

# Diel population dynamics and mortality of *Prochlorococcus* in the North Pacific Subtropical Gyre

Stephen J. Beckett<sup>1,2,†,\*</sup>, David Demory<sup>1,2,†,\*</sup>, Ashley R. Coenen<sup>2,3</sup>, John R. Casey<sup>4,5,6</sup>, Mathilde Dugenne<sup>4,5</sup>, Christopher L. Follett<sup>6,7</sup>, Paige Connell<sup>8,9</sup>, Michael C.G. Carlson<sup>10</sup>, Sarah K. Hu<sup>8,11</sup>, Samuel T. Wilson<sup>4,5</sup>, Daniel Muratore<sup>1,2,12</sup>, Rogelio A. Rodriguez-Gonzalez<sup>1,2,12</sup>, Shengyun Peng<sup>1,2,13</sup>, Kevin W. Becker<sup>11,14</sup>, Daniel R. Mende<sup>4,5,15</sup>, E. Virginia Armbrust<sup>16</sup>, David A. Caron<sup>8</sup>, Debbie Lindell<sup>10</sup>, Michael J. Follows<sup>6</sup>, Angelicque E. White<sup>4,5</sup>, François Ribalet<sup>16</sup>, and Joshua S. Weitz<sup>1,2,3,\*</sup>

<sup>1</sup>School of Biological Sciences, Georgia Institute of Technology, Atlanta, GA, USA

<sup>2</sup>Center for Microbial Dynamics and Infection, Georgia Institute of Technology, Atlanta, GA, USA

<sup>3</sup>School of Physics, Georgia Institute of Technology, Atlanta, GA, USA

<sup>4</sup>Daniel K. Inouye Center for Microbial Oceanography: Research and Education, University of Hawai'i at Mānoa, Honolulu, Hawai'i, HI, USA

<sup>5</sup>Department of Oceanography, University of Hawai'i at Mānoa, Honolulu, Hawai'i, HI, USA

<sup>6</sup>Department of Earth, Atmospheric, and Planetary Sciences, Massachusetts Institute of Technology, Boston, MA, USA

<sup>7</sup>Cardiff University, Wales, UK

<sup>8</sup>Department of Biological Sciences, University of Southern California, Los Angeles, CA, USA

<sup>9</sup>Biology Department, San Diego Mesa College, San Diego, CA, USA

<sup>10</sup>Faculty of Biology, Technion – Israel Institute of Technology, Haifa, Israel

<sup>11</sup>Department of Marine Chemistry and Geochemistry, Woods Hole Oceanographic Institution, Woods Hole, MA, USA

<sup>12</sup>Interdisciplinary Graduate Program in Quantitative Biosciences, Georgia Institute of Technology, Atlanta, GA, USA

<sup>13</sup>Adobe, San Jose, CA, USA

<sup>14</sup>GEOMAR Helmholtz Centre for Ocean Research, Kiel, Germany

<sup>15</sup>Laboratory of Applied Evolutionary Biology, Department of Medical Microbiology, Academic Medical Centre, University of Amsterdam, Amsterdam, The Netherlands

<sup>16</sup>School of Oceanography, University of Washington, Seattle, Washington, WA, USA

<sup>†</sup>These authors contributed equally.

\*Corresponding authors: [sjbeckett@gatech.edu](mailto:sjbeckett@gatech.edu), [david.demory@biosci.gatech.edu](mailto:david.demory@biosci.gatech.edu), [jsweitz@gatech.edu](mailto:jsweitz@gatech.edu)

## Abstract

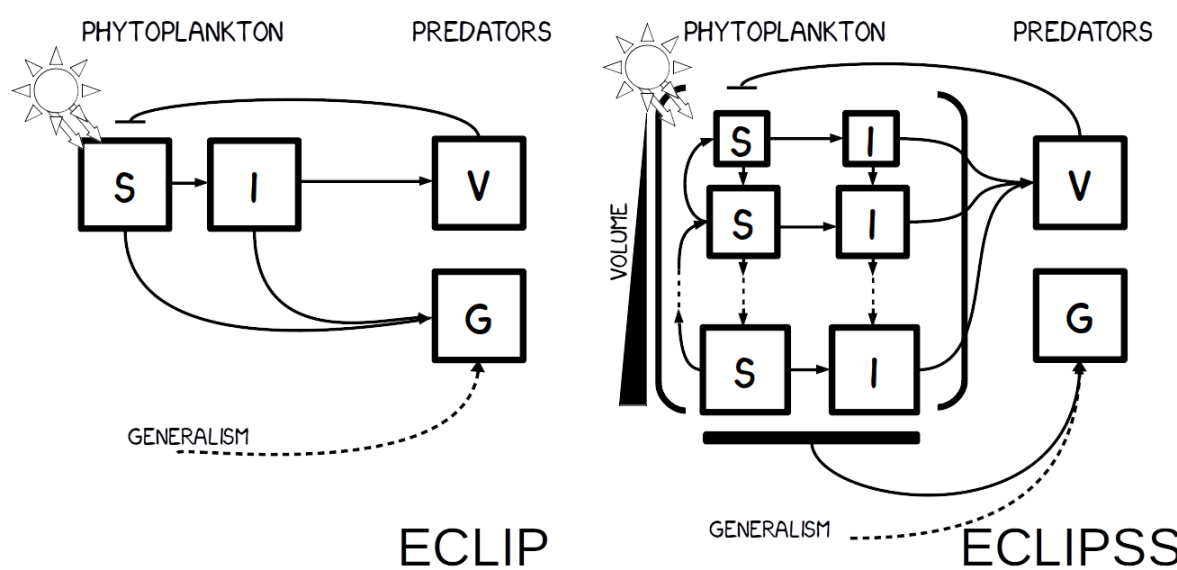
Marine ecosystem models often consider temporal dynamics on the order of months to years, and spatial dynamics over regional and global scales as a means to understand the ecology, evolution, and biogeochemical impacts of marine life. Large-scale dynamics are themselves driven over diel scales as a result of light-driven forcing, feedback, and interactions. Motivated by high-frequency measurements taken by Lagrangian sampling in the North Pacific Subtropical Gyre, we develop a hierarchical set of multitrophic community ecology models to investigate and understand daily ecological dynamics in the near-surface ocean including impacts of light-driven growth, infection, grazing, and phytoplankton size structure. Using these models, we investigate the relative impacts of viral-induced and grazing mortality for *Prochlorococcus*; and more broadly compare *in silico* dynamics with *in situ* observations. Via model-data fitting, we show that a multi-trophic model with size structure can jointly explain diel changes in cyanobacterial abundances, cyanobacterial size structure, viral abundance, viral infection rates, and grazer abundances. In doing so, we find that a significant component (between 5% to 55%) of estimated *Prochlorococcus* mortality is not attributed to either viral lysis (by T4- or T7-like cyanophage) or grazing by heterotrophic nanoflagellates. Instead, model-data integration suggests a heightened ecological relevance of other mortality mechanisms – including grazing by other predators, particle aggregation, and stress-induced loss mechanisms. Altogether, linking mechanistic multitrophic models with high-resolution measurements provides a route for understanding of diel origins of large-scale marine microbial community and ecosystem dynamics.

