# **Dynamic macromolecular composition and high exudation rates in**

# 2 **Prochlorococcus**

3

4 Dalit Roth-Rosenberg<sup>1</sup>, Dikla Aharonovich<sup>1</sup>, Anne-Willem Omta<sup>2</sup>, Michael J. Follows<sup>2</sup> and Daniel
5 Sher<sup>1</sup>

<sup>1</sup> Department of Marine Biology, Leon H. Charney School of Marine Sciences, University of Haifa,
 Israel, <sup>2</sup> Department of Earth, Atmospheric and Planetary Sciences, Massachusetts Institute of
 Technology, USA

9

# 10 Abstract

Every living cell is composed of macromolecules such as proteins, DNA, RNA and pigments or 11 12 cofactors. The ratio between these macromolecular pools depends on the allocation of resources within the organism to different physiological requirements, and in turn determines the elemental composition 13 of the organism and, potentially, how it may affect biogeochemical cycles of elements such as carbon, 14 15 nitrogen and phosphorus. Here, we present detailed measurements of the macromolecular composition of *Prochlorococcus* MIT9312, a representative strain of a globally abundant marine primary producer, 16 17 as it grows and declines due to N starvation in laboratory batch cultures. As cells reached stationary stage and declined, protein/cell decreased by ~30% and RNA/cell and pigments/cell decreased by an 18 order of magnitude. The decline stage was associated with the appearance of chlorotic cells which had 19 higher forward scatter (a proxy for cell size) but lower chlorophyll autofluorescence, as well as with 20 changes in photosynthetic pigment composition. Specifically, during culture decline divinyl-21 22 chlorophyll-like pigments emerged, which were not observed during exponential growth. These

23	divinyl-chlorophyll-like pigments were also observed in natural samples from the Eastern
24	Mediterranean. Around 80-85% of the carbon fixed by Prochlorococcus MIT9312 (but not of a
25	different strain, NATL2A) was released into the growth media as dissolved organic carbon under these
26	laboratory conditions. Broadly defined, the macromolecular composition of Prochlorococcus
27	MIT9312 is more similar to eukaryotic phytoplankton than to marine heterotrophic bacteria, suggesting
28	a different set of physiological constraints determines the macromolecular composition of these two
29	broad classes of marine microorganisms.

30

- 31 Keywords: Prochlorococcus, Photosynthetic pigments, Macromolecular pools, exudation,
- 32 ribosomes
- 33

34

#### 35 Introduction

36 Like every cell on Earth, phytoplankton and bacterioplankton are composed of several classes of 37 biological macromolecules that function together to maintain the cells structure and activity (Geider and La Roche 2002, Finkel et al. 2016). Proteins are the most abundant macromolecule (in terms of 38 mass), responsible for most of the metabolic functionality of the cells. DNA stores hereditary 39 information, whereas RNA (as mRNA or other regulatory RNAs) accesses this stored data and (as 40 41 rRNA and tRNAs) uses it to produce protein. Lipids form the membrane(s) that separate the cell from 42 its environment and, in the case of photosynthetic organisms, host the molecular machinery used to 43 harvest light for energy. This molecular machinery includes, in addition to proteins, also photosynthetic 44 pigments and other metabolites. Finally, various storage molecules (including some lipids, carbohydrates, polyphosphate and other storage molecules) allow cells to retain energy and elements 45 46 for future use, and a host of other metabolites are involved in all forms of cellular function. The relative 47 amount of each of these classes of macromolecules changes between different phytoplankton groups (Vargas et al. 1998, Geider and La Roche 2002, Finkel et al. 2016), and may also change within a 48 single type of organism in response to environmental conditions (Vargas et al. 1998, Liefer et al. 2019). 49 For example, the relative amount of chlorophyll-a per cell changes in response to light intensity (e.g. 50 (Moore et al. 1995)) and may also change in response to nutrient starvation (Lourenço et al. 1998, 51 52 Liefer et al. 2019). This affects the ability of photosynthetic cells to harvest light and fix carbon, 53 impacting cell physiology and likely the interactions of the cell with other organisms, e.g. through the 54 release of fixed organic carbon (Dubinsky and Berman-Frank 2001). Similarly, it has been suggested 55 that the number of ribosomes per cell is a strong determinant of cellular growth rate, and is itself 56 affected by temperature and nutrient (particularly phosphorus, P) availability (e.g. (Elser et al. 2003, 57 Garcia et al. 2016, Martiny et al. 2016)). Thus, to some extent, the macromolecular composition of the 58 cell is affected by, and may shed light on, the allocation of cellular resources to different functions.

The basic building blocks of biological macromolecules such as amino acids and nucleic acids have 59 distinct elemental composition, and the macromolecular composition of the cell is thus intimately 60 61 linked with its elemental composition (Geider and La Roche 2002, Finkel et al. 2016, Garcia et al. 62 2016, Liefer et al. 2019). In phytoplankton, proteins generally comprise the largest pool in terms of 63 cell biomass (dry weight) as well as nitrogen and carbon (N and C) pools (Lourenço et al. 1998, Vargas 64 et al. 1998, Geider and La Roche 2002, Finkel et al. 2016). Other macromolecules such as Nucleic 65 acids and pigments contribute less to total cell biomass yet contribute significantly to the pools of N 66 and phosphorus (P) (Geider and La Roche 2002, Finkel et al. 2016, Liefer et al. 2019). Thus, the 67 macromolecular composition of the cell will determine its elemental composition, which in turn will 68 determine the effect of the cell on biogeochemical cycles of elements. Over the last ~85 years (Redfield 69 1934) significant advancements have been made in our understanding of how the elemental ratios in 70 marine organisms change over time and space, how they are affected by cell physiology, and how they 71 are related to surrounding nutrient element concentrations (e.g. (Martiny et al. 2013, Gruber and 72 Deutsch 2014)). Significantly less is known about the dynamics of the macromolecular composition of 73 marine organisms, which underlie elemental composition. Similarly, most of the models of ocean 74 communities and their effect on biogeochemistry are formulated using elements as the model currency 75 rather than macromolecules (Omta et al. 2009, Goebel et al. 2010, Talmy et al. 2016). In order to 76 obtain a better understanding of the dynamics of the macromolecular composition of organisms, and 77 to link these to elemental ratios, experimental measurements are needed, particularly in abundant 78 marine organisms.

Here, we measure the cellular pools of major macromolecules (protein, DNA, RNA and pigments) in
laboratory batch cultures of *Prochlorococcus*, a highly abundant pico-cyanobacterium which is
responsible for ~8.5% of the oceanic photosynthesis (Partensky and Garczarek 2010, Flombaum et al.
2013, Biller et al. 2014). *Prochlorococcus*, as a clade, comprise multiple genotypes, each different in

its physiology, genome structure and oceanic niche (Johnson et al. 2006, Biller et al. 2014). Many 83 aspects of the physiology of the *Prochlorococcus* clade have bene intensively studied, ranging from its 84 85 elemental composition (Bertilsson et al. 2003, Martiny et al. 2013) through its photophysiology (Moore et al. 1995, Moore and Chisholm 1999, Ting et al. 2002, Steglich et al. 2003, Komatsu et al. 2016) to 86 its system-wide transcriptomic response to changes in environmental conditions (e.g. (Martiny et al. 87 88 2006, Tolonen et al. 2006, Thompson et al. 2011, Aharonovich and Sher 2016). Yet, detailed analyses 89 of the macromolecular composition of the cells are lacking, even though changes in the elemental 90 composition of *Prochlorococcus* that are observed in response to environmental conditions are in fact 91 due to changes in the macromolecular composition of the cells (E.g. (Bertilsson et al. 2003, Van Mooy 92 et al. 2006, Martiny et al. 2013, Martiny et al. 2016)). The strain we focus on, MIT9312, represents the 93 high-light adapted HL-II clade which is the most abundant clade in the surface waters of large parts of 94 the ocean (Bouman et al. 2006, Johnson et al. 2006). We chose to follow batch cultures of 95 Prochlorococcus MIT9312 under conditions where entry into stationary stage occurs when the N 96 source in the media is depleted for several reasons. First, N stress affects Prochlorococcus in large 97 parts of the ocean (e.g.(Saito et al. 2014)), and significant information is available as to the 98 physiological, transcriptomic and evolutionary responses of the cells to acute N starvation (Steglich et 99 al. 2001, Moore et al. 2002, Tolonen et al. 2006, Gilbert and Fagan 2011, McDonagh et al. 2012, Read 100 et al. 2017, Berube et al. 2019, Szul et al. 2019). Second, long-term analyses of batch cultures may 101 identify cellular physiological processes not seen when cells are growing exponentially or exposed to 102 sudden or short-term nutrient starvation (e.g. (Christie-Oleza et al. 2017, Roth-Rosenberg et al. 2019)). 103 Indeed, it is an open question to what extent stationary-phase and culture decline in batch cultures 104 represent conditions found in nature, yet we show that a pigment identified in our cultures during the 105 culture decline stage is found in the oceans, raising the possibility that some cells in nature are indeed 106 found in these physiological stages. Finally, we measured, in addition to the cellular macromolecular

L07	pools, also the concentrations of TOC in the media, extracting from these measurements estimates of
L08	the release of DOC from the cells. This release may be due to passive exudation, active excretion and
L09	cell lysis (reviewed by (Thornton 2014)). Some of the released macromolecules are available for

subsequent utilization by other organisms, whereas another part contributes to a huge pool of long-

111 lived oceanic DOM, potentially impacting global biogeochemical cycles (Hansell et al. 2009).

