one presented here, which allow quantification of the broader effects of these interactions across the plankton size spectrum.

Bottom-up resource controls and top-down predator controls conspire to set the biogeography of plankton in the sea. The simplest, size-based theories suggest that, as nutrient supplies increase, larger phytoplankton grow in while the smaller phytoplankton remain. For the smallest, and arguably most critical, phytoplankton *Prochlorococcus*, this prediction is broken, as *Prochlorococcus* populations collapse at high latitudes. The location of this collapse in the North Pacific is difficult to explain with temperature but emerges naturally if shared predation with similarly sized heterotrophic bacteria is considered. Under the theory posed here, heterotrophic bacteria increase with nutrient supplies along with higher biomass of larger phytoplankton and higher primary production. If heterotrophic bacteria and *Prochlorococcus* share a grazer, this increase in bacterial

populations can lead to a collapse of *Prochlorococcus*. This study highlights that the marine ecosystem needs to be understood, and modeled, as an interconnected network of organisms where both direct and indirect interactions control community structure.

Materials and Methods

Observational Data. Cell count data used in Fig. 5 were provided by discrete samples collected in surface water (~5 m depth), fixed with glutaraldehyde (0.25% final concentration), and analyzed by Influx's cell sorter. *Prochlorococcus* was discriminated based on forward light scattering and red autofluorescence emitted by pigments. Total bacteria counts were obtained by staining the sample with SYBR Green I (0.01% final concentrations), and heterotrophic bacteria counts were calculated by subtracting *Prochlorococcus* cell counts from total bacteria counts (51). Biomass estimates in Fig. 6 use diameters and carbon quotas of individual cells estimated by the Influx (heterotrophic bacteria) and SeaFlow flow cytometers (*Prochlorococcus* and total phytoplankton) from forward light scatter measurements by the application of Mie theory for spherical particles, calibrated using particles of known refractive indices and several phytoplankton species of known carbon content (4). Data were averaged across tenth-of-a-degree bins. All data used in this study are publicly available. Data for the COOK-BOOK cruises [April and October 2003 (8)] can be found at <https://hahana.soest.hawaii.edu/cookbook/cookbook.html>; data for the SCOPE-Gradients cruises (April 2016, May/June 2017) are available via the open access data repository Zenodo: Influx 2016: 10.5281/zenodo.4085858; Influx 2017: 10.5281/zenodo.4085873; SeaFlow: 10.5281/zenodo.3994953.

Steady-State Solutions of Theoretical Frameworks. An informative set of nonzero steady-state results for the size-based model without bacteria (Eqs. 1–3) are

$$R_{pj}^* = \frac{\kappa_j(m_p + g_j Z_j^*)}{\mu_{pj}^{\max} - (m_p + g_j Z_j^*)} \quad [M1]$$

$$P_j^* = \frac{m_z}{\gamma g_j} \quad [M2]$$

$$Z_j^* = \frac{1}{g_j} \left(\frac{\mu_{pj}^{\max} R_p^*}{R_p^* + k_j} - m_p \right) \quad [M3]$$

$$\sum Z_j^* = \frac{\gamma S_{R_p}}{m_z} - \sum \frac{m_p}{g_j}. \quad [M4]$$

Solutions for the more complex system including heterotrophic bacteria and an organic detrital pool (Eqs. 6–10) are

$$R_{pj}^* = \frac{\kappa_{pj}(m_p + g_j Z_j^*)}{\mu_{pj}^{\max} - (m_p + g_j Z_j^*)} \quad [M5]$$

$$P_j^* + H_j^* = \frac{m_z}{\gamma g_j} \quad [M6]$$

$$Z_j^* = \begin{cases} \frac{1}{g_j} \left(\frac{\mu_{hj}^{\max} R_h^*}{R_h^* + \kappa_{hj}} - m_h \right) \\ \frac{1}{g_j} \left(\frac{\mu_{pj}^{\max} R_p^*}{R_p^* + \kappa_{pj}} - m_p \right) \end{cases} \quad [M7]$$

$$R_{hj}^* = \frac{\kappa_{hj}(m_h + g_j Z_j^*)}{\mu_{hj}^{\max} - (m_h + g_j Z_j^*)}. \quad [M8]$$

And, if we assume $m_h = m_p$, we additionally find

$$\sum Z_j^* = \frac{\gamma}{m_z} \left(S_{R_p} + \frac{S_{R_h}^*}{\delta} \right) - \sum \frac{m_p}{g_j}. \quad [M9]$$

Allometric Scaling of Plankton Growth and Grazing. In both zero-dimensional and the global three-dimensional model, we follow the allometric scaling used in ref. 35. Phytoplankton and heterotrophic maximum growth rate, nutrient affinity, grazing, and sinking (in the three-dimensional model), as well as zooplankton maximum grazing rates, are parameterized as power law functions of cell volume V : aV^b . Thus, many size classes can be described by just two coefficients (a , b) per parameter (SI Appendix, Table S1). Plankton smaller than 3 μm have an increase of growth rate with size, and those larger than 3 μm have a decrease of growth rate with size. This unimodal distribution has been observed (30, 52–55) and explained as a trade-off between replenishing cell quotas versus synthesizing new biomass (56, 57). Thus the b for maximum growth rates are different below and above 3 μm ESD. The intercept a depends on the functional group of

phytoplankton in the three-dimensional model (see discussion below, and SI Appendix, Table S1). The smallest phytoplankton have the highest affinity for nutrients (21), as a result of the lower surface to volume ratio found in larger cells (52, 58).