# 1 Introduction

Highly resolved surface ocean observations suggest that diel light rhythms drive repeatable changes in the abundance of ubiquitous cyanobacteria at the base of the microbial food web, including both *Prochlorococcus* and *Synechococcus* [1, 2]. Cyanobacterial population dynamics are influenced by both nutrients and light availability [2, 3, 4, 5, 6, 7, 8] as well as by density- and size-dependent feedback processes with other components of the community [9, 10, 11, 12, 13, 14, 15]. These interactions lead to diel oscillations, including in grazing rates, viral abundances, viral infection rates, and viral activity [16, 17, 18, 19, 20, 21]. The emergence of diel rhythms in abundances and process rates raises the question: how does light-driven growth of photoautotrophs over day-night cycles influence oscillations of other microbial populations and ecosystem processes? Resolving this question is challenging because it is often hard to disentangle process from pattern, e.g., reduced population growth and/or increased mortality can have the same net effect on cyanobacterial abundances. This type of identifiability problem is inherent to prior efforts to model size-structured phytoplankton growth with generic loss terms [2, 9, 10]. The identifiability problem is compounded by the fact that daily rhythms of microbes are also entrained, in part, with oscillatory day-night light forcing. In this study, we set out to use an ecological modeling framework, combined with field observations that go beyond cyanobacterial population measurements, as a means to understand how observed rhythms in *Prochlorococcus* dynamics influence and are influenced by rhythms of other populations in the microbial community in the North Pacific Subtropical Gyre (NPSG).

The NPSG is an oligotrophic system numerically dominated by the unicellular cyanobacteria *Prochlorococcus*, the most abundant phototroph in the global oceans [22]. Grazers and viruses are hypothesised to be the dominant drivers of *Prochlorococcus* loss [23, 24]. However estimating the relative contribution of viral-induced mortality and grazing-induced loss remains challenging [20, 25, 26, 27, 28]. We focus our study on a near-surface Lagrangian parcel of water in the NPSG, which was sampled at 15m depth over the course of ten days in summer 2015. Sampling on this expedition was performed at high time resolution (i.e., at four hour intervals for some components and nearly continuously for others). Related work from this cruise using the cellular iPolony method estimates that viruses, despite being highly abundant, may contribute to approximately 5% or less of total *Prochlorococcus* cell losses per day [20]. Congruently, using quota based reasoning, grazing by heterotrophic nanoflagellates could potentially account for the majority of *Prochlorococcus* cell losses per day [21], but this method cannot rule out the potential for significantly lower rates of grazing. We note that cruise data show clear repeatable periodicity in average *Prochlorococcus* cell volume, which peaks close to dusk, and cell abundance, which peaks during nighttime. Population abundances of heterotrophic nanoflagellates oscillate as does the fraction of cells infected by of T4- and T7-like viruses [20, 21]; suggesting the potential for emergent synchronization.

Here, we synthesise and integrate data from the 2015 cruise; augmented by the development, fitting and comparison with multi-trophic models. Our primary aim is to understand how mechanistic interactions between cyanobacteria, viruses, and grazers explain the observed diel variation in population abundances, process rates, and aggregate community dynamics in a model ocean ecosystem. To do so, we developed a hierarchical set of nonlinear dynamic models of microbial interactions spanning the cyanobacterium *Prochlorococcus*, heterotrophic nanoflagellate grazers and T4- and T7-like cyanophages in an oscillating light environment, and including phytoplankton size structure. These models combine principles of microbial interactions (including virus-microbe interactions) [28, 29, 30, 31, 32, 33] with size-structured models of phytoplankton growth [2, 7, 9, 10, 12, 13, 34, 35, 36]. We use these models to examine a series of related questions: how is the system expected to behave over diel timescales and how does light driven forcing of cyanobacterial growth affect size-dependent interactions between marine prey and predators? To do so, we compare *in silico* ecological rhythms with *in situ* ecological rhythms as a means to identify whether mechanistic representations of feedback can recapitulate observed rhythms and, if so, to what extent we can disentangle the relative contribution of viral-induced lysis and grazing to total *Prochlorococcus* mortality. Indeed, as we show, model-data integration suggests that the tight coupling of *Prochlorococcus* growth and loss over diel cycles may not be controlled only by cyanophages and heterotrophic grazers, but that additional factors are at play, raising new questions on governing mechanisms at the base of the marine food web.



**Figure 1: Community ecological models of viral and grazer mediated predation.** Phytoplankton are structured by viral infection status; either susceptible ( $S$ ) or infected ( $I$ ). Susceptible phytoplankton can become infected upon encountering viruses ( $V$ ). Infected cells produce virions which are released into the environment upon viral-induced lysis. Both susceptible and infected cells are prone to grazing pressure from grazers ( $G$ ). Additionally, grazers may have a generalist strategy (e.g., grazing on heterotrophs, mixotrophs and phytoplankton not represented by  $S$  and  $I$ ) denoted by the dotted line. a) the ECLIP model b) the ECLIPSS model in which phytoplankton are additionally structured by size. Three size classes are graphically presented; more could be included as denoted by the dotted lines.

## 2 Modeling Framework

### 2.1 Overview of Model Structure

We develop two related models of an Ecological Community driven by Light including Infection of Phytoplankton, with and without Size Structure, which we term ECLIP and ECLIPSS, respectively. The first model includes dynamics of *Prochlorococcus*, grazers, and viruses (ECLIP). The second model extends this model to include *Prochlorococcus* size structure (ECLIPSS). These models include phytoplankton division and loss, where the loss arises due to a combination of grazing, viral lysis, and other factors. The ECLIPSS model also includes a phytoplankton growth term and explicit accounting for variation in cell size. In both models, viruses denote abundances of a focal set of T4- and T7-like cyanophages which are known to primarily infect *Prochlorococcus*. The grazers represent heterotrophic nanoflagellates which can feed on multiple prey types [37], however the primary food source for heterotrophic nanoflagellates may be *Prochlorococcus* [21]. As a result, we introduce flexibility in our model framework to account for whether or not *Prochlorococcus* constitutes the primary source of food for the observed grazer class. To assess this uncertainty we investigated models of two grazer strategies, with either a “specialised” grazer class that only consumes *Prochlorococcus* or a “generalised” grazer class that includes additional prey sources, e.g., heterotrophic bacteria which are not explicitly integrated into the models. Mixotrophic nanoflagellates [38] were observed at the study site, but are thought to contribute less to the grazing pressure on the bacterial community compared to heterotrophic nanoflagellates [21]. As it was not possible to differentiate measurements of mixotrophic nanoflagellates from measurements of phototrophic nanoflagellates [21] we choose to focus our attention on grazing by heterotrophic nanoflagellates. In both models, we search for sets of ecological parameters that are biologically feasible and use model-data integration approaches to identify parameter sets that generate dynamics consistent with the observed population dynamics in the NPSG. Details of ECLIP, ECLIPSS, and size structured assumptions follow. A complete description of the model structure and justifications for functional and parameter choices are found in the Supplementary Methods.