112

## 113 Materials and Methods

## 114 Growth conditions and experimental procedure

To follow the growth and macromolecular composition of Prochlorococcus over the various stages of 115 116 laboratory batch culture, axenic Prochlorococcus MIT9312 cultures were grown in 1 liter bottles of 117 Pro99 media (Moore et al. 2007) where the  $NH_4$  concentration was lowered to 100 $\mu$ M, leading to cessation of cell growth due to N starvation (Grossowicz et al. 2017). The experiment included 15 118 119 bottles, three of which were used for routine culture monitoring and the other 12 were used to collect samples at four time-points, corresponding to exponential growth (t=6 days), early stationary stage 120 121 (t=10 days), late stationary stage (t=13 days) and culture decline (t=15 days). Samples were also 122 collected from the starter culture to represent t=0. The number and timing of the samples collected for 123 full analysis were selected in order to provide appropriate data for subsequent modeling (Omta et al. 124 2017). This experimental design was used to minimize the chance for contamination of the axenic cultures with heterotrophic bacteria. The cell numbers of the cultures used for macromolecular analysis 125 were never statistically different from those of the cultures used for monitoring (Students t-test). 126

Prior to the beginning of the experiment, cells from a mid-exponential culture were counted by flow cytometry (FACSCantoII, BD), and diluted in the new media to an initial cell density of  $10^{6}$ /ml. Cultures were maintained under constant light (22 µE), at 24.5 ±1°C. 1mM NaHCO<sub>3</sub> was supplemented

130	at three different time points (T=0, T=7, T=11 days) to verify that the cultures are not carbon-limited
131	(Moore et al. 2007, Grossowicz et al. 2017). Bulk culture fluorescence was measured in samples
132	collected aseptically from the three monitoring bottles using a Carey Eclipse spectrofluorometer (Ex
133	440nm/Em 680nm). Samples for cell counting were fixed in 0.25% glutaraldehyde, kept in the dark

134 for 10 minutes and transferred to a -80°C freezer for subsequent flow cytometry analysis.

135

## 136 Measurements of macromolecular pools and photosynthetic pigments.

At each of the five time-points selected, cells were collected by filtration. For protein measurements, 137 138 cells were gently filtered on 0.22µm polycarbonate filters and kept in -80°C until extraction. For protein extraction, the filters were incubated in 500µl lysis buffer (50mM Tris pH6.8, 5mM EDTA, 2%SDS) 139 on a lab rotator at 37°C for 20'. Following incubation, samples were sonicated in water bath for 10', 140 centrifuged at 12,000g in a table-top micro-centrifuge for 5', and the supernatant liquid taken for 141 142 measurement using the Bicinchoninic acid assay (BCA, Sigma). Nucleic acids were collected similarly 143 on 0.22 µm Supor-200 Membrane Disc Filters (25 mm; Pall Corporation) and preserved in storage buffer (40 mM EDTA, 50 mM Tris pH 8.3, 0.75 M sucrose) and RNA Save (Biological Industries) for 144 145 DNA and RNA measurements, respectively. Samples were kept at -80°C until extraction. DNA was extracted from the filters according (Massana et al. 1997). Total RNA was extracted using the mirVana 146 miRNA kit (Ambion, Austin, TX, USA) as described in with lysozyme treatment for better lysis 147 148 (Tolonen et al. 2006), follow by removal of contaminating DNA from total RNA by Turbo DNase 149 (Ambion). Nucleic acids quantity measured using Qubit 2.0 (Invitrogen). For pigment analysis, samples were collected on Glass Fiber filters (25 mm GF/F, Whatman) and kept in -80C until 150 151 extraction. Glass fiber filters were used, rather than polycarbonate filters, as the latter are not 152 compatible with many extraction methods for pigments. Nevertheless, preliminary experiments using 153 a culture of Prochlorococcus MED4, which is similar in size to MIT9312, showed that with the vacuum pumps we used approximately 99.9% of the cells were retained on the GF/F membrane (Marmen et al. 154 155 2018), and thus the difference in pore size has minimal effect on the results. Cells were extracted in 100% methanol at 25°C for 2h. The cell extracts were clarified with Syringe filters (Acrodisc CR, 13 156 mm, 0.2 µm PTFE membrane, Pall Life Sciences), and kept at -20°C until analysis by column 157 158 chromatography on a C8 column (1.7 µm particle size, 2.1 mm internal diameter, 50 mm column 159 length, ACQUITY UPLC BEH, 186002877) using an ACQUITY UPLC system equipped with a 160 Photodiode Array detector (Waters). The chromatography protocol was based on LOV method 161 (Hooker et al. 2005) modified for UPLC. A linear gradient was applied over 14 minutes, where solvent 162 A was 70:30 methanol:0.5 M ammonium acetate, and solvent B was B. 100% methanol. The flow rate 163 was 0.5 ml/min, and the column and injection heating were set to 50°C and 30°C, respectively. To 164 identify and calculate pigments concentration, the following standards were used: Divinyl chlorophyll A (DVchlA), chlorophyll B (chlB), chlorophyll C (chlC), zeaxanthin (Zea), α- carotene (α-car), Divinyl 165 166 Protochlorophyllide (MGDVP), Pheophorbide a, Pheophythin a and Chlorophyllide a. All standards 167 were from DHI, Denmark.

#### 168 Measurements of inorganic nutrients and TOC

169 The GF/F filtrates (see above) were kept in polypropylene at -20°C until analysis. For measurements, 170 samples were adjusted to 25°C and diluted 4-fold in DDW to reduce the salt level. NH<sub>4</sub> (NH 3-N) was 171 measured using an HI 96715 meter (Hanna instrument), and PO<sub>4</sub> was measured by HI 96713 meter 172 (Hanna instrument). In both cases, low concentration kits were used. For Total Organic Carbon (TOC) 173 analysis, samples were collected into cleaned, pre combusted, acid washed (10% HCL) 40ml vials and 174 HCl was added to remove dissolved inorganic C before preservation at -20°C. Before analysis, samples 175 were combusted at 680°C using catalytic (platinum) oxidation method in oxygen-rich environment. 176 TOC analysis was performed using TOC-L analyzer (Shimadzu ASI-L Autosampler, Columbia, MD).

177 For inorganic carbon (DIC), samples were collected in 14ml dark glass vials with screw caps and 75µl (0.05% v/v) saturated HgCl<sub>2</sub> solution (Dickson et al. 2007) was added to the samples. For alkalinity, 178 60 ml samples were collected in glass vials. All samples were kept in  $4^{\circ}$ C until measurement. CO<sub>2</sub> was 179 extracted from the samples by acidification with phosphoric acid ( $H_3PO_4$ , 10%) using a custom, 180 automated CO<sub>2</sub> extractor and delivery system (AERICA by MARIANDA) using high grade  $N_2$  as a 181 182 carrier gas, connected online with a LiCor 6252 IR CO<sub>2</sub> analyzer. Measurements were calibrated using 183 seawater Certified Reference Materials (CRMs) from Dickson's lab. In order to check and correct for drift the CRM was run after every 4 samples. 184

185

### 186 Model of *Prochlorococcus* resolving key macromolecular pools

We formulated a mathematical model of the growth of a generalized axenic phytoplankton cell, 187 which resolves the macromolecular pools measured in the experiments – protein, RNA, DNA and 188 photosynthetic pigments, as well as N storage and dissolved organic carbon (Figure S3). The 189 model is formulated in units of N in each macromolecular pool, with the C in each pool determined by 190 191 the elemental ratios in each type of macromolecule (Geider and La Roche 2002). In this model, proteins 192 act as enzymes in the synthesis of RNA, DNA and Chl, whereas RNA acts as an enzyme for the synthesis of proteins. The total organic carbon in the media (TOC, the sum of cellular C and DOC) is 193 determined by the balance between photosynthesis and respiration. Since the cells are defined in terms 194 of N, and the TOC is defined in units of C, the concentration of DOC is an emergent property of the 195 196 model, and enables us to test under what conditions the model can reproduce high DOC production 197 rates. We fit the model to the experimental data presented below, using the Metropolis-Hastings 198 algorithm (Metropolis et al. 1953). A more detailed description of the model, including equations, can 199 be found in the supplementary information, and the model itself (written in FORTRAN) is available 200 on https://github.com/AWO-code/DalitDaniel\_Model.