The choice of allometric scaling parameters for the heterotrophic bacteria is less clear, as fewer laboratory studies have examined the size dependence of growth as compared to phytoplankton and zooplankton. Additionally, different bacteria may consume a variety of types of organic matter (59, 60). However, it is generally recognized that larger bacteria (copiotrophs) grow significantly faster than smaller oligotrophs (59–61), and also that smaller oligotrophs have high nutrient affinities (27, 60). Without better constraints, we chose to use the same power exponent for growth rate and uptake as for phytoplankton (although this is not a requirement; see additional extra zero-dimensional experiment discussed below). The intercept a value was chosen to provide the smallest bacteria types with maximum growth rates on the order of 0.1 per day (62, 63).

Zero-Dimensional Model. The model simulation results presented in Fig. 3 B and D represent a parcel of water (i.e., zero-dimensional) with an influx of resource S_{R_p} . We integrate the two sets of equations (Eqs. 1–3 and 6–10) from initial conditions (set by the grazing-free steady-state analytical solutions) until a (quasi-) steady state is reached using a fourth-order Runge–Kutta scheme (64). We include 10 size classes (j) in the simulations, but show only the 4 smallest for simplicity. Size spacing was the same as in the global model (Fig. 4D). Plankton growth and grazing parameters are allometrically based, as discussed above, and are provided in SI Appendix, Table S1. We run individual simulations with different values of S_{R_p} spanning from very low values through values typical of the subtropical gyres (10^{-7} mmol $\text{N} \times \text{m}^{-3} \times \text{s}^{-1}$, as suggested by the global numerical model) to high values more typical of seasonal higher latitudes. Each grid in Fig. 3 B and D shows the final solution of each of these experiments. At low values of S_{R_p} , these results are the steady-state solution. At higher supply rates, the simulations develop predator–prey oscillations, and here we present the average of these cycles.

We perform a number of additional simulations to test parameter choices as well as sensitivity to resource supply assumptions. We show three additional results in SI Appendix. We repeat the experiment seen in Fig. 3D with the smallest size class only consisting of bacteria, and find, qualitatively, the same results (SI Appendix, Fig. S1B). Additionally, for ease in the theoretical proof showing that only one size class of phytoplankton and heterotrophic bacteria can coexist, we assumed that there is no external supply rate (i.e., S_{R_p} was entirely due to remineralization of detrital matter) and that the allometric relationships for bacteria and phytoplankton growth parameters were identical. We perform two additional numerical experiments that show that these assumptions are not critical to the qualitative behavior described above (SI Appendix).

The Global Biophysical Ocean Model. The biogeochemical/ecosystem model captures the cycling of C, N, P, Si, and Fe as they pass through inorganic and (dead and living) organic pools (35, 65). The specific details of the ecosystem follow from ref. 34 (see SI Appendix for the exact differences from that model) and resolves 31 phytoplankton (2 picoprokaryotes, 2 picoeukaryotes, 5 coccolithophores, 5 diazotrophs, 9 diatoms, 8 mixotrophic dinoflagellates), 16 zooplankton, and 3 heterotrophic bacteria. Phytoplankton have size resolution spanning from 0.6 μm to 140 μm ESD, zooplankton spanning 4.5 μm to 1,636 μm , and bacteria spanning 0.4 μm to 0.9 μm (Fig. 4C). The new component of the model relative to ref. 34 is the inclusion of the heterotrophic bacteria component. We follow ref. 26 in parameterizing the bacteria as consuming dissolved organic matter with a Michaelis–Menten function. A fraction of the uptake goes to bacteria growth, and the remainder is remineralized to inorganic nutrients.

Parameters influencing phytoplankton growth, grazing, and sinking are related to size (35, 66), with specific differences between the six functional groups (35) (SI Appendix, Table S1, and discussion above). Phytoplankton growth is limited by multiple nutrients (N, P, Fe, and Si in the case of diatoms), light (following ref. 67), and temperature (following refs. 68 and 69). Grazing is parameterized using a Holling II function (70) and is size specific such that grazers can prey upon plankton 5 to 15 times smaller than themselves, with an optimal size of 10 times smaller (36–38). The emergent size distribution of the simulated plankton populations is strongly controlled both by the rate of supply of limiting nutrients (bottom up) and by grazing (top down) (19, 35).

