

Light-driven synchrony of *Prochlorococcus* growth and mortality in the subtropical Pacific gyre

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Theoretical studies predict that competition for limited resources reduces biodiversity to the point of ecological instability, whereas strong predator/prey interactions enhance the number of coexisting species and limit fluctuations in abundances. In open ocean ecosystems, competition for low availability of essential nutrients results in relatively few abundant microbial species. The remarkable stability in overall cell abundance of the dominant photosynthetic cyanobacterium *Prochlorococcus* is assumed to reflect a simple food web structure strongly controlled by grazers and/or viruses. This hypothesized link between stability and ecological interactions, however, has been difficult to test with open ocean microbes because sampling methods commonly have poor temporal and spatial resolution. Here we use continuous techniques on two different winter-time cruises to show that *Prochlorococcus* cell production and mortality rates are tightly synchronized to the day/night cycle across the subtropical Pacific Ocean. In warmer waters, we observed harmonic oscillations in cell production and mortality rates, with a peak in mortality rate consistently occurring ~6 h after the peak in cell production. Essentially no cell mortality was observed during daylight. Our results are best explained as a synchronized two-component trophic interaction with the per-capita rates of *Prochlorococcus* consumption driven either directly by the day/night cycle or indirectly by *Prochlorococcus* cell production. Light-driven synchrony of food web dynamics in which most of the newly produced *Prochlorococcus* cells are consumed each night likely enforces ecosystem stability across vast expanses of the open ocean.

cyanobacteria | cell division | mortality | flow cytometry | SeaFlow

Potential interdependencies between species diversity and ecosystem stability have gained increased focus due to global changes in species distributions and abundances (1). Strong predator–prey interactions are predicted to enhance the number of coexisting species and limit fluctuations in abundances (2, 3), whereas competition for limited resources is predicted to reduce biodiversity, in some instances, to the point of ecological instability (3, 4). Mechanisms underlying ecosystem stability remain challenging to characterize on relevant temporal and spatial scales, in part because few empirical data are available to test these theories.

Our focus is on the microbial communities within surface waters of the vast oligotrophic gyre of the north Pacific Subtropical Ocean. Here, competition for low concentrations of essential nutrients is hypothesized to result in relatively few abundant microbial species, typified by their extremely small cell sizes and streamlined genomes (5). The cyanobacterium *Prochlorococcus* numerically dominates the photosynthetic community in these regions, with a relatively constant cell abundance close to half a billion cells per liter despite a population doubling time of approximately one day (6). Such constant cell numbers are predicted when both prey and predators are abundant and broadly dispersed and when the predator–prey interactions are tightly coupled (7). Understanding the stability of this coupling is important as loss processes determine how *Prochlorococcus* cell

production is transferred into the marine food web. Estimates of cell production and cell mortality rates for *Prochlorococcus* remain sparse, however, as they are commonly derived from incubations in bottles (8) or measures of cell cycle progression over the day/night cycle (9), both of which are labor intensive and thus limited in their broad-scale applicability (10). Here we estimated *Prochlorococcus* cell production and cell mortality rates continuously across a 2,900-km transect in the northeast Pacific Ocean in winter (October–November 2011) using a shipboard-based flow cytometer, SeaFlow (11) and show that, throughout the gyre, rates of *Prochlorococcus* cell production and cell mortality are both tightly synchronized to the day/night cycle, which likely helps stabilize open ocean ecosystems.

Results and Discussion

Temperature Regulates the Rates of *Prochlorococcus* Cell Division.

Prochlorococcus division rates were estimated using a size-structured population model (12, 13) with the continuously measured size distribution of *Prochlorococcus* as input (*SI Appendix*, Figs. S1–S4). Continuous, daily averaged division rates of natural *Prochlorococcus* populations along the transect increased from 0.2 to 0.9 d⁻¹ (Fig. 1A) and were linearly correlated with temperature ($R^2 = 0.97$, $P < 0.001$; Fig. 1B). These computed division rates were similar to our independent estimates based on cell cycle analyses (*SI Appendix*, Figs. S2 and S3). Lower division rates were encountered in cooler waters (16–19 °C) at the edge of the north Pacific gyre and further inshore where the

Significance

The cyanobacterium *Prochlorococcus* is the most abundant photosynthetic organism in the oceans, driving marine food webs and biogeochemistry. Estimates of *Prochlorococcus* cell mortality and cell production are critical to determine how organic matter is transferred into the food web. Using novel high-resolution sampling methods on two winter-time cruises, we show that the daily production of *Prochlorococcus* cells in surface waters of the subtropical Pacific gyre is consistently balanced by their nightly consumption by other organisms. These synchronized loss processes suggest that *Prochlorococcus*-derived organic matter stabilizes multiple species interactions, from viruses to grazers. The observed resilience of this synchronized food web dynamic as temperature increased within the gyre suggests that ecosystem stability may persist in future warmer oceans.