#### 201 **Results**

# 202 Dynamics of cell numbers and macromolecular composition during growth and nitrogen 203 starvation

To determine to what extent the macromolecular composition of *Prochlorococcus* changes between 204 the different physiological states of a batch culture, we grew strain MIT9312 in laboratory batch 205 cultures where the N:P ratio of the media was set to 2, thus leading to cessation of growth due to N 206 207 starvation (Grossowicz et al. 2017). The bulk culture fluorescence, often used to monitor phytoplankton 208 growth in a non-invasive way, increased exponentially until day 10, after which the culture fluorescence declined rapidly, with no clearly observable stationary stage (Fig 1a). The decline of the 209 culture fluorescence coincided with the reduction of soluble NH4 to below detection threshold 210 211 (<10 uM), while PO<sub>4</sub> and DIC levels remained high throughout growth and decline (Fig 1c, d). Cell 212 counts by flow cytometry also showed an increase in the cell numbers until day 10, with a growth rate (µ) of 0.353 day<sup>-1</sup> (±0.0239) and doubling time of 1.968 days (±0.129). However, unlike the bulk 213 214 culture fluorescence that decline rapidly after this day, the cell numbers remained relatively stable for 215 an additional 3-4 days, before starting to decline. During this period, a clear population of cells with low chlorophyll autofluorescence (low-fl) emerged, identified by flow cytometry (Fig 1b, 67% low-fl 216 cells on day 13 and 88% on day 15, (Roth-Rosenberg et al. 2019)). These low-fl ("chlorotic") cells 217 218 were also stained much more weakly than the high-fl ones with the dye Sybr Green, which binds 219 nucleic acids (primarily dsDNA but also ssDNA and RNA, Fig S1b). At the same time, an increase in Sybr-Green staining was observed in the high-fl cells, potentially related to the arrest of the cells in G2 220 221 stage (Sup Fig. S1, (Zinser et al. 2009)). At the same time, the median forward scatter, a proxy for cell 222 size, increased for both the high-fl and low fl cells. (Fig S1c).

Based on the observed fluorescence curves (Fig 1a), we sampled cells during exponential stage (day6), early stationary stage (day 10), late stationary stage (after bulk culture fluorescence had started to

225 decline but while cell numbers were still stable, day 13) and culture decline (day 15) (Table S1). An additional estimate of the macromolecular pools from exponentially-growing cells was obtained from 226 227 the starter cultures used to start the experiment (treated here as day 0). The per-cell content of proteins and RNA were relatively stable during exponential phase (0-10 days), but was reduced during the 228 stationary and decline phases (days 13 and 15) (Fig 1e-f). The reduction in RNA/cell (a drop of 229 approximately 75% between days 10 and 13) was much larger than that of protein/cell (10-30%). The 230 231 total photosynthetic pigment concentration per cell started declining earlier than RNA or protein, with a drop of approximately 75% between days 6 and 10, as the cells moved from exponential growth to 232 233 early stationary stage (Fig 1g).



fluorescence curves of MIT9312 population shows a log-phase growth until day 10 than stationary and decline of the culture. (a) Flow cytometry scatter gram of specific measurements points. (c-d) Analysis of external inorganic nutrients shows that NH<sub>4</sub><sup>+</sup> drops below detection limit, while PO<sub>4</sub> and DIC are still available. Thus, the cultures enter stationary stage due to N starvation. Results are means and ranges of triplicate cultures. (e-g): Changes in the cell quotas of major macromolecules-DNA and RNA (e) proteins (f) and total pigments (g). While protein/cell is relatively stable, pigment and RNA pools drop as cells become starved and the culture declines. (h) Accumulation of large amounts of DOC. Triangles show measured Total Organic Carbon (TOC), squares show calculated C in cell biomass, based on flow cytometry counts and on per-cell C quotas. The error bars shown represents uncertainty due to the differences in estimates of per-cell C quotas between studies (Table 1), with the squares showing a value of 50 fg cell<sup>-1</sup>.

234

235

#### 236 Changes in photosynthetic pigment composition across the different growth stages

237 The reduction in chlorophyll autofluorescence per cell suggests changes to the photosynthetic machinery of the cells as they enter stationary stage and during culture decline. Indeed, as shown in 238 Fig 2a, in addition to a reduction in the total pigment quota per cell (Fig 1g), the ratio of some accessory 239 240 pigments to divinyl chlorophyll A (DVchlA) changed over time. Chlorophyll B and a chlorophyll-C 241 like pigment were reduced compared to DVchlA after day 10, whereas the ratio of zeaxanthin (and also α-carotene) to DVchlA increased after this day (Fig 2b). In addition, on day 10 and later, UPLC 242 243 several peaks were observed close to that of DVchlA that differed from it in their retention time. These 244 peaks were not due to overloading of the UPLC column (were not seen when higher amounts of pigments were injected), and were reproducibly observed in other cultures from multiple 245

246 *Prochlorococcus* strains. One of these peaks (marked with an arrow in Fig 2c), which was especially prominent, had an absorption spectrum similar to that of DVchlA (Fig 2d) but was not one of the known 247 degradation products (pheophytin-A eluted later, whereas phaeophorbide A, chlorophyllide A and 248 divinyl protochlorophillide A all eluted much earlier, consistent with the loss of the phytol moiety). 249 Assuming a similar molar absorption rate as DVchlA, this pigment (which may be a DVchlA', an 250 epimerization product of DVchlA, (Komatsu et al. 2016)) comprised as much as 21% of the total 251 252 DVchlA pigments on day 15, at the end of the decline stage. Importantly, pigments with the same elution time and absorbance were repeatedly observed in samples collected from the oligotrophic 253 254 Eastern Mediterranean Sea, as demonstrated in Fig S2.

255



**Figure 2. Pigments content and relative composition change during growth.** (a) Changes in the major per-cell pigment quotas. Divinyl chlorophyll A - DVchlA, chlorophyll B - chlB, a chlorophyll C-like pigment - chlC, zeaxanthin - Zea, and  $\alpha$ - carotene -  $\alpha$ -car. (b) The relative abundance of zea to DVchlA increases, while that of chlB to DVchlA decreases. (c) UPLC chromatograms at different times show the appearance of additional pigments during stationary phase. The pigment marked with an arrow has an absorption spectrum similar to that of DVchlA (panel d), but differs from it in its retention time.

256

257

#### 258 Reproducibly high rates of DOC exudation in *Prochlorococcus* MIT9312.

Part of the organic carbon fixed by phytoplankton is released from the cell due to exudation and cell 259 260 mortality, and this organic carbon provides sustenance for co-occurring heterotrophic bacteria. To assess how much organic C is released by Prochlorococcus, we measured the total organic C in our 261 262 cultures, and compared it to the amount of C expected to be in the particulate fraction based on the 263 number of cells and the carbon quota of *Prochlorococcus* from previous studies (10-130 µg/cell, Table 264 1). During exponential stage (day 6), the measured TOC was within the range expected based on cell 265 numbers and C quota, however, during the stationary and decline phases the amount of TOC was up 266 to 23-fold higher than could be explained by cell biomass, suggesting a large amount of organic C was released into the growth media (Fig 1g). To check if the high release of organic carbon is common to 267 different Prochlorococcus strain, we repeated this experiment using both strain MIT9312 and strain 268 269 NATL2A which belongs to the Low Light I clade (Biller et al. 2014). High extracellular concentrations 270 of organic C (up to 13-fold higher than the predicted cellular biomass) were observed again for

- 271 MIT9312, whereas strain NATL2A that grew under the same N starvation conditions showed TOC
- values much lower than MIT9312, within the expected range of cell quota (Fig 3).