The biogeochemical and biological tracers are transported and mixed by the Massachusetts Institute of Technology (MIT) general circulation model (MITgcm) (71), constrained to be consistent with altimetric and hydrographic observations (72). This three-dimensional configuration has coarse

resolution (1° by 1° horizontally) and 23 levels ranging from 10 m in the surface to 500 m at depth.

Two simulations are presented: one with explicit bacteria and the other where organic matter remineralized to inorganic nutrients with a temperature-dependent rate, as used in previous versions of this model (e.g., ref. 35). The two simulations were run for 10 y. The ecosystem quickly (within 2 y) reaches a quasi-steady state. Here we show results from the 10th year of the simulations.

The latitudinal distribution and seasonality of bulk ecosystem properties such as chlorophyll (SI Appendix, Fig. S2), as well as distributions of size classes and functional groups, are plausible in comparison with satellite and in situ observations (73, 74) (SI Appendix, Figs. S2 and S3). The model results are similar to those in previous studies, with slight modification in the model (e.g., with a slightly different number of larger size classes) and different physical frameworks that have conducted additional evaluation (e.g., refs. 35, 75, and 76). As in those studies, we find that our model Chl a compares well to satellite estimates, although we note that the satellite estimates have large uncertainties (77) (more than 35% errors). The coarse resolution of the model does not capture important physical processes near coastlines, and lack of sedimentary and terrestrial supplies of nutrients and organic matter lead to Chl a being too low in these regions. Chl a is underestimated by the model in the subtropical gyres, likely due to lack of mesoscale processes in the model that would supply additional nutrients in these regions (78). The model results have the observed ubiquitous picophytoplankton and the limitation of the larger size classes to the more productive regions. The model picophytoplankton size class Chl a is potentially slightly too low, and the nano size class is potentially too high. However, we note that, if we set the pico/nano break at the model fifth size class (just under $3 \mu\text{m}$) instead of at the fourth ($2 \mu\text{m}$) size class, the relative values are much more in line with the satellite product. We suggest that the satellite product division might not be that accurate.

We also compare to a synthesis of phytoplankton abundances (world atlas of MARINE Ecosystem DATA [MAREDAT]) (74) (SI Appendix, Fig. S3). Although the observations are sparse, we do capture the ubiquitous nature of the picophytoplankton, the limited domain of the diazotrophs (including

observed lack of diazotrophs in the South Pacific gyre), and the pattern of enhanced diatom biomass in high latitudes and low diatom biomass in subtropical gyres. We overestimate the coccolithophore biomass relative to MAREDAT in many regions, but note that the conversion from cells to biomass in that compilation was estimated to have uncertainties of several 100% (79). The MAREDAT compilation did not include a category for dinoflagellates.

Model code and parameters (80) and model biomass output (81) are available online (see Data Availability).

Data Availability. The csv files from flow cytometer data have been deposited in Zenodo (10.5281/zenodo.4085858; 10.5281/zenodo.4085873; 10.5281/zenodo.3994953). Data for the COOK-BOOK cruises [April and October 2003 (8)] can be found at <https://hahana.soest.hawaii.edu/cookbook/cookbook.html>. All data used can be accessed directly through the Simons CMAP project, <https://simonscmmap.com/>, for download (82). The physical model used here is available through mitgcm.org, and the generic ecosystem code used in this study is available through <https://github.com/darwinproject/darwin3>. The specific modifications for the setup used here, and all parameter values, are available via Harvard Dataverse <https://doi.org/10.7910/DVN/ISJQ1W>, and the model output used in this manuscript is available at <https://doi.org/10.7910/DVN/FEWXB4>.

ACKNOWLEDGMENTS. We thank all the members of the Simons Collaboration on Ocean Processes and Ecology (SCOPE)-Gradients program for conversation about this region of the ocean, and the identification of the “Propocalypse.” We also thank John MacMillan, Kelsy Cain, and Annette Hynes for processing and curating flow cytometry data from SCOPE-Gradient cruises. This work was supported by multiple grants from the Simons Foundation. The Simons Collaboration on Computational Biogeochemical Modeling of Marine Ecosystems (CBIOMES) supported S.D. and M.J.F. on CBIOMES Grant 549931 and supported C.L.F. on CBIOMES Grants 827829 and 553242. F.R. was funded on Simons Life Sciences Project Award 574495. SCOPE funded D.C., E.V.A., and M.J.F. on SCOPE Award 329108. E.Z. was funded by the Simons Postdoctoral Fellowship in Marine Microbial Ecology. Additional support came from NASA (Grant 80NSSC17K0561, S.D., F.R., E.V.A., and M.J.F.).

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