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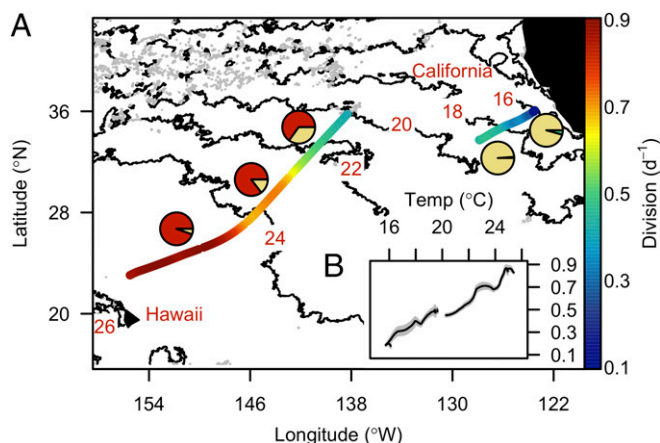


Fig. 1. Temperature determines the rate of *Prochlorococcus* cell division in surface waters of the Northeast Pacific Ocean. (A) Daily averaged division rate of *Prochlorococcus* (division, d^{-1} , color scale), monthly averaged satellite-based sea surface temperature for October ($^{\circ}C$, contour line), and *Prochlorococcus* ecotype composition (pie charts based on percent of each *Prochlorococcus* ecotype relative to total *Prochlorococcus* ecotypes represented in 16S rRNA gene amplicons from TAG sequencing; yellow, red, and green for eHLI, eHLII, and eLLI, respectively) during the northward cruise to California (4–9 October 2011) and southward cruise to Hawaii (29 October–2 November 2011). Daily division rates were calculated as the sum of hourly division rates over a 24-h period using a 1-h rolling window. (B) Relationship between daily division rates and surface temperature during the survey. Vertical gray bars represent SDs ($n = 24$).

high-light-adapted *Prochlorococcus* ecotype I (eHLI) dominated (>95% of all *Prochlorococcus* 16S rDNA barcoded sequences), and were similar to those previously observed in that region (<0.4 d^{-1}) (14) (Fig. 1A and *SI Appendix*, Fig. S5). Higher division rates were observed in the warmer subtropical gyre waters, which were dominated by the high-light-adapted *Prochlorococcus* ecotype II (eHLII) (65–94% of all *Prochlorococcus* 16S rDNA barcoded sequences) and were similar to, or higher than, the few individual measurements available from the Sargasso Sea (>0.7 d^{-1}) (14), another warm water ecosystem.

The highest in situ division rate (0.9 d^{-1}) occurred at 25 $^{\circ}C$ in low-latitude waters (25 $^{\circ}N$; Fig. 1B) and is remarkably similar to the highest division rate reported for a cultured representative of eHLII (MIT9312) grown under nutrient-replete conditions (*SI Appendix*, Fig. S6). Thus, the eHLII-dominated field population in the subtropical gyre was likely dividing at near-maximal rates, despite low ambient nutrient availability commonly found in this region (15). Similar to other ecotypes (16, 17), *Prochlorococcus* eHLII must possess efficient nutrient acquisition mechanisms and reduced physiological nutrient demands that allow it to maintain such high division rates in the face of nutrient scarcity. Under laboratory conditions, the growth rate of *Prochlorococcus* is positively correlated with temperature across the range of conditions tested in this study (18). In the field, *Prochlorococcus* is most abundant at warmer low latitudes (6). Both observations suggest a central role of temperature in regulating the rate of *Prochlorococcus* cell division in surface waters.