273

# 274 Table 1: Prochlorococcus Carbon biomass estimated in different studies, expanded from

275 (Bertilsson et al. 2003)

Strain	fg C cell <sup>-1</sup>	Analytical method	Reference
Natural population, Atlantic Ocean	10-70	X ray microanalysis	(Grob et al. 2013)
PCC9511, EQPAC, SB, GP2, SARG, NATL1	17-34	X ray microanalysis	(Heldal et al. 2003)
PCC9511	17-38	POC	(Claustre et al. 2002)
MED4	17-40	POC	(Fu et al. 2007)
VOL7(MED4), VOL8(MIT9515), VOL29, VOL4 (MIT9312), VOL1(MIT9215), UH18301	18-55	POC	(Martiny et al. 2016)
MIT9301, MED4, NATL2A, MIT9313	30-33 (HL), 45 (LLI), 80 (LLIV)	Macromechanical mass sensor, assuming C is 50% of wet biomass	(Cermak et al. 2016)
MIT9301, MED4, MIT9313	44-51 (HL), ~180 (LLIV)	POC	(Becker et al. 2014)

MED4	46-61	POC	(Bertilsson et al. 2003)
CCMP1378(MED4)	49+/-9	POC	(Cailliau et al. 1996)
Generic (0.6 µm diameter cell)	53	Cell size and volume- carbon conversion (470 fg C/µm <sup>3)</sup>	(Campbell et al. 1994)
Natural population, Atlantic Ocean	56	Cell size and volume- carbon conversion (325 fg C/mm3)	(DuRand et al. 2001)
Generic (0.8 µm diameter cell)	59	Cell size and volume- carbon conversion (220 fg C/mm3)	(Li et al. 1992)
MIT9211, MIT9215, MIT9312, MED4, SS120, MIT9302, MIT9303, MIT9313	61-94	Primaryproduction,potential overestimation(does not account forrespirationandexudation)	(Moore 1997)
MIT9312	67-123	Macromolecular pools	This study
MED4, MIT9401, SS120	78+/-27	Dry weight, assuming C is 50% of biomass	(Shaw et al. 2003)
Generic (0.8 μm diameter cell)	124	Cell size and volume- carbon conversion (294 fg C/cell)	(Veldhuis and Kraay 1990)

276

277

278

279

#### 280



**Figure 3: Differences in exudation rates between two axenic** *Prochlorococcus* **strains, MIT9312** (**a**) **and NATL2A** (**b**). The two strains were grown under identical conditions. Triangles show measured Total Organic Carbon (TOC), squares show calculated C in cell biomass, based on flow cytometry counts and on per-cell C quotas. The range shown represents uncertainty due to the differences in estimates of per-cell C quotas between studies (Table 1), with the square markers showing a value of 50 fg cell<sup>-1</sup>. Panel a shows an independent experiment from that shown in Fig 1h, yet the high release of DOC is consistent.

281

#### Figure 3: Differences in exudation rates between *Prochlorococcus* strains.

283

# 284 Discussion

Quantitative measurements of cellular parameters, including macromolecular composition andelemental quotas, are fundamental to our understanding (and modeling) of microbial systems.

287 However, measurements of the per-cell C quota of *Prochlorococcus*, performed using different 288 techniques, range over an order of magnitude (Table 1, (Bertilsson et al. 2003)). In this study, we 289 directly measured the cell quotas of DNA, RNA and photosynthetic pigments at different physiological 290 stages of laboratory batch culture in Prochlorococcus MIT9312. With some assumptions, these 291 measurements can be used to constrain upper and lower bounds of the total cell quotas of C, N and P 292 (Supplementary text, Supplementary Table 1). Measuring these macromolecular pools requires less 293 biomass and can be performed at higher throughput compared to measurements of total particulate or dissolved C and N (Supplementary text). Assuming a lower bound on the C:N ratio of 6 (Bertilsson et 294 295 al. 2003, Martiny et al. 2013, Martiny et al. 2016), the cellular C quota we estimate (~67-123 fg cell<sup>-1</sup>, 296 Supplementary Table 1) is at the upper range of measurements from other studies (Table 1), and is 297 driven primarily by the high (and mostly stable) measurements of protein/cell. Like any other 298 experimental result, these measurements are sensitive to the experimental design and the methodology 299 used. For example, while the method we used for protein determination (BCA) is relatively insensitive 300 to the effects of salts and detergents (if these are incorporated into the standard curve), the quantitative 301 results can be affected by, for example, the protein standards used (e.g. Bovine Serum Albumin or 302 Immunoglobulin). Similarly, our estimates of DNA/cell are about one-third those expected from cell 303 number and the genome size of MIT9312 (1.7Mbp, (Kettler et al. 2007)). Experimental measurements 304 of DNA/cell that are not consistent with genome size have been observed also in other studies 305 (Zimmerman et al. 2014), and may be caused by differences in the DNA extraction efficiency across 306 organisms or culture conditions. Finally, the Prochlorococcus population is not homogenous, 307 containing at times two distinct sub-populations of cells, with different flow cytometry signatures (as 308 discussed in detail below, Fig S1, see also (Coe et al. 2016, Roth-Rosenberg et al. 2019)). We did not 309 measure the amount of the macromolecular pools in each sub-population separately, due to the 310 difficulty in obtaining sufficient biomass from non-fixed cells sorted by fluorescence-activated cell

sorting. With these uncertainties in mind, in the following sections we use the measurements of macromolecular pools at different growth stages, as well as the estimates of cellular C, N and P, to discuss the relationship between cell physiology, macromolecular pools and elemental budgets. Doing so, we note the need for an in-depth comparison of measurements between labs, growth stages and methods, perhaps along the lines of similar studies used to calibrate pigment measurements (e.g. (Claustre et al. 2004)).

317

# 318 Dynamic changes in the macromolecular composition of *Prochlorococcus* in batch culture – 319 changes in the average cell or sub-populations?

320 In our laboratory batch cultures, *Prochlorococcus* grow exponentially until they run out of their 321 nitrogen source (NH<sub>4</sub>), at which stage they stop growing, enter a short stationary stage and finally die. 322 During this time, the cultures lose their green color, the cell population became heterogeneous, and populations of low-fl cells emerge, forming the majority of the population as the cultures decline (Fig 323 324 1a, Fig S1a). The low-fl cells also stain weaker with the dye Sybr-green (Fig S1), which binds both 325 DNA and RNA (albeit with a higher affinity for DNA, (Martens-Habbena and Sass 2006)). We propose 326 that a significant part of the changes in the mean per-cell concentration of chlorophyll and RNA is 327 caused by the presence of these low-fl cells, rather than by a gradual change in the concentration of these pools per cell. In the model cyanobacteria Synechococcus elegantus PCC7942 and Synechocystis 328 329 PCC6803, the loss of culture chlorophyll (chlorosis) is part of a developmental program to generate 330 resting stages, that can survive long-term starvation (e.g. (Sauer et al. 2001, Klotz et al. 2016)). In 331 contrast, we have recently shown that *Prochlorococcus* cultures that become chlorotic cannot revive 332 when nutrients are added, unless co-cultured with a heterotrophic bacterium (Roth-Rosenberg et al. 333 2019). Nevertheless, some of the chlorotic cells are still active (photosynthesize and take up NH<sub>4</sub>,

(Roth-Rosenberg et al. 2019)). Thus, we interpret the changes observed in the macromolecular pools
of the MIT9312 cultures, and primarily those that can be attributed to the low-fl (chlorotic) population,
as physiological responses to N starvation that nevertheless are not sufficient to enable the cells to
survive long-term nutrient stress.

338 In parallel to the observed loss of culture fluorescence and the appearance (and dominance) of low-fl 339 cells, changes were also in the per-cell concentration and composition of the photosynthetic pigments 340 (Fig 1h, Fig 2). Specifically, the mean relative amount of zeaxanthin increases compared to DVchlA, 341 whereas the relative amount of chlorophyll B decreases (Fig 2b). Similar changes were observed in 342 nutrient-replete batch cultures of other *Prochlorococcus* (strains MED4 and SS120) in response to 343 increasing light intensities (Moore et al. 1995). One possible interpretation, which we deem less likely, 344 is that the changes in the pigment ratio as the cells enter stationary phase and start declining are due to 345 increased light levels as many of the cells become chlorotic and the cultures become clear. An 346 alternative interpretation is that the increase in relative abundance of zeaxanthin is a general stress 347 response, as these pigments may act as antioxidants in cyanobacteria (Zhu et al. 2010). Indeed, an increase in the zeaxanthin-DVchlA ratio was observed upon nitrogen starvation also in an additional 348 *Prochlorococcus* strain, SS120 (Steglich et al. 2001). The decrease in chlorophyll B can also be 349 interpreted independently from a potential change in light levels. The reduction in DVchlB could 350 351 simply be related to the deactivation of the photosystems, in agreement with previous studies that 352 showed a decrease in photosynthetic activity (e.g. Fv/Fm) and psbA protein levels upon nitrogen starvation (Steglich et al. 2001, Tolonen et al. 2006). Alternatively, the reduction in DVchlB may be 353 354 part of a cellular mechanism to reduce the use of N-containing pigments under N starvation, as chlorophylls contain nitrogen, whereas xanthophylls do not. This is consistent with other mechanisms 355 356 whereby *Prochlorococcus* conserve N, including changes in the transcriptional profiles (Tolonen et al.