## 2.2 Ecological model of phytoplankton communities with viral and grazer mediated predation (ECLIP)

The ECLIP model represents *Prochlorococcus* cell division as a light-driven process (where cell division is expected to occur at night [2, 39]) and *Prochlorococcus* cell losses are controlled by viral lysis, grazing, and other density-dependent factors (Figure 1a). The *Prochlorococcus* population is structured by two states of infection: cells that are susceptible to viral infection ( $S$ ) and cells that are infected ( $I$ ) by viruses ( $V$ ). Grazers ( $G$ ) feed indiscriminately on both on  $S$  and  $I$  classes. The dynamics of the abundances of  $S$ ,  $I$ ,  $V$  and  $G$  over time are described by the following system of equations:

$$\begin{aligned} \frac{dS}{dt} &= \overbrace{\mu(t)S}^{\text{division}} - \overbrace{m_{p1}S - m_{p2}S(S+I)}^{\text{losses}} - \overbrace{\phi SV}^{\text{infection}} - \overbrace{\psi SG}^{\text{grazing}} \\ \frac{dI}{dt} &= \overbrace{\phi SV}^{\text{infected}} - \overbrace{m_{p1}I - m_{p2}I(S+I)}^{\text{losses}} - \overbrace{\eta I}^{\text{viral-lysis}} - \overbrace{\psi IG}^{\text{grazing}} \\ \frac{dV}{dt} &= \overbrace{\beta \eta I}^{\text{viral production}} - \overbrace{\phi(S+I)V}^{\text{adsorption}} - \overbrace{m_{v1}V}^{\text{viral losses}} - \overbrace{m_{v2}V^2}^{\text{higher-order losses}} \\ \frac{dG}{dt} &= \epsilon \frac{N_P}{N_G} \overbrace{\psi(S+I)G}^{\text{grazing}} - \overbrace{m_{g1}G}^{\text{specialist/generalist grazing}} - \overbrace{m_{g2}G^2}^{\text{higher-order losses}}, \end{aligned} \quad (1)$$

where

$$\mu(t) = \mu_{ave} ( 1 + \delta_{\mu} \sin( 2\pi(t + \delta_t) ) ). \quad (2)$$

*Prochlorococcus* have a diel-driven population division rate  $\mu(t)$  whose proportional amplitude and phase are set by parameters  $\delta_{\mu}$  and  $\delta_t$ , and  $t=0$  represents 06:00:00 local time (where  $t$  is measured in days). To aid model fitting we constrain  $\delta_t$  such that peak division rate occurs at night. *Prochlorococcus* have a linear loss rate  $m_{p1}$  and a nonlinear loss rate,  $m_{p2}$ , dependent on total phytoplankton abundance to implicitly represent niche competition [40]. Viruses infect susceptible *Prochlorococcus* at a rate  $\phi$  and release  $\beta$  new virions into the environment upon cellular lysis following the latent period (defined by the reciprocal of the lysis rate as  $\frac{1}{\eta}$ ). Grazing upon *Prochlorococcus* is non-preferential with respect to infection status and occurs at a rate  $\psi$  with a Gross Growth Efficiency (GGE)  $\epsilon$  proportional to the fraction of nitrogen contents in a *Prochlorococcus* cell ( $N_P$ ) and a grazer ( $N_G$ ). We introduce  $m_{g1}$  as a parameter to denote whether grazers act as specialists or generalists, where the term represents net additional gains to the grazer from non-*Prochlorococcus* prey sources after accounting for respiratory costs. We represent a specialist strategy via  $m_{g1} \geq 0$ , assuming that *Prochlorococcus* cells are the main source of grazer sustenance. In contrast, we represent a generalist strategy via  $m_{g1} < 0$ , implying that grazers have a net positive growth rate even in the absence of  $S$  or  $I$  via the consumption of other phytoplankton, heterotrophic bacteria, or grazers (via intraguild predation). Grazer and viral losses are both characterised by a nonlinear loss term to avoid structurally biasing the model to favour one of these *Prochlorococcus* predators [28] and to avoid competitive exclusion. A full list of parameters and further details are shown in the Supplementary Information.

## 2.3 Ecological model of phytoplankton communities including size structure (ECLIPSS)

The ECLIPSS model is illustrated in Figure 1b; it represents a *Prochlorococcus* population structured by differences in cell size and infection status: cells that are susceptible to viral infection ( $S$ ) and cells that are infected ( $I$ ) by viruses ( $V$ ). Both  $S$  and  $I$  are grazed upon by a general grazing class ( $G$ ).  $S$ ,  $I$ ,  $V$  and  $G$  represent population abundances. The ecological dynamics can be written, for a discrete number of size classes in matrix form as:

$$\begin{aligned}
 \frac{d\vec{S}}{dt} &= \overbrace{2\vec{B}\vec{S}}^{\text{birth}} - \overbrace{\vec{D}\vec{S}}^{\text{division}} + \overbrace{\vec{R}(E)\vec{S}}^{\text{growth}} - \overbrace{\vec{\lambda}(E)\vec{S}}^{\text{respiration}} - \overbrace{m_{p1}\vec{S} - m_{p2}\vec{S}M}^{\text{losses}} - \overbrace{\vec{\phi}\vec{S}V}^{\text{infection}} - \overbrace{\vec{\psi}\vec{S}G}^{\text{grazing}} \\
 \frac{d\vec{I}}{dt} &= \overbrace{\vec{R}(E)\vec{I}}^{\text{growth}} - \overbrace{\vec{\lambda}(E)\vec{I}}^{\text{respiration}} - \overbrace{m_{p1}\vec{I} - m_{p2}\vec{I}M}^{\text{losses}} + \overbrace{\vec{\phi}\vec{S}V}^{\text{infected}} - \overbrace{\vec{\psi}\vec{I}G}^{\text{grazing}} - \overbrace{\eta\vec{I}}^{\text{viral-lysis}} \\
 \frac{dV}{dt} &= \overbrace{\beta[\eta\vec{I}]}^{\text{viral production}} - \overbrace{\vec{\phi}\vec{S}V - \vec{\phi}\vec{I}V}^{\text{adsorption}} - \overbrace{m_{v1}V}^{\text{viral losses}} - \overbrace{m_{v2}V^2}^{\text{higher-order losses}} \\
 \frac{dG}{dt} &= \overbrace{\epsilon[\vec{N}_S\vec{\psi}(\vec{S} + \vec{I})]}^{\text{grazing}} \frac{G}{N_G} - \overbrace{m_{g1}G}^{\text{specialist/generalist grazing}} - \overbrace{m_{g2}G^2}^{\text{higher-order losses}}
 \end{aligned} \tag{3}$$

where vectors are shown in bold and accented by right arrows, and matrices are capitalised, bold and double underlined. Total *Prochlorococcus* carbon biomass is defined as  $M = \sum (\vec{C}_S(\vec{S} + \vec{I}))$ , where the carbon biomass of a *Prochlorococcus* cell is denoted as  $\vec{C}_S$ . Both  $\vec{C}_S$  and the nitrogen biomass of a cell,  $\vec{N}_S$ , are cell size-dependent (see Supplementary Information S1). The nitrogen biomass of a grazer cell is denoted as  $N_G$ . For simplicity, time dependencies on incoming photosynthetically active radiation,  $E$ , and biological state variables  $S$ ,  $I$ ,  $V$  and  $G$  are suppressed. A full list of parameters and further details are detailed in Supplementary Information S1. Key differences between the ECLIP and ECLIPSS models are that ECLIP has a sinusoidal diel-driven division rate, while in ECLIPSS we incorporate a light response that drives cellular volumetric growth and respiration (both absent in ECLIP). In ECLIPSS division rate is size-dependent such that a division event creates two daughter cells each with a volumetric size equal to half that of the mother cell. Additionally, in ECLIPSS, we model size-dependent encounter rates between *Prochlorococcus* and heterotrophic nanoflagellate grazers ( $\vec{\psi}$ ), and T4- and T7-like cyanophages ( $\vec{\phi}$ ).