Synchronized Oscillations in the Rates of *Prochlorococcus* Cell Production and Cell Mortality. Despite the significant differences in daily division rates along the transect, maximum cell abundance ($\sim 400 \times 10^6$ cells/L) was comparable in both cooler and warmer waters (Fig. 2A–D). In warm waters (>19 $^{\circ}C$), cell numbers were tightly synchronized to the day/night cycle, with a net increase in abundance during the day and a net decrease at night (Fig. 2C, day 1, and D). No such synchrony was observed in the colder waters (<19 $^{\circ}C$; Fig. 2C, day 3–6). The day/night pattern observed in warm

waters differs from those observed in other warm water regions such as the equatorial Pacific, where cell abundances increase during the first half of the night and decrease or remain stable during the day (19). These different patterns of cell abundance over the day/night cycle indicated an underlying difference in the timing of cell division and/or cell loss processes among these regions.

Our use of continuous measurements of cell size allowed us to examine hourly cell division over the daily cycle. The peak in cell division consistently occurred near dusk, regardless of ecotype composition (Fig. 1A), and was reflected in the declining mean cell size of the *Prochlorococcus* population (*SI Appendix*, Fig. S7). Although most previous studies report essentially stable or decreasing cell abundance in *Prochlorococcus* with essentially no cell division during the day (6, 19), our data revealed that a significant proportion of the population underwent cell division (*SI Appendix*, Fig. S7C and D), similar to populations in the Indian Ocean (20). Our estimates of cell production (Fig. 2E and F), computed by multiplying our independent measures of cell abundance by our estimates of division rates, accounted for most of the observed changes in *Prochlorococcus* cell abundances during the day (Fig. 2D) and indicated that loss processes were negligible during this period.

The net loss rate of *Prochlorococcus* cells from a given location is a combination of biological factors (i.e., consumption by grazers and/or lysis by viruses) and physical factors (i.e., the removal or addition of cells through physical transport). In our study, the greatest source of physical transport was due to sampling from a moving ship, with other sources of physical transport considered as negligible for our calculations. Physically driven rates of changes in *Prochlorococcus* abundance was estimated based on the spatial gradient of *Prochlorococcus* abundance along the cruise track (*SI Appendix*, *SI Materials and Methods*). Biologically driven rates of cell loss (i.e., cell mortality) were based on the difference between the cell production and the rates of change of cell abundance over time, after correcting for physical transport (*SI Appendix*, Fig. S8). Hourly mortality rates were near zero during the day and peaked at night in subtropical gyre waters (<34 $^{\circ}N$; Fig. 2E, day 1, and F), with a more stochastic pattern observed further inshore (>34 $^{\circ}N$; Fig. 2E, day 2–6). In the subtropical gyre, the peak in mortality consistently occurred about 6 h after dusk (Fig. 2F) such that the bulk of the daily *Prochlorococcus* cell production (ranging from 88×10^6 to 289×10^6 cells/L per day; *SI Appendix*, Fig. S9) was consumed over the course of the night. This 90 $^{\circ}$ -phase difference (i.e., 6 h in a 24-h cycle) between the rates of cell production and cell mortality was consistent across 2,240 km of the gyre, regardless of water temperature or *Prochlorococcus* ecotype composition (Fig. 1A). The consistency of the phase and period of the oscillations in the rates of cell mortality was particularly surprising across such a large spatial scale given the diversity of potential sources of *Prochlorococcus* mortality (e.g., viruses and grazers) (21, 22). A similar diel pattern of cell production and cell mortality was observed in the same region the previous year (November 2010; *SI Appendix*, Fig. S10), suggesting that these synchronized oscillations may be relatively stable, at least over an annual cycle.

Light-Driven *Prochlorococcus* Cell Mortality. In dynamical systems theory, a synchrony between two coupled oscillators indicates that the periodicity of one signal entrains the phase and period of the other. The periodic biological signal in the subtropical gyre is the light-driven photosynthesis and division rate of *Prochlorococcus* that produces a daily pulse of new carbon. This pulse entrains a synchronous, phase-coupled transfer of the carbon into the food web. Synchronized predator–prey cycles such as those observed here are predicted to promote food web stability (7). No such synchrony was observed in the colder waters where the abundance of *Prochlorococcus* was significantly lower (< 150×10^6 cells/L), and the phytoplankton assemblage was dominated by larger sized