2006), use of shorter transcripts (Read et al. 2017) and, potentially, "thrifty" N use in proteins expressed
in response to N starvation (Gilbert and Fagan 2011).

The emergence of the DVchlA-like pigment is currently unexplained, although this pigment may be 359 an epimerization product of DVchlA (DVchlA'). DVchlA' has been previously observed in several 360 361 Prochlororoccus strains (Komatsu et al. 2016), and in our study the DVchlA-like pigment is associated with culture decline. Generally speaking, harvesting cells from late exponential stage or early 362 363 stationary stage is a common practice when high biomass is required, yet as discussed above the 364 physiology of the cells can change during stationary phase in comparison with exponentially-growing 365 cells. Additionally, it is currently unclear to what extent processes observed in late stages of laboratory 366 batch cultures, such as nutrient starvation and the subsequent changes in cell physiology, occur in 367 nature. Nevertheless, the observation that a pigment with the same retention time and absorption spectrum as the DVchlA-like pigment observed at-sea suggests that at least some of the cells in nature 368 369 may be undergoing processes similar to those we observe in the laboratory batch cultures.

370 The high-fl cells also exhibit an increase in mean forward scatter, a proxy for cell size. This occurs 371 despite the overall reduction in RNA, pigments and (to a lesser extent) protein (Fig 1e, f), which are 372 the major N-containing macromolecular pools. Assuming the increase in forward scatter represents an increase in cell biomass, this suggests that the cells are accumulating primarily C-rich macromolecules 373 374 such as storage carbohydrates and lipids, leading to an increased C:N ratio. A similar response to 375 nitrogen starvation has been predicted by mathematical models of N-starved cells (Grossowicz et al. 2017) and observed in other phytoplankton species (Hu et al. 2008, Breuer et al. 2012, Liefer et al. 376 377 2019). If these processes occur in nature, they could impact the C:N ratio of DOM released from dead 378 Prochlorococcus cells (e.g. (Agusti and Sanchez 2002, Llabrés et al. 2011, Ribalet et al. 2015)) as well as POM exported by these cells (Richardson and Jackson 2007, Lomas and Moran 2012, Zhao et al. 379 380 2017).

381

# 382 The macromolecular composition of *Prochlorococcus* MIT9312 is more akin to eukaryotic 383 phytoplankton than to heterotrophic bacteria

Phytoplaknton and heterotorphic bacteria are inherently different in their life histories, and this likely 384 385 determines and constrains the relative allocation of energy and elements to different molecular functions. Phytoplankton need to allocate resources to the photosynthetic and carbon fixation 386 machineries and are likely less constrained by the availability of carbon (which can be derived from 387 388 photosynthesis) compared to other elements such as nitrogen. Eukaryotic phytoplankton are also often larger than bacteria, and thus are less constrained by the internal volume of the cell. Heterotrophic 389 bacteria, in contrast, do not need to allocate resources to photosynthesis but are potentially limited by 390 391 C and by cell size. As shown in Fig 4, the limited number of measurements of macromolecular pools 392 in marine microorganisms available in the literature supports the idea phytoplankton and bacteria 393 allocate resources differently.

Generally speaking, phytoplankton have a lower RNA/protein ratio than heterotrophic bacteria with 394 395 the same RNA/DNA ratio. Assuming that the ribosomes of phytoplankton and bacteria have similar maximal rates of protein production (chain elongation), this suggests that phytoplankton ribosomes are 396 working at a higher relative capacity. In phytoplankton, a significant amount of protein production 397 needs to be invested in maintaining the photosynthetic apparatus, for example in the replacement of 398 399 the core proteins (Zavřel et al. 2019). An alternative (but non-exclusive) explanation is that 400 heterotrophic bacteria (primarily copiotrophic ones) maintain more ribosomes than needed, using the "spare" production capacity to allow rapid response to changes in growth conditions. Such a strategy 401 402 has been demonstrated experimentally for E. coli (e.g. (Li et al. 2018). Notably, some bacteria, such as 403 Psychrobacter Mor119, Halomonas Hal146, Vibrio Vib2d and Ruegeria Oce241, are more similar in 404 the RNA/protein ration to phytoplankton (Fig 4). The reason for this similarity, which may represent a lower ability to allocate ribosome resources to rapid changes in gene expression, is unclear, and may 405 depend on the specific experimental conditions employed in their study (Zimmerman et al. 2014). 406 407 While *Prochlorococcus* is a bacterium (prokaryote), its RNA/protein and RNA/DNA ratios at different stages of the growth curve all fall within the range of those from eukaryotic phytoplankton. The single 408 409 measurement form Synechococcus is also closer to other phytoplankton (Fig 4). This could be the result 410 of both the need to invest in maintaining the photosystem and, potentially, that *Prochlorococcus* may have a lower ability compared to fast-growing heterotrophs to "ramp up" their translation in order to 411 412 rapidly respond to changes in environmental conditions.

413 Nevertheless, *Prochlorococcus* may still maintain some unused ribosome capacity. In our study, 414 protein/cell remained relatively stable, with a decline of no more than  $\sim 30\%$  as the culture declined. 415 RNA/cell, in contrast, dropped by more than 80% over the same period (Fig 1). The decline in 416 RNA/cell also started earlier. This suggests either that processes resulting in the loss of protein (e.g. degradation, exudation or excretion) were strongly reduced in stationary and decline phase cells, or 417 418 that the actual rate of protein production per ribosome increased in declining cultures compared to exponentially-growing ones. These two explanations - decrease in loss processes or increase in 419 420 ribosome efficiency - are not mutually exclusive, but evidence is lacking for either of them in most 421 organisms, including Prochlorococcus.

#### 422 Figure 4. RNA/DNA and RNA/protein compared to those of other phytoplankton and bacteria.



**Figure 4. RNA/DNA and RNA/protein compared to those of other phytoplankton and bacteria.** The ratios in phytoplankton (from (Finkel et al. 2016)), presented in cross markers , show a high diversity of RNA/DNA, while relatively low diversity of RNA/Proteins compositions. Heterotrophs bacteria however ( squares) are highly diverse in both RNA/Proteins and RNA/DNA composition (Churchward et al. 1982, Neidhardt et al. 1990, Zimmerman et al. 2014). *Prochlorococcus* MIT9312 at different growth phases (spheres of different sizes) are in fact similar to microalgae rather to bacterial macromolecule composition. The most phylogenetically related cyanobacteria, Synechococcus WH7803, marked in circle, shows similarity to MIT9312 in the RNA/DNA but not in the RNA/Protein ratios.

423

# 424 DOC accumulation and its potential mechanism

One of the surprising observations of this study was the high rate of accumulation of DOC during the
exponential growth of MIT9312. By the time the cultures reached stationary stage (day 13 in Fig 1h)

427 and day 11 in Fig 3a) the particulate organic carbon (i.e. cell biomass ) was only 3-7% of the total organic carbon (considering the two experiments shown in Fig 1h and 3h, and assuming a cell C quota 428 429 of 120 fg cell<sup>-1</sup>). This suggests that 80-85% of the organic carbon fixed by *Prochlorococcus* in two 430 separate experiments was released as dissolved organic carbon. Pervious experimental studies where DOC release was measured using  $^{14}$ C as a tracer suggested that 2-24% of the primary productivity is 431 432 released (Bertilsson et al. 2005, Lopez-Sandoval et al. 2013). However, indirect assessments suggest 433 that a somewhat higher fraction of the fixed carbon is lost from the cells. First, a comparison of carbon 434 uptake through photosynthesis and growth rate in Prochlorococcus MED4 over a diel cycle suggested 435 that ~30% of the fixed carbon is released or respired (Zinser et al. 2009). Second, the theoretical growth 436 rate of MIT9312 based on the expected yield of photosynthesis (taking into account published values 437 for illumination, cross section, efficiency and chlorophyll/cell, (Moore and Chisholm 1999)) is almost 438 two-fold higher than measured (expected  $\mu$ =0.64 day<sup>-1</sup>, compared to 0.35±0.02 day<sup>-1</sup> measured during exponential stage, see Supplementary Information for more details). Third, we used a mathematical 439 model of a generalized phytoplankton, resolving the macromolecular pools measured in this 440 441 experiment, to test whether the exudation rates we observed are feasible. The model was able to 442 reproduce quite well some of the experimental results, including the accumulation of DOC (observed 443 as an increase in TOC, Figs S3-S5, Supplementary Table S2). This model reproduced well the cell 444 density, the uptake of inorganic N and the reduction after stationary stage in the cellular quotas of RNA and chl-a, although the timing in the decline of the protein quota per cell was predicted to be earlier 445 446 than observed experimentally. Importantly, while the values of C fixation per unit chl-a in most of 447 the model simulations were within the range of the maximum photosynthesis measured for 448 *Prochlorococcus* (up to 10-20mol C[g divinyl Chl-a]<sup>-1</sup>day<sup>-1</sup>, (Moore and Chisholm 1999, Bruyant et 449 al. 2005, Felcmanová et al. 2017)), this value had to vary over the dynamics of batch culture for the 450 model to reproduce well the accumulation of TOC, increasing transiently as the cells entered the