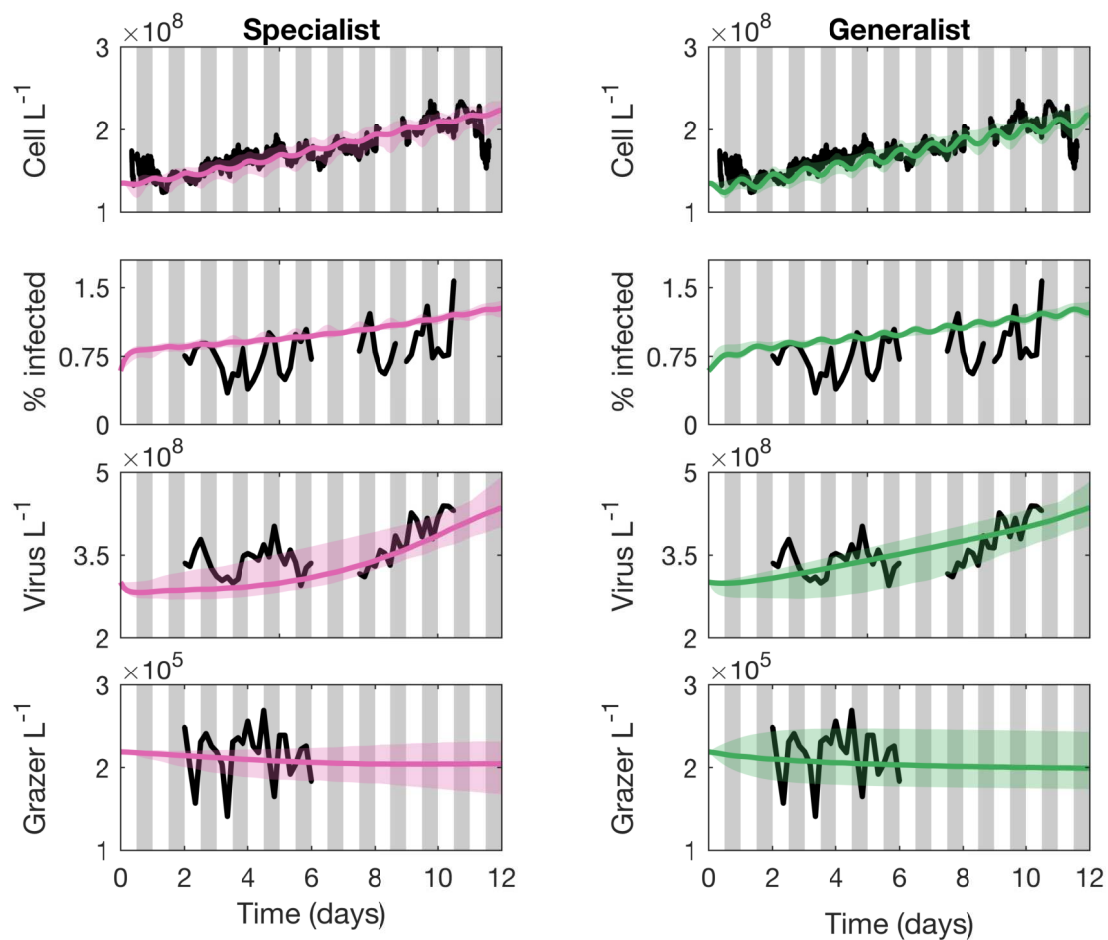
### 3 Results

#### 3.1 Ecological model of phytoplankton communities with viral and grazer mediated predation (ECLIP) and model-data integration

To explore the potential coexistence dynamics of *Prochlorococcus*, viruses and grazers, we first attempted to fit the ECLIP model using a Markov chain Monte Carlo (MCMC) approach given parameter bounds in biologically realistic ranges (see Table S2 for parameters and the Supplementary Information S2 for information on the MCMC fitting procedure). Since our data spans several orders of magnitude, and to avoid preferential fitting of a particular timeseries, we define error as the sum of the average relative differences between the model and empirical timeseries for each signal (*Prochlorococcus* cells, %-infected *Prochlorococcus* cells, T4- and T7- like cyanophages, and heterotrophic nanoflagellates), subject to a quadratic penalty (see details in Supplementary Information, equation S25). The models are fitted against detrended empirical data, so for visualisation we add this trend to the model simulations. The results from the best-fit specialist and generalist ECLIP model are shown in Figure 2. Both classes of ECLIP models were able to reproduce the magnitudes of the different data signals and produced fits with similar fitting error (see Figure S5). Hence, we find that a relatively simple multitrophic model can recapitulate multi-trophic populations dynamics in the NPSG.

Despite the strong agreement with population dynamics, the specialist and generalist models differ significantly in the posterior parameter values identified by the MCMC procedure (see Figure S5). Some life-history traits converge to similar values in parameter space, such as the phasing of division, which corresponds to the peak division rate occurring between 00:00:00 and 00:30:00. However, modeled *Prochlorococcus* abundances did not fit the phasing of the empirical data without constraining the model to ensure peak division rate occurs at night. *Prochlorococcus* division is expected to peak at night after daytime photosynthesis-driven growth. The failure of ECLIP to identify this phasing provides motivation for including a more mechanistically driven model of division (explored in the next section). In other cases, MCMC parameter distributions appear bimodal (e.g. adsorption rates, lysis rates), multi-modal (e.g. grazing encounter rates), some with no clear mode (e.g. GGE), and others that differ between specialist and generalist models. For example, specialist models tend to have a larger lysis rate and grazing gross growth efficiency than do generalist models (see Supporting Information Section S2). Altogether, these findings suggest that the ECLIP model exhibits parameter entanglement and “sloppiness” [41], i.e.