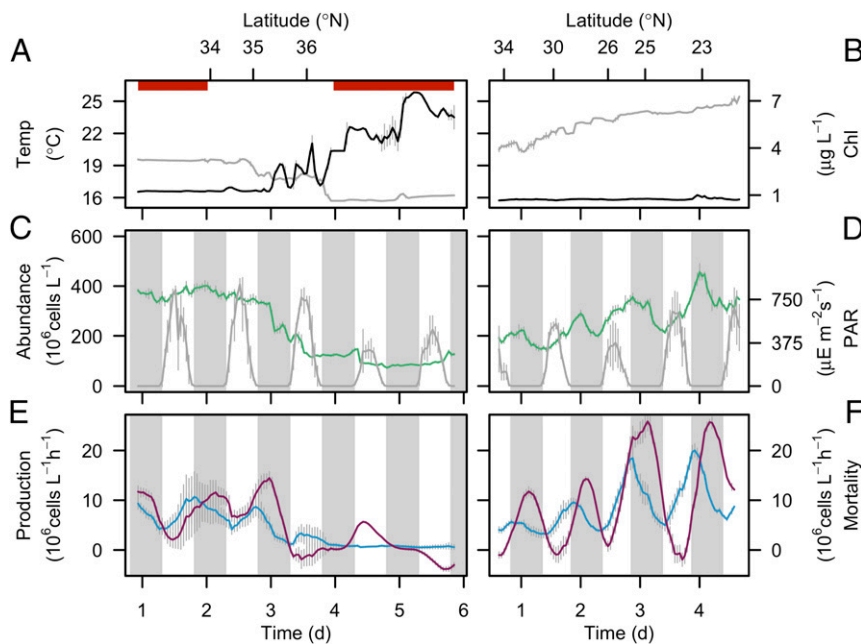


Fig. 2. Synchrony of *Prochlorococcus* mortality rates at high cell production rates. (A and B) Hourly averaged temperature ($^{\circ}\text{C}$, gray line) and bulk chlorophyll a concentrations ($\mu\text{g/L}$, black line), (C and D) hourly averaged cell abundances (10^6 cells/L, green line) and hourly averaged PAR at 5 m depth ($\mu\text{E}/\text{m}^2$ per second, gray line), and (E and F) hourly rates of cell production (10^6 cells/L per hour, blue line) and cell mortality (10^6 cells/L per hour, purple line) during the northward cruise to California (4–9 October 2011) (Left) and southward cruise to Hawaii (29 October–2 November 2011) (Right). Vertical gray bars represent SDs ($n = 20$ for temperature, chlorophyll a concentrations, PAR, and cell abundances, $n = 24$ for cell production and mortality rates). The gray regions indicate night. Horizontal red bars in A represent periods on station. Note that the left panels are shown according to chronology of sampling, as opposed to ascending sea surface temperature in Fig. 1B.

phytoplankton (SI Appendix, Fig. S11). As predicted by systems theory, changes in *Prochlorococcus* cell abundance over time were more stochastic in this region (Fig. 2E, day 3–6).

The synchronized oscillations in the rates of cell production and cell mortality observed in the subtropical gyre mirrors two-component models of virus/host or predator/prey systems, although compressed over the course of a daily cycle. We explored different models to determine which best emulates the observed oscillations in *Prochlorococcus* mortality rates characterized by limited cell loss during daylight. Classical models commonly assume that mortality rate depends exclusively on prey abundance, with a constant per-capita rate of host infection or prey consumption (23, 24). These models produce a diel oscillation in mortality rates but do not explain the negligible cell loss during the day (Fig. 3A and B). There is growing evidence that viral infection rates in the cyanobacteria *Synechococcus* and other marine phytoplankton depend on light either directly or indirectly via the host's cell cycle (25, 26). Mixotrophic protists, such as caryophytes, can graze at night and photosynthesize during the day (27), some of which consume *Prochlorococcus* in oligotrophic regions (22). Modeled cell mortality emulates our observations if we assume that the per-capita rate of viral infection varies over the day/night cycle or that grazing only occurs at night (Fig. 3C and D). The relative contribution of grazers and viruses to overall *Prochlorococcus* mortality in natural communities has not been quantified. However, *Prochlorococcus* abundances in the Sargasso Sea were observed to decrease dramatically during the latter half of the night, similar to the timing observed here; this nearly identical period of cell loss was experimentally linked to grazing activity rather than viruses (14). Regardless of whether the mortality we observed in the subtropical gyre is caused by viruses or by grazers, our results indicate that the day/night cycle, either directly or indirectly, restricts mortality to the night, allowing *Prochlorococcus* abundance to recover each day and maintain a stable population over vast regions of the ocean.