451 decline phase (Fig S5). This phenomenon – an increase in photosynthetic efficiency in cells entering chlorosis (potentially caused by a reduction in shelf-shading, (Felcmanová et al. 2017)) - has not, to 452 453 the best of our knowledge, been described previously. An alternative explanation is that the respiration 454 rates decrease in relation to growth rate as the cells enter chlorosis (Fang et al. 2019). The current 455 model formulation cannot distinguish between these two possibilities, and thus experimental measurements of photosynthesis and respiration across batch culture are required to test these two 456 alternative model predictions. Taken together, these considerations show that while the rates of DOC 457 release recorded here are 2-3 fold higher than those demonstrated experimentally (e.g. using <sup>14</sup>C release 458 459 assays), they are within the range of growth rates possible based on theoretical considerations.

460

The differences between our results, based on accumulation of DOC over time, and estimates of 461 exudation based on <sup>14</sup>C partitioning between the particulate and dissolved phases, may also be due to 462 loss of volatile substances during the acidification and subsequent venting of inorganic <sup>14</sup>C during 463 464 radioactive incorporation experiments. Indeed, Prochloroccoccus produce several volatile substances (e.g. organic acids, organohalogens, methane and isoprene, (Shaw et al. 2003, Bertilsson et al. 2005, 465 Hughes et al. 2011, Bižić-Ionescu et al. 2019)), although to what extent these substances would be lost 466 from the media during the protocol for measuring <sup>14</sup>C incorporation is unclear. Thus, our results 467 showing high release of DOC are consistent with theoretical calculations and (some) previous studies, 468 but also highlight the need for additional studies, specifically addressing DOC release using different 469 analytical methods. 470

The DOM released by *Prochlorococcus* can include, in addition to C, also other elements such as N and P. While we did not measure this experimentally, a rough estimate of the loss processes N and P can be obtained by comparing the total amount of N and P in the original media and the estimated

474 amount of N and P in the cells at their maximum density. Since Prochlorococcus do not fix dinitrogen or perform denitrification, the N "missing" from the cell biomass when reach stationary stage under 475 476 N-starved conditions should represents organic N that were lost from the cells in a format that is not 477 available for re-uptake (i.e. refractory to *Prochlorococcus*). At the peak of the *Prochlorococcus* growth in our experiment, on day 10, the cellular N accounted for 79% of the total N in the media. Given that, 478 on that day, NH<sub>4</sub> in the media was depleted, this suggests that ~20% of the N had already been released 479 480 from the cells in a form that is not available for re-utilization. Thus, while C rich, the DOM released by *Prochlorococcus* MIT9312 contains significant amounts of other elements, likely (at least in part) 481 482 in the form of proteins, amino acids, DNA, RNA and nucleotides. This is consistent with studies in 483 Synechococcus (Christie-Oleza et al. 2015, Christie-Oleza et al. 2017, Zhao et al. 2017) and with the 484 effect of *Prochlorococcus*-derived DOM on gene expression in natural communities (Sharma et al. 485 2014). With regard to P, a similar calculation shows that only ~3.2% of the total P in the media was 486 found in cellular biomass, while the PO<sub>4</sub> concentration in the media dropped from 50µM to 34-36µM, a reduction of  $\sim 30\%$ . Thus, either we have significantly underestimated total cellular P (i.e. there is an 487 additional, unmeasured pool besides RNA, DNA and phospholipids), or a significant amount of P has 488 489 been converted from PO<sub>4</sub> to extracellular DOP.

490 While our understanding of biogeochemistry is framed in terms of elements, cells actually release the same macromolecules from which they are built, e.g. the proteins, DNA and RNA measured in this 491 492 study. Previous studies have suggested that *Prochlorococcus* release, in addition to the volatile 493 compounds described above, also TEP (Iuculano et al. 2017), and that at least part of the produced 494 DOM (the hydrophobic fraction that can be analyzed following solid-phase extraction) differs from 495 strain to strain, and even between replicate cultures (Becker et al. 2014). Prochlorococcus have also 496 been shown to release vesicles containing proteins and nucleic acids (Biller et al. 2014). Furthermore, 497 DOC loss from cultures can be due not only to passive leakage from the cells (exudation) or active

release (excretion) but also to lysis of dead cells (reviewed by (Thornton 2014)), with the latter process
likely releasing most or all of the cellular macromolecular content. Several studies have indicated that
up to ~20% of the cells in exponentially-growing *Prochlorococcus* cultures may be dead (Agusti and
Sanchez 2002, Hughes et al. 2011). Therefore, it is likely that all of these processes (exudation,
excretion, vesicle production and mortality) contribute to DOM formation.

503 Several studies have suggested that high rates of DOC release occur in oligotrophic regions of the 504 oceans, and that DOC, rather than POC, contributes significantly to carbon export from the photic zone 505 in these areas (Guyennon et al. 2015, Roshan and DeVries 2017). It is tempting to speculate that 506 *Prochlorococcus* may be at least partially responsible for this DOC accumulation, based on the high rates of DOC production observed here by Prochlorococcus MIT9312, a member of the HL-II clade 507 508 that numerically dominates large areas of the oligotrophic ocean (Bouman et al. 2006, Johnson et al. 509 2006). In support of this speculation, DOM released by *Prochlorococcus* in the surface waters of the 510 Pacific Ocean (which was dominated by the HL-II clade) has been suggested to provide as much as 511 75% of the daily organic carbon produced (Ribalet et al. 2015), although it is unclear to what extent 512 mortality compared to exudation feeds this organic pulse. It has also been suggested that DOM derived 513 from *Prochlorococcus* contributes a significant amount to deep sea DOM (Zhao et al. 2017). We note, however, that DOC production rates may differ based on the physiological states of the cells and the 514 515 surrounding nutrient conditions. For example, similar to our study, Bertilsson and co-workers also 516 noted differences between two strains of *Prochlorococcus*, MIT9312 and MED4, with the former 517 releasing approximately twice as much DOC (Bertilsson et al. 2005), and Becker and co-workers 518 documented differences in the composition of released DOM (Becker et al. 2014). Future work is 519 required to determine the to what extent DOM production changes between strains and across culture 520 conditions, what are the mechanisms of this process, and to what extent assessments of DOM release 521 under lab conditions can be extended to the ocean.

522

#### 523 Acknowledgements

- 524 We thank Hila Elifantz, Ilana Berman-Frank, Eyal Geisler and Edo Bar-Zeev for the TOC analyses
- and Jack Silberman for the DIC analysis. This study was supported by the Gordon & Betty Moore
- 526 Foundation (grant number GBMF #3778, to MJF), by the Human Frontiers Science Program (grant
- 527 number grant RGP0020/2016, to DS) and by the United States-Israel Binational Science Foundation
- 528 (Grant number 2010183 to DS and MJF and grant number 2016532 to DS).

529

# 530 Author contributions

531 DR, DA, AWO, MJF and DS conceived the study and designed the experiments. DR and DA

performed the experiments and analyzed the samples. AWO performed the modelling. DR and DS

533 wrote the manuscript with contributions from all authors.

534

# 535 **Competing interests**

536 The authors declare no competing interests

537

# 538 Materials and Correspondence

539 Please send requests for materials or other correspondence to Daniel Sher, dsher@univ.haifa.ac.il

540

541

5	л	2
J	4	۷.