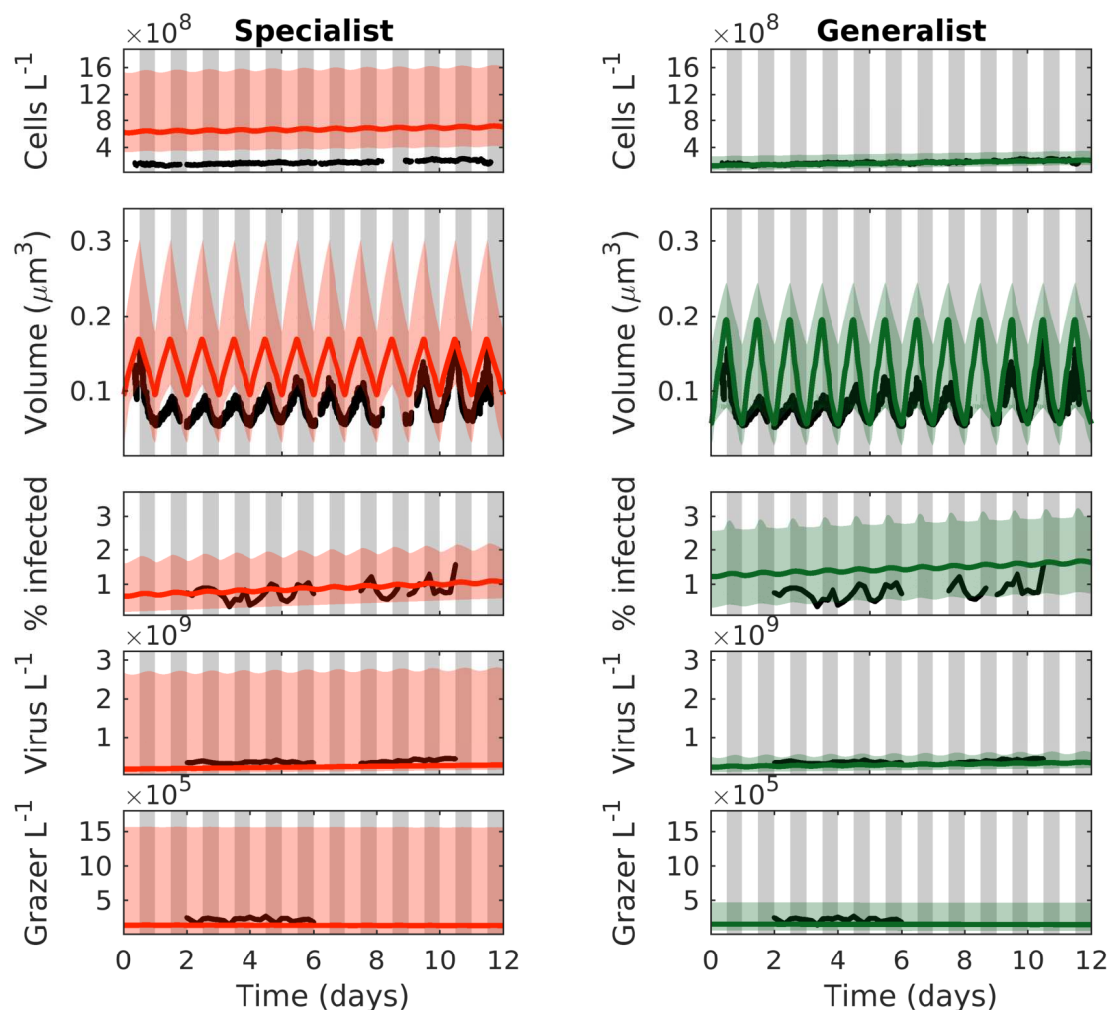
**Figure 2: Specialist and generalist ECLIP model fit to empirical data.** The best fitting specialist (left) in pink and generalist (right) in green ECLIP models are compared against empirical data in black. Model lines represent the median MCMC solution within the full range of solutions found by the converged chains shown as bands. Data signals include *Prochlorococcus* cell abundances (top), the percentage of infected *Prochlorococcus* cells, the abundance of free viruses and the abundance of heterotrophic nanoflagellate grazers (bottom). The models were fitted against detrended data; for visualisation we have added these trends to the model solutions. Grey bars indicate hours of darkness.

many parameter sets can lead to similar system behaviour. Further evidence for sloppiness is suggested by the parameter variability between different MCMC chains which all exhibited small and similar fitting errors. We interpret these findings to mean that despite the ability of ECLIP to recapitulate baseline dynamics (but not necessarily oscillation features of cells and infected cells over the diel cycle), we must be cautious in conflating fits with mechanism. Instead, the parameter sloppiness suggests the need to include additional data types (e.g., *Prochlorococcus* size structure) in order to differentiate between biological mechanisms (e.g., the relative contributions of lysis, grazing, or other factors to phytoplankton loss).

### 3.2 Size-structured ecological model (ECLIPSS) and model-data integration

The size-structured ECLIPSS model includes dynamics of size-structured susceptible and infected *Prochlorococcus* cells, cyanophages and heterotrophic nanoflagellates. Hence, in identifying parameters within ECLIPSS we augment the abundances of total cells, viruses, and heterotrophic nanoflagellates with the mean average volume of *Prochlorococcus* (as measured using SeaFlow; see Materials and Methods). We simulate the models using 20,000 parameter sets obtained by Latin Hypercube sampling (LHS) in biologically realistic ranges. Of these, 400 simulations produced feasible results (those that lead to stable coexistence of *Prochlorococcus*, cyanophages and grazers) under the specialist model and 1,094 simulations produced feasible results under the generalist model. These low rates of feasibility given random

parameter sampling are consistent with prior analyses of a multitrophic model without light-driven dynamics [29]. We used a subset of model runs with the smallest total error to constrain our parameter ranges; and used these to generate 10,000 new parameter sets (see Supplementary Text). Of these, 1,886 simulations produced feasible results under the specialist model and 2,512 simulations produced feasible results under the generalist model. The best fit from the generalist model has an error of 0.1584, whilst the best fit from the specialist model has an error of 1.0414 (Supplementary Information, Figure S12). We cannot directly compare fitting error between ECLIP and ECLIPSS models due to the addition of fitting ECLIPSS to cell volume, however the errors in the ECLIPSS models are larger per population type on average than those found in the optimised ECLIP models. We note these fits improve upon those found in the initial parameterisation of ECLIPSS (Supplementary Information, Table S1) shown in Supplementary Information Figure S4.



**Figure 3: The best fitting specialist and generalist ECLIPSS models.** The best fitting specialist (left) in red and generalist (right) in green ECLIPSS models are denoted in solid lines and enveloped in the bounds of the best ten fitting models and compared against empirical data in black. Data signals that are compared to the data for fitting are *Prochlorococcus* cell abundance (top), mean *Prochlorococcus* cell volume, the percentage of infected *Prochlorococcus* cells, the abundance of free viruses and the abundance of heterotrophic nanoflagellate grazers. The figure shows the bounds and mean of the top ten specialist and generalist models. The models were fitted to detrended data; for visualisation we have added these trends to the model solutions. Grey bars indicate hours of darkness.

The agreement between model and data in Figure 3 suggest that the core feedback mechanisms are

sufficient to jointly explain *Prochlorococcus* abundances and size structure, viral infection rates, viral abundances, and grazer abundances at sub-daily resolution. Nonetheless, differences appear between the best fitting specialist and generalist model states and the data; as well as between the model outcomes themselves. In the specialist model *Prochlorococcus* cell abundances appear about an order of magnitude larger than those observed and the model population has a larger average volume than that found in the data. On the other hand, the generalist model broadly captures the magnitude of *Prochlorococcus* cell abundance; and captures the diel variations in average cell volume. Both ECLIPSS models appear close to the observed magnitudes of  $\approx 1\%$  infected cells [20]. Both ECLIPSS models were broadly able to characterise the magnitudes of free T4- and T7-like cyanophages and heterotrophic nanoflagellate grazer abundances, though there was large variation amongst the best fitting specialist models. We note that the model predicted dynamics of *Prochlorococcus*, free virus abundance, heterotrophic nanoflagellate abundance, and the percentage of infected cells do not always recapitulate the magnitude of diel oscillations observed. This gap may point to direct light-mediated modulation of life history traits and behavior [16, 17, 18, 19, 20, 21], and/or unaccounted for feedback mechanisms in the model structure. Like our analysis of ECLIP, we find differences in parameter sets between the best fitting generalist and specialist ECLIPSS models that is then recapitulated via analysis of the full suite of coexisting LHS parameter sets (Supplementary Information, Figure S9 and S12-S15) Hence, rather than conclude that a particular parameter set provides the most explanatory power to observed patterns, we seek to identify mechanisms common to those parameter sets that fit the data nearly equally well.