Concluding Remarks. The observed synchrony between cell production and cell mortality means that each day a pulse of *Prochlorococcus*-derived organic matter enters the ecosystem, either as dissolved organic matter derived from viral lysis, or as particulate organic matter grazed by higher trophic level organisms, or a combination of both. We estimate that *Prochlorococcus* pulses of carbon could account for more than 75% of the daily photosynthetic organic carbon production in the surface layer of the subtropical gyre during this time of year (SI Appendix, Fig. S9).

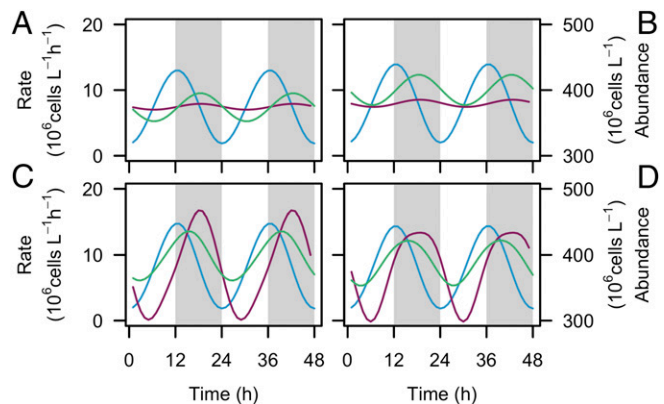


Fig. 3. Light dependency of *Prochlorococcus* mortality forces. Modeled *Prochlorococcus* mortality rate (purple line), production rate (blue line), and cell abundance (green line). (A and C) Stable solution for a virus/host model (23) where the per-capita rates of virus infection (A) remain constant over the day/night cycle or (C) vary according to *Prochlorococcus* cell division rates. (B and D) Stable solution for a predator–prey model (24) where the per-capita rates of predation (B) remain constant over the day/night cycle or (D) vary according to light intensity. The gray regions indicate night.

A recent study by Ottesen et al. (28) documented synchronous gene expression patterns across bacterial communities within the subtropical gyre and hypothesized that these patterns could result from light-driven pulses of dissolved organic matter released by *Prochlorococcus*. Our results indicate that a mortality-driven pulsed delivery of *Prochlorococcus*-derived organic compounds through the food web may serve as a stabilizing factor of additional trophic levels across vast expanses of the subtropical ecosystem. The next step is to determine whether similar patterns are observed in the subtropical gyre throughout the year and whether they are conserved across different oceanic regions.

Materials and Methods

Phytoplankton Abundance and Composition. Flow cytometry samples were collected underway from the continuous seawater flow-through system (5 m depth) during the northward cruise to California (4–9 October 2011) and during the southward cruise to Hawaii (16–18 November 2010 and 29 October–2 November 2011). Continuous underway measurements of *Prochlorococcus* abundances and cell size were made using SeaFlow (11), and data were analyzed using the R package *flowPhyto* (29) version 2.4.1, which uses statistical clustering methods to discriminate phytoplankton populations. Forward light scatter measured by the instrument was converted to cell volume using an empirical relationship between light scatter measured by SeaFlow and cell volume measured by a Coulter counter for different exponentially growing phytoplankton cultures (SI Appendix, SI Materials and Methods).

Estimated Cell Division Rates of *Prochlorococcus*. Hourly division rates of *Prochlorococcus* were estimated using the R package *ssPopModel* version 0.1.1, a modified version of the size-structured matrix population model developed by Sosik et al. (12). The model is based on the assumptions that (i) cell growth is determined by light exposure, (ii) the probability of a cell dividing depends on size, (iii) all cells within a discrete size class have the same probability to change to another size class, and (iv) a cell divides into two daughter cells, each half the size of the mother cell. The model predicts the cell size distribution over the course of the day using the cell size/cell division relationships and the light dependence of cell division (SI Appendix, SI Materials and Methods). We assumed that there was no differential mortality of *Prochlorococcus* based on cell size. Size distribution-based division rate estimates of *Synechococcus* have been shown to be similar between undiluted incubations (higher grazing pressure) and diluted incubations (lower grazing pressure) (30), supporting the assumption that size-selective grazing is not important. To establish the accuracy of our estimates of size distribution-based division rate estimates, we compared size-based estimates of *Prochlorococcus* division rates (h^{-1}) with rates of changes in cell number for laboratory cultures and with cell cycle-based estimates of division rates for natural *Prochlorococcus* populations sampled near the beginning and end of the transect (SI Appendix, SI Materials and Methods). Estimates of *Prochlorococcus* cell production was obtained by multiplying hourly averaged cell abundances by the hourly estimates of cell division rates.