### 543

### 544 **References**

#### 545

- Agusti, S. and M. C. Sanchez (2002). "Cell viability in natural phytoplankton communities quantified by a
- 547 membrane permeability probe." <u>Limnology and Oceanography</u> **47**(3): 818-828.
- 548 Aharonovich, D. and D. Sher (2016). "Transcriptional response of Prochlorococcus to co-culture with a marine
- Alteromonas: differences between strains and the involvement of putative infochemicals." <u>ISME J</u> 10(12):
   2892-2906.
- 551 Becker, J. W., P. M. Berube, C. L. Follett, J. B. Waterbury, S. W. Chisholm, E. F. DeLong and D. J. Repeta
- (2014). "Closely related phytoplankton species produce similar suites of dissolved organic matter." <u>Frontiers</u>
   in Microbiology 5.
- 554 Bertilsson, S., O. Berglund, D. M. Karl and S. W. Chisholm (2003). "Elemental composition of marine
- 555 Prochlorococcus and Synechococcus: Implications for the ecological stoichiometry of the sea." <u>Limnology and</u> 556 Oceanography **48**(5): 1721-1731.
- 557 Bertilsson, S., O. Berglund, M. J. Pullin and S. W. Chisholm (2005). "Release of dissolved organic matter by 558 Prochlorococcus." Vie Et Milieu-Life and Environment **55**(3-4): 225-231.
- 559 Berube, P. M., A. Rasmussen, R. Braakman, R. Stepanauskas and S. W. Chisholm (2019). "Emergence of trait 560 variability through the lens of nitrogen assimilation in Prochlorococcus." eLife **8**: e41043.
- 561 Biller, S. J., P. M. Berube, J. W. Berta-Thompson, L. Kelly, S. E. Roggensack, L. Awad, K. H. Roache-Johnson, H.
- 562 Ding, S. J. Giovannoni, G. Rocap, L. R. Moore and S. W. Chisholm (2014). "Genomes of diverse isolates of the 563 marine cyanobacterium Prochlorococcus." <u>Scientific Data</u> **1**.
- 564 Biller, S. J., P. M. Berube, D. Lindell and S. W. Chisholm (2014). "Prochlorococcus: the structure and function 565 of collective diversity." <u>Nature Reviews Microbiology</u> **13**: 13.
- Biller, S. J., F. Schubotz, S. E. Roggensack, A. W. Thompson, R. E. Summons and S. W. Chisholm (2014).
- 567 "Bacterial vesicles in marine ecosystems." <u>Science</u> **343**(6167): 183-186.
- 568 Bižić-Ionescu, M., T. Klintzsch, D. Ionescu, M. Y. Hindiyeh, M. Günthel, A. M. Muro-Pastor, W. Eckert, F.
- 569 Keppler and H.-P. Grossart (2019). "Widespread methane formation by <em>Cyanobacteria</em> in aquatic 570 and terrestrial ecosystems." bioRxiv: 398958.
- 571 Bouman, H. A., O. Ulloa, D. J. Scanlan, K. Zwirglmaier, W. K. Li, T. Platt, V. Stuart, R. Barlow, O. Leth, L.
- 572 Clementson, V. Lutz, M. Fukasawa, S. Watanabe and S. Sathyendranath (2006). "Oceanographic basis of the
- 573 global surface distribution of Prochlorococcus ecotypes." <u>Science</u> **312**(5775): 918-921.
- 574 Breuer, G., P. P. Lamers, D. E. Martens, R. B. Draaisma and R. H. Wijffels (2012). "The impact of nitrogen
- 575 starvation on the dynamics of triacylglycerol accumulation in nine microalgae strains." <u>Bioresource</u>
- 576 <u>Technology</u> **124**: 217-226.
- 577 Bruyant, F., M. Babin, B. Genty, O. Prasil, M. J. Behrenfeld, H. Claustre, A. Bricaud, L. Garczarek, J.
- 578 Holtzendorff, M. Koblizek, H. Dousova and F. Partensky (2005). "Diel variations in the photosynthetic
- 579 parameters of Prochlorococcus strain PCC 9511: Combined effects of light and cell cycle." Limnology and
- 580 <u>Oceanography</u> **50**(3): 850-863.
- 581 Cailliau, C., H. Claustre, F. Vidussi, D. Marie and D. Vaulot (1996). "Carbon biomass, and gross growth rates as
- estimated from <sup>14</sup> C pigment labelling, during photoacclimation in Prochlorococcus CCMP
- 583 1378." <u>Marine Ecology Progress Series</u> **145**(1/3): 209-221.

- Campbell, L., H. A. Nolla and D. Vaulot (1994). "The importance of Prochlorococcus to community structure in
   the central North Pacific Ocean." <u>Limnology and Oceanography</u> **39**(4): 954-961.
- 586 Cermak, N., J. W. Becker, S. M. Knudsen, S. W. Chisholm, S. R. Manalis and M. F. Polz (2016). "Direct single-
- cell biomass estimates for marine bacteria via Archimedes' principle." <u>The Isme Journal</u> **11**: 825.
- 588 Christie-Oleza, J. A., D. J. Scanlan and J. Armengaud (2015). ""You produce while I clean up", a strategy
- 589 revealed by exoproteomics during Synechococcus–Roseobacter interactions." <u>PROTEOMICS</u>: n/a-n/a.
- 590 Christie-Oleza, J. A., D. Sousoni, M. Lloyd, J. Armengaud and D. J. Scanlan (2017). "Nutrient recycling
- 591 facilitates long-term stability of marine microbial phototroph–heterotroph interactions." <u>Nature</u>
- 592 <u>Microbiology</u> **2**: 17100.
- 593 Churchward, G., H. Bremer and R. Young (1982). "Macromolecular composition of bacteria." <u>Journal of</u>
   594 <u>Theoretical Biology</u> **94**(3): 651-670.
- 595 Claustre, H., A. Bricaud, M. Babin, F. Bruyant, L. Guillou, F. Le Gall, D. Marie and F. Partensky (2002). "Diel 596 variations in Prochlorococcus optical properties." <u>Limnology and Oceanography</u> **47**(6): 1637-1647.
- 597 Claustre, H., S. B. Hooker, L. Van Heukelem, J. F. Berthon, R. Barlow, J. Ras, H. Sessions, C. Targa, C. S.
- 598 Thomas, D. van der Linde and J. C. Marty (2004). "An intercomparison of HPLC phytoplankton pigment
- 599 methods using in situ samples: application to remote sensing and database activities." Marine Chemistry
- 600 **85**(1-2): 41-61.
- 601 Coe, A., J. Ghizzoni, K. LeGault, S. Biller, S. E. Roggensack and S. W. Chisholm (2016). "Survival of
- 602 Prochlorococcus in extended darkness." <u>Limnology and Oceanography</u> **61**(4): 1375-1388.
- Dickson, A. G., C. L. Sabine and J. R. e. Christian (2007). Guide to best practices for ocean CO2 measurement.
   Sidney, British Columbia, North Pacific Marine Science Organization: 191.
- 605 Dubinsky, Z. and I. Berman-Frank (2001). "Uncoupling primary production from population growth in
- 606 photosynthesizing organisms in aquatic ecosystems." <u>Aquatic Sciences</u> **63**(1): 4-17.
- 607 DuRand, M. D., R. J. Olson and S. W. Chisholm (2001). "Phytoplankton population dynamics at the Bermuda
- Atlantic Time-series station in the Sargasso Sea." <u>Deep Sea Research Part II: Topical Studies in Oceanography</u>
   **48**(8–9): 1983-2003.
- 610 Elser, J. J., K. Acharya, M. Kyle, J. Cotner, W. Makino, T. Markow, T. Watts, S. Hobbie, W. Fagan, J. Schade, J.
- 611 Hood and R. W. Sterner (2003). "Growth rate-stoichiometry couplings in diverse biota." <u>Ecology Letters</u>
- 612 **6**(10): 936-943.
- Fang, X., Y. Liu, Y. Zhao, Y. Chen, R. Liu, Q.-L. Qin, G. Li, Y.-Z. Zhang, W. Chan, W. R. Hess and Q. Zeng (2019).
- 614 "Transcriptomic responses of the marine cyanobacterium Prochlorococcus to viral lysis products."
- 615 Environmental Microbiology **21**(6): 2015-2028.
- 616 Felcmanová, K., M. Lukeš, E. Kotabová, E. Lawrenz, K. H. Halsey and O. Prášil (2017). "Carbon use efficiencies
- and allocation strategies in Prochlorococcus marinus strain PCC 9511 during nitrogen-limited growth."
- 618 <u>Photosynthesis Research</u> **134**(1): 71-82.
- Finkel, Z. V., M. J. Follows, J. D. Liefer, C. M. Brown, I. Benner and A. J. Irwin (2016). "Phylogenetic Diversity in
  the Macromolecular Composition of Microalgae." <u>PLOS ONE</u> **11**(5): e0155977.
- Flombaum, P., J. L. Gallegos, R. A. Gordillo, J. Rincón, L. L. Zabala, N. Jiao, D. M. Karl, W. K. W. Li, M. W.
- 622 Lomas, D. Veneziano, C. S. Vera, J. A. Vrugt and A. C. Martiny (2013). "Present and future global distributions
- 623 of the marine Cyanobacteria <em>Prochlorococcus</em> and <em>Synechococcus</em>." <u>Proceedings of</u>
- 624 the National Academy of Sciences **110**(24): 9824-9829.
- 625 Fu, F.-X., M. E. Warner, Y. Zhang, Y. Feng and D. A. Hutchins (2007). "EFFECTS OF INCREASED TEMPERATURE
- AND CO2 ON PHOTOSYNTHESIS, GROWTH, AND ELEMENTAL RATIOS IN MARINE SYNECHOCOCCUS AND
   PROCHLOROCOCCUS (CYANOBACTERIA)1." Journal of Phycology 43(3): 485-496.
- 627 PROCHLOROCOCCUS (CYANOBACTERIA)1. Journal of Phycology 43(3): 485-496.
- 628 Garcia, N. S., J. A. Bonachela and A. C. Martiny (2016). "Interactions between growth-dependent changes in 629 cell size, nutrient supply and cellular elemental stoichiometry of marine Synechococcus." The Isme Journal
- 630 **10**: 2715.
- 631 Geider, R. J. and J. La Roche (2002). "Redfield revisited: variability of C : N : P in marine microalgae and its
- 632 biochemical basis." <u>European Journal of Phycology</u> **37**(1): 1-17.