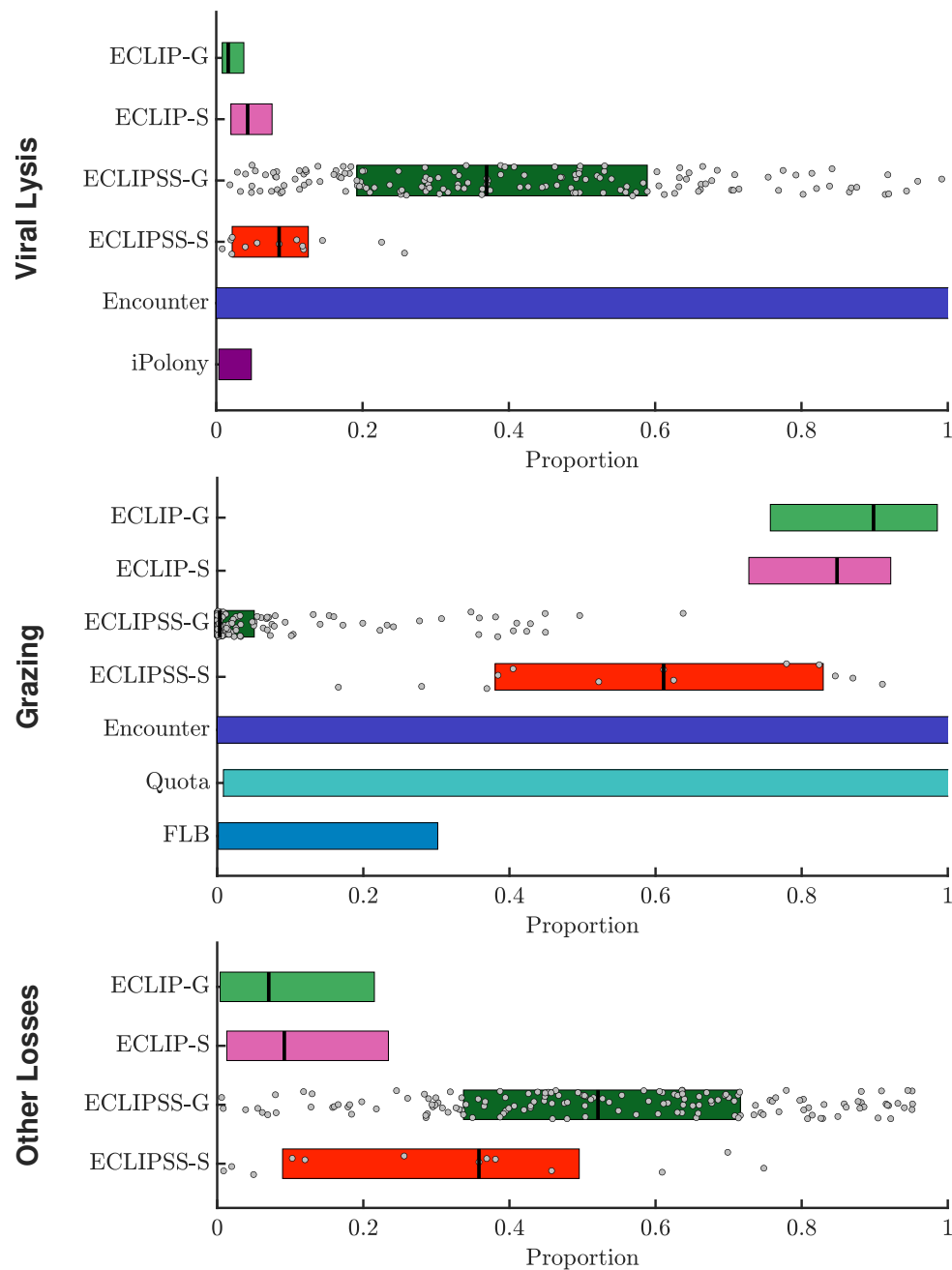
### 3.3 Partitioning the daily losses of *Prochlorococcus* between top-down and other effects

We revisit the best fitting specialist and generalist ECLIP and ECLIPSS models shown in Figures 2 and 3 to analyse the predicted partitioning of mortality among grazing by heterotrophic nanoflagellates, viral-induced lysis by T4- and T7-like cyanophages, and other sources. We expect that grazing may represent a larger proportion of losses in the models in which heterotrophic nanoflagellates act as specialists such that *Prochlorococcus* are their only prey source. For ECLIP we evaluate models from across the best chains, which all had small fitting errors; while in ECLIPSS, due to a paucity of model solutions with low error, we focus our analysis on models with a fitting error less than 2 (the top 13 specialist and 154 generalist models).

In Figure 4 we compared our model-inferred estimates of relative mortality with additional independent estimates from field based (iPolony measurements and fluorescently labelled bacterial (FLB) uptake estimates) or theoretical (encounter and quota) methods (see details in Supplementary Information S4). Theoretical estimates via the encounter and quota methods have significant variability due in part to uncertainties in life history traits. As such, viral lysis or grazing could potentially explain daily *Prochlorococcus* losses. However, direct estimates suggest that both viral-induced lysis by T4- and T7-like cyanophages and grazing by heterotrophic nanoflagellates only contribute a comparatively small amount to *Prochlorococcus* cell losses. We estimate that less than 35% of *Prochlorococcus* cell losses are explained by the combination of viral-induced lysis by T4- and T7-like cyanophages (estimated by iPolony) and grazing by heterotrophic nanoflagellates (estimated by FLB). Across models, we find that grazing makes up a larger proportion of *Prochlorococcus* cell losses in models in which heterotrophic nanoflagellates act as specialist consumers. In general, we find that viral-induced lysis is responsible for only a small proportion of losses in both models in agreement with Mruwat et al. [20]. Grazing by heterotrophic nanoflagellates, by contrast, made up the majority of *Prochlorococcus* cell losses across both the ECLIP models and in the specialist grazer ECLIPSS model. However, viral-induced lysis was inferred to be larger in the generalist ECLIPSS model, while grazing was much smaller – and more in line with estimates made via FLB measurements – relative to the other models. Combining model-data integration with direct measurements of top-down effectors, we found that the model-inferred combination of viral-induced lysis by T4- and T7-like cyanophages and grazing by heterotrophic nanoflagellates typically do not sum up to equal total *Prochlorococcus* cell losses on a daily basis.

Instead, our model-data integration suggests that other sources of *Prochlorococcus* cell losses are non-negligible, accounting for between 5% and 55% across the different model types. One possibility is that mortality partitioning could be linked to how quickly *Prochlorococcus* cells are expected to turnover in the NPSG. We note that the best set of models supports a large range of per capita *Prochlorococcus* cell loss rates of between nearly 0 and 1.1 per day; suggesting that perhaps the variability in other loss rates are coupled to magnitude of the estimated birth/death rates. However, contrary to this hypothesis, we do not have evidence linking total loss rates and mortality partitioning between viral-induced lysis, heterotrophic





**Figure 4: Relative importance of viral lysis, grazing, and other effects on total *Prochlorococcus* mortality.** The proportion of mortality partitioned between viral-induced lysis (top panel), grazing (middle panel), and other (bottom panel) sources for the best ECLIP specialist (ECLIP-S) and generalist (ECLIP-G) and ECLIPSS specialist (ECLIPSS-S) and generalist (ECLIPSS-G) models and other measures of relative mortality. For ECLIP the results from all chains are shown. For ECLIPSS only models with a fitting error less than 2 are included (and are additionally shown as individual points). Bars in these panels denote mortality rate proportions associated with the best ECLIP and ECLIPSS models marked by their 25, 50 and 75% quantiles. Other measures of relative mortality are given via theoretical encounter rates and quota based estimates, direct measurements of viral infection (iPolony), and Fluorescently Labelled Bacteria (FLB) incubation measurements (see Supplementary Information Section S4 for details).

nanoflagellate grazing and other processes in either the ECLIP or ECLIPSS models (Supplementary Information, Figure S16). Taken together, model-data integration suggests that viral-induced lysis accounts for a relatively small proportion of *Prochlorococcus* cell losses in the NPSG. Model results also suggest that the sum of losses via viral-induced lysis and heterotrophic nanoflagellate grazing do not account for

the majority of *Prochlorococcus* cell losses - in contrast to prior work that generally aims to partition loss rates in terms of established top-down drivers. Together, both model-data fits and independent estimates of top-down mortality suggest that other forms of losses beyond grazing by heterotrophic nanoflagellates and lysis by T4/T7-like cyanophages may be critical in shaping daily phytoplankton rhythms.