Estimated Cell Mortality Rates of *Prochlorococcus*. The rate of change in the abundance of *Prochlorococcus*, C , measured by the instrument from a moving ship reflects changes in C due to local time-dependent biological processes (i.e., division and mortality) and physical processes (i.e., advection and diffusion) occurring at a particular location and an advective component due to the ship moving through a spatial gradient of C . Given that the greatest source of physical transport was due to sampling from a moving ship, with other sources of physical transport considered as negligible for our calculations, mortality rate is given by

$$gC = \mu C - \frac{dC}{dt} + v_{ship} \frac{\partial C}{\partial x},$$

where g is the mortality rate due to biological processes such as grazing and viral infection, μ is the division rate, C is the cell abundance, v_{ship} is the velocity of the ship and $\partial C/\partial x$ is the spatial gradient of C along the cruise track, x . To approximate the spatial gradient of C , we assumed that *Prochlorococcus* abundances remain constant from day to day at a given

location, as observed from previous studies (31, 32) and during the first day on station (Fig. 2C, from day 1 to day 2). Daily averaged C was calculated using a 24-h running mean of hourly averaged C . The daily distance was calculated as the distance between the location at t_0 and the location at $t + 24$ h. A 1-h rolling window was then used to determine the start of each 24-h period in the time series.

Idealized Ecological Models. We used two different mathematical models that describe two-trophic-level microbial systems (i.e., virus/host and predator/prey systems) in continuous culture, based on our estimates of *Prochlorococcus* division rates (SI Appendix, SI Materials and Methods). We modeled the dynamics of host/virus interactions according to Levin's model (23) where the per capita adsorption rate of the viruses was set either as a constant or defined to maximize cell lysis 8 h after the peak of *Prochlorococcus* cells in S phase (i.e., 2 h before dusk). The dynamics of the interactions between predator and prey was modeled using Lotka–Volterra equations (24), where the per-capita predation rate was set either as a constant or defined such that the predation rate was inversely correlated to irradiance. Abundances were simulated for 100 d to reach a system at equilibrium.

Estimated Daily Carbon Production of *Prochlorococcus*. Cell volume was converted to carbon per cell using a conservative estimate of 325 fg C/ μm^3 (33). This yields median values of carbon content per cell of 25 and 50 fg C/cell in the subtropical gyre and near the California coast, respectively. These estimates are in agreement with direct measurements made on cultures (49 fg/cell) (34). The carbon per cell was multiplied by the estimated daily cell production at the end of the southward transect (near Hawaii) to obtain the mean daily carbon production (0.59 mmol C/ m^3 per day). Our estimate of daily carbon production represents more than 75% of the seasonal gross primary production at station ALOHA (0.75 mmol C/ m^3 per day) (35), which is based on ^{18}O -labeled O_2 in bottles over a 24-h incubation period.

***Prochlorococcus* Ecotype Composition.** DNA samples were collected underway from the continuous seawater flow-through system. Genomic DNA was extracted from 500-mL water samples filtered through a 0.2- μm Supor filter following the protocol described in Demir-Hilton et al. (36) for the northward cruise and through a 0.2- μm GTPP Isopore membrane filter using the QIAGEN DNeasy tissue kit according to the manufacturer's protocol for gram-negative bacterial cells for the southward cruise. Filters were first thawed and vortexed vigorously with lysis buffer using a bead beater (Bio-Spec Products) for 30 s at 2,100 $\times g$ (Eppendorf 5418 with rotor FA-45-18-11), and the buffer with cells was then collected and processed according to the manufacturer's protocol. Barcoded V1-V2 region 16S rRNA data were produced and analyzed using a previously published method (37) that incorporates *pplacer* (38) version 1.1.15. The number of *Prochlorococcus* amplicons produced were 1,382, 34,351, 40,046, 30,738, and 38,153 (ordered by ascending sea surface temperature).

Auxiliary Data. Monthly averaged fields of satellite-based sea surface temperature were obtained from the NASA website (disc.sci.gsfc.nasa.gov/giovanni). Photosynthetic active radiation (PAR) was measured using an underwater Biospherical Instruments light sensor with log amplifier. Underway sea surface temperature was measured using a shipboard Sea-Bird Electronics SBE-21 thermosalinograph, and underway chlorophyll *a* fluorescence was measured using a shipboard WET Labs ECO FLRTD chlorophyll fluorometer.

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