- 633 Gilbert, J. D. J. and W. F. Fagan (2011). "Contrasting mechanisms of proteomic nitrogen thrift in
- 634 Prochlorococcus." <u>Molecular Ecology</u> **20**(1): 92-104.
- Goebel, N. L., C. A. Edwards, J. P. Zehr and M. J. Follows (2010). "An emergent community ecosystem model
- applied to the California Current System." Journal of Marine Systems 83(3): 221-241.
- Grob, C., M. Ostrowski, R. J. Holland, M. Heldal, S. Norland, E. S. Erichsen, C. Blindauer, A. P. Martin, M. V.
- 638 Zubkov and D. J. Scanlan (2013). "Elemental composition of natural populations of key microbial groups in
- 639 Atlantic waters." <u>Environmental Microbiology</u> **15**(11): 3054-3064.
- 640 Grossowicz, M., D. Roth-Rosenberg, D. Aharonovich, J. Silverman, M. J. Follows and D. Sher (2017).
- 641 "Prochlorococcus in the lab and in silico: The importance of representing exudation." <u>Limnology and</u>
  642 Oceanography 62(2): 818-835.
- 643 Gruber, N. and C. A. Deutsch (2014). "Redfield's evolving legacy." <u>Nature Geoscience</u> **7**: 853.
- 644 Guyennon, A., M. Baklouti, F. Diaz, J. Palmieri, J. Beuvier, C. Lebaupin-Brossier, T. Arsouze, K. Béranger, J. C.
- Dutay and T. Moutin (2015). "New insights into the organic carbon export in the Mediterranean Sea from 3-D
   modeling." <u>Biogeosciences</u> 12(23): 7025-7046.
- Hansell, D. A., C. A. Carlson, D. J. Repeta and R. Schlitzer (2009). "Dissolved Organic Matter in the Ocean a
  Controversy Stimulates New Insights." <u>Oceanography</u> 22(4): 202-211.
- Heldal, M., D. J. Scanlan, S. Norland, F. Thingstad and N. H. Mann (2003). "Elemental composition of single
- cells of various strains of marine Prochlorococcus and Synechococcus using X-ray microanalysis." <u>Limnology</u>
   <u>and Oceanography</u> 48(5): 1732-1743.
- Hu, Q., M. Sommerfeld, E. Jarvis, M. Ghirardi, M. Posewitz, M. Seibert and A. Darzins (2008). "Microalgal
- triacylglycerols as feedstocks for biofuel production: perspectives and advances." <u>The Plant Journal</u> 54(4):
  621-639.
- Hughes, C., D. J. Franklin and G. Malin (2011). "Iodomethane production by two important marine
- cyanobacteria: Prochlorococcus marinus (CCMP 2389) and Synechococcus sp. (CCMP 2370)." <u>Marine</u>
   <u>Chemistry</u> 125(1–4): 19-25.
- Iuculano, F., I. P. Mazuecos, I. Reche and S. Agustí (2017). "Prochlorococcus as a Possible Source for
   Transparent Exopolymer Particles (TEP)." <u>Frontiers in Microbiology</u> 8(709).
- Johnson, Z. I., E. R. Zinser, A. Coe, N. P. McNulty, E. M. Woodward and S. W. Chisholm (2006). "Niche
- 661 partitioning among Prochlorococcus ecotypes along ocean-scale environmental gradients." <u>Science</u>
- 662 **311**(5768): 1737-1740.
- Kettler, G. C., A. C. Martiny, K. Huang, J. Zucker, M. L. Coleman, S. Rodrigue, F. Chen, A. Lapidus, S. Ferriera, J.
  Johnson, C. Steglich, G. M. Church, P. Richardson and S. W. Chisholm (2007). "Patterns and implications of
  gene gain and loss in the evolution of Prochlorococcus." PLoS Genet **3**(12): e231.
- 666 Klotz, A., J. Georg, L. Bučinská, S. Watanabe, V. Reimann, W. Januszewski, R. Sobotka, D. Jendrossek,
- 667 Wolfgang R. Hess and K. Forchhammer (2016). "Awakening of a Dormant Cyanobacterium from Nitrogen 668 Chlorosis Reveals a Genetically Determined Program." Current Biology **26**(21): 2862-2872.
- 669 Komatsu, H., K. Wada, T. Kanjoh, H. Miyashita, M. Sato, M. Kawachi and M. Kobayashi (2016). "Unique
- 670 chlorophylls in picoplankton Prochlorococcus sp. "Physicochemical properties of divinyl chlorophylls, and the
- discovery of monovinyl chlorophyll b as well as divinyl chlorophyll b in the species Prochlorococcus NIES-
- 672 2086"." Photosynth Res 130(1-3): 445-467.
- Li, S. H.-J., Z. Li, J. O. Park, C. G. King, J. D. Rabinowitz, N. S. Wingreen and Z. Gitai (2018). "Escherichia coli
- translation strategies differ across carbon, nitrogen and phosphorus limitation conditions." <u>Nature</u>
   <u>microbiology</u> **3**(8): 939-947.
- Li, W. K. W., P. M. Dickie, B. D. Irwin and A. M. Wood (1992). "Biomass of bacteria, cyanobacteria,
- 677 prochlorophytes and photosynthetic eukaryotes in the Sargasso Sea." <u>Deep Sea Research Part A.</u>
- 678 <u>Oceanographic Research Papers</u> **39**(3): 501-519.
- Liefer, J. D., A. Garg, M. H. Fyfe, A. J. Irwin, I. Benner, C. M. Brown, M. J. Follows, A. W. Omta and Z. V. Finkel
- 680 (2019). "The Macromolecular Basis of Phytoplankton C:N:P Under Nitrogen Starvation." <u>Frontiers in</u>
- 681 <u>Microbiology</u> **10**(763).

- 682 Llabrés, M., S. Agustí and G. J. Herndl (2011). "DIEL IN SITU PICOPHYTOPLANKTON CELL DEATH CYCLES
- 683 COUPLED WITH CELL DIVISION1." Journal of Phycology **47**(6): 1247-1257.
- Lomas, M. W. and S. B. Moran (2012). "Evidence for aggregation and export of cyanobacteria and nanoeukaryotes from the Sargasso Sea euphotic zone." Biogeosciences **7**(5): 7173-7206.
- 686 Lopez-Sandoval, D. C., T. Rodriguez-Ramos, P. Cermeno and E. Maranon (2013). "Exudation of organic carbon
- 687 by marine phytoplankton: dependence on taxon and cell size." Marine Ecology Progress Series **477**: 53-60.
- 688 Lourenço, S. O., E. Barbarino, U. M. L. Marquez and E. Aidar (1998). "DISTRIBUTION OF INTRACELLULAR
- 689 NITROGEN IN MARINE MICROALGAE: BASIS FOR THE CALCULATION OF SPECIFIC NITROGEN-TO-PROTEIN 690 CONVERSION FACTORS." Journal of Phycology **34**(5): 798-811.
- 691 Marmen, S., L. Blank, A. Al-Ashhab, A. Malik, L. Ganzert, M. Lalzar, H.-P. Grossart and D. Sher (2018). "The
- 692 role of land use types and water chemical properties in structuring the microbiome of a connected lakes693 system." <u>bioRxiv</u>.
- 694 Martens-Habbena, W. and H. Sass (2006). "Sensitive determination of microbial growth by nucleic acid 695