## 4 Discussion

We developed and analysed two multitrophic community ecology models, ECLIP and ECLIPSS, consisting of *Prochlorococcus*, viruses, and grazers to investigate the feedback mechanisms and ecological drivers of oligotrophic ocean microbial population dynamics on diel timescales. Both ECLIP and ECLIPSS include light-driven phytoplankton growth, where phytoplankton can be infected and lysed by viruses, or grazed upon by nanoflagellate grazers. ECLIPSS explicitly incorporates phytoplankton size-structure including size-mediated division and encounter with viral and grazer predators, while ECLIP considers the ecological interactions independent of size-structure. We found that both ECLIP and ECLIPSS were capable of generating coexisting populations of cyanobacteria, viruses infecting cyanobacteria, and grazers, and that the suite of model solutions encompasses the population abundances observed in the NPSG (Supplementary Information, Figure S11). Despite strong agreement with data, we found that underlying process rates and life-history traits differed significantly amongst best-fitting ECLIP models. In an effort to resolve identifiability problems, we found that ECLIPSS was able to jointly explain the abundance dynamics of *Prochlorococcus*, virus particle abundances, infected cells, and grazers. By combining model-data fits with direct measurements of mechanistic interactions we infer that the combination of lysis and grazing does not account for daily *Prochlorococcus* losses. Instead, we find that a significant - and potentially majority - of *Prochlorococcus* losses are due to other mechanisms beyond that of lysis by T4- and T7-like cyanophage and nanoflagellate grazing.

The development of a multitrophic model focused in on a portion of a complex, surface ecosystem. In doing so, we integrated our models with field observations to understand interactions of T4- and T7-like cyanophages as well as nanoflagellate grazers with *Prochlorococcus*, the most abundant phytoplankton at our study site. Prior work has assumed that *Prochlorococcus* is the predominant prey for heterotrophic nanoflagellate grazers - but such an assumption had the potential to bias interpretations of direct ingestion experiments (and the interpretation of total loss rates). Instead, it is likely that heterotrophic nanoflagellate grazers rely on other food sources as part of a generalist feeding strategy. To evaluate this possibility, the ECLIP framework included the potential for model outcomes to span generalist to specialist strategies. However, we found a critical identifiability problem: a broad range of model classes could explain population dynamics in the absence of size structure. These identifiability problems within the ECLIP framework prevented us from distinguishing whether heterotrophic nanoflagellates act primarily as specialist grazers of *Prochlorococcus*, or ingest a broader range of prey items. Turning to ECLIPSS, we found that assuming grazers are generalist, rather than specialists, leads to improvements in fit quality. Notably, ECLIPSS was able to recapitulate daily dynamics of total *Prochlorococcus* cell abundances and size structure, as well as top-down populations including the fraction of infected cells, T4- and T7-cyanophage abundances, and heterotrophic nanoflagellate grazer abundances.

Overall, the fit of model dynamics to NPSG measurements enabled us to examine how much of *Prochlorococcus* mortality can be ascribed to viral lysis, heterotrophic nanoflagellate grazing, or other sources. In partitioning *Prochlorococcus* mortality we found different outcomes across the different model scenarios and independent auxiliary estimates (Figure 4). Indirect estimates via encounter rate or quota-based theory are poorly constrained and are limited by our current knowledge of microbial ecological life-history traits. However, fitting dynamical models, ECLIP and ECLIPSS with the same initial parameter ranges, to the data resulted in more constrained mortality estimates. Both the ECLIP and ECLIPSS models predicted that other sources of mortality were likely important in explaining community diel patterns. Assuming that grazers function as generalists, we found that compatible model-data fits predict that other sources of mortality can account for over 50% of daily *Prochlorococcus* mortality. We found that direct estimates of mortality from grazing incubation experiments and infected cell measurements also provide evidence that heterotrophic nanoflagellate grazing and T4- and T7- like viral-induced mortality do not account for all *Prochlorococcus* losses in the NPSG. Quantifying the relative importance of mortality processes beyond conventional top-down effects (grazing and lysis) is critical for understanding how grazers and viruses contribute to mortality and energy transfer in marine microbial communities [2, 23, 24, 28].

## Box 1 | Potential mechanisms to explain other losses of *Prochlorococcus*

### Undercharacterised ecological interactions

Mixotrophic nanoflagellates are expected to contribute to the “other” loss of *Prochlorococcus*, but measurements suggest this is a small contribution relative to the heterotrophic nanoflagellates [21, 42]. Unknown viruses (like RNA viruses), as well as other grazer types including larger consumers may be contributing to the ‘other mortality’ terms in our models. In addition, the model includes viral-induced and grazing-induced losses; however in both cases we utilize a contact-driven model, analogous to ‘Type I’ functional responses. Mechanistic changes in functional responses and/or responses to light may drive distinct interaction rates (and aggregate mortality) even given the same set of viruses and grazers.

### Aggregation and sinking of picoplankton

Some losses may be due to *Prochlorococcus* aggregation, particle attachment, and subsequent sinking out. Picoplankton (and their viruses) have been suggested to be an important contributor to export in large scale analyses e.g. [43, 44], though Guidi et al. [44] suggest *Synechococcus* rather than *Prochlorococcus* abundances appear correlated to export. Additionally, less is known about the microscale processes that might lead to export contributions from picoplankton – *Prochlorococcus* cells can be grown and sustained in laboratory culture for month-long experiments, suggesting sedimentation of *Prochlorococcus* cells may be limited. Generally, conceptual models of ocean ecology do not include sinking out of picoplanktonic populations. However, particle aggregation could be stimulated via processes such as sloppy feeding or viral lysate [28, 45]. As viral lysate from picocyanobacteria is partially labile [46, 47] aggregation and sinking could be stimulated indirectly by viral lysate via heterotrophic bacterial growth. Particle attachment and aggregation could also be stimulated by TEP (Transparent exopolymer particles), which is produced in xenic *Prochlorococcus* cultures [48, 49]. TEP production appears to be stimulated by high light intensity, characteristic of the surface ocean, and appears to be linked to loss rates [48].

### Physiological stress(es)

Other potential loss mechanisms include physiological stress which can be induced, for example, by high-level irradiance associated with ultraviolet radiation [50, 51] and refraction of light through surface waves that could lead to photodamage via the flashing effect [52, 53]. Other abiotic factors such as nutrient deficiency [54], metal toxicity [55, 56], and thermal variations [57] can also contribute to increased stress, though we do not expect thermal variations to be important in the NPSG. Stresses often result in the generation of reactive oxygen species, that cause oxidative stress and can lead to DNA damage. Reactive oxygen species are also used as a signalling pathway for programmed cell death in photosynthetic microbes [58, 59, 60], though there is a lack of evidence for a programmed cell death pathway in *Prochlorococcus* and sympatric heterotrophic bacteria are thought to alleviate this type of stress [61, 62]. We note that there is also a lack of evidence for toxin-antitoxin systems, which are typically common across bacteria and archaea and could act as a pathway to additional cell loss, in *Prochlorococcus* [63].

### Population heterogeneity

Microbes experience senescence and aging, leading to intracellular accumulation of damage through their life cycle [64], which may lead to asymmetric division [65]. Unlike other microorganisms, *Prochlorococcus* does not have resting stages and rely on heterotrophic bacteria to survive nutrient starvation in cultures [66], though the types of heterotrophic bacteria they form associations with *in situ* differ from those found in culture which tend to be copiotrophic [67]. Heterogeneity within the *Prochlorococcus* population may mean there are differing mortality responses at the individual or strain-level vs. at the scale of total population. *Prochlorococcus* populations are combinations of ecotypes which respond differently to environmental stressors [68, 69, 70, 71, 72, 73]. Competition between these ecotypes and with other phytoplankton in the same niche (like *Synechococcus* or picoeukaryotes) may also increase stresses, especially in an oligotrophic environment like the NPSG where nutrients are limited.

The finding that model-data fits impute other sources of mortality as quantitatively significant suggests that we may be missing important feedback mechanisms in our representation of marine surface community dynamics. Indeed, phytoplankton loss from the surface oceans is likely to include factors beyond grazing and viral-induced lysis [74, 75]. In Box 1 we review some potential mechanisms that may contribute to the unaccounted losses of *Prochlorococcus*, including ecological feedback that leads to distinct functional and/or light-driven responses, aggregation and/or sinking, stress, population heterogeneity, as well as the possibility of having missed other sources of top-down mortality (e.g., uncharacterised viral infections or grazing by larger organisms). Similarly, our measurements may miss heterogeneities within the *Prochlorococcus* population that mask our ability to interpret mortality. Alternatively, we may be missing entire mechanisms of loss such as via particle aggregation and sinking out, or via physiological stresses on cells. Investigating whether alternative mechanisms of *Prochlorococcus* losses could be significant warrants future research, and may be critical to improving understanding of how biomass and nutrients are transferred through marine food webs.

Both the ECLIP and ECLIPSS modeling frameworks come with caveats, despite the inclusion of multiple populations and interaction types. First, we have focused on the impacts of direct, light-driven forcing of cyanobacterial division (in ECLIP) and growth leading to division (in ECLIPSS). Hence, oscillations arising in other components reflect a combination of instabilities that can arise in nonlinear population models as well as the cascading impacts of such oscillations on the community. Direct incorporation of diel impacts on grazing [16, 18, 21] or viral traits (e.g., adsorption) [76, 19] may be required to mechanistically understand population dynamics on sub-daily timescales. Second, we have used two focal processes to examine ways that carbon and nutrients in basal picoplankton are transferred, e.g., up through the food chain via grazing or retained in the microbial loop due to viral lysis (aka the viral shunt) [77, 78]. This dichotomy reflects a potential tension in the extent to which primary production stimulates the biological pump. Extensions to the current model might also examine the mechanistic process of export explicitly, whether through coupling grazing to the generation of particles and/or examining the extent to which viral lysis generates aggregates which can be exported to the deep oceans via the viral shuttle [45, 79]. Finally, our work has implicated an accounting challenge in quantifying the balance of *Prochlorococcus* growth and losses. Despite the daily growth and division of cells, the overall abundance remains tightly constrained – our work suggests that this constraint may in fact depend on other factors beyond conventional mechanisms of loss ascribed to T4- and T7-like cyanophage and nanoflagellate grazers.

In summary, this multi-trophic modeling framework provides an opportunity to test the limits of our understanding of feedback mechanisms of microbial communities in the surface open ocean. In doing so, the model has provided additional support for the hypothesis that relatively high viral abundances can be compatible with relatively low infection (and mortality) rates, how grazer generalism may be key to understanding *Prochlorococcus* abundances, while also identifying a key direction for future work: identifying the potentially ‘missing mortality’ of phytoplankton at the base of the marine food web. In moving forward, *in situ* observations will be needed to probe aggregation and sinking, autolysis, programmed cell death, or other forms of loss of *Prochlorococcus* in the surface ocean and to understand the feedbacks of variation in cyanobacterial growth and loss in a changing ocean.

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## Materials and Methods

### Model availability

Both the ECLIP and the ECLIPSS models are coded in MATLAB. Core code for running model simulations, analysis and plotting is archived on Zenodo ([80]).

### Empirical data

We use data collected from the HOE-Legacy 2A cruise in Summer 2015. In all Figures, the 12 days shown represent the time between 06:00:00 24th July 2015 and 06:00:00 August 5th 2015 local time.

- SeaFlow measurements of *Prochlorococcus* cell abundance, size-structure and carbon biomass:  
The SeaFlow continuously measures forward scattering, red and orange fluorescence intensities of particles ranging in size from ~0.4 to 4  $\mu\text{m}$  in diameter from underway samples every 3 minutes. A combination of manual gating and statistical methods was used to identify *Prochlorococcus* based on forward scatter (457/50 bandpass filter), red fluorescence (572/28 bandpass filter) and orange fluorescence (692/40 band-pass filter) relative to 1- $\mu\text{m}$  calibration beads (Invitrogen F8823). Cell diameter of individual cells was estimated from SeaFlow-based light scatter by the application of Mie light scatter theory to a simplified optical model, using an index of refraction of 1.38 [81, 82]. Data was obtained via Simons CMAP [83].

- Virus abundance and infection:

Samples for virus abundance and infection were collected every 4 hours at 15 m depth using a CTD-rosette equipped with 12 L niskin bottles [20]. Samples for cyanophage abundances (40 mL) were filtered through a 0.2  $\mu\text{m}$  syringe top filter, flash frozen, and stored at  $-80^{\circ}\text{C}$ . Samples for infected cells (40 mL) were filtered through a 20  $\mu\text{m}$  nylon mesh, fixed with electron microscopy grade glutaraldehyde (0.125% final abundance), incubated for 30 minutes in the dark at  $4^{\circ}\text{C}$ , flash frozen, and stored at  $-80^{\circ}\text{C}$ . Cyanophage abundances were analysed using the polony method for T7-like [84] or T4-like [85] cyanophage families. Virally infected *Prochlorococcus* was quantified using the iPolony method [20] in which *Prochlorococcus* cells were sorted with a BD Influx cytometer and screened for intracellular T4-like and T7-like cyanophage DNA.

- Measurements of heterotrophic nanoflagellates:

Samples for nanoplankton (protists 2-20  $\mu\text{m}$  in diameter) abundances were collected every 4 hours at 15 m depth [21]. Preserved samples were stored at  $4^{\circ}\text{C}$ . Slides were prepared from preserved samples within 12 hours of sampling by filtering 100 mL subsamples down to 1 mL onto blackened 2  $\mu\text{m}$ , 25 mm polycarbonate filters and staining the samples with 50  $\mu\text{L}$  of a 4'-6'diamidino-2-phenylindole (DAPI, Sigma-Aldrich, St. Louis, MO) working solution ( $1\text{ mg mL}^{-1}$ ) for 5-10 minutes in the dark [86]. Stained samples were filtered and rinsed; filters were placed on glass slides with a drop of immersion oil and coverslip, then sealed with clear nail polish. Slides were stored at  $-20^{\circ}\text{C}$  until analysis. Heterotrophic nanoplankton abundances were counted using epifluorescence microscopy from triplicate slides, and differentiated from photo/mixotrophic nanoplankton by the lack of chlorophyll a autofluorescence in plastidic structures when viewed under blue-light excitation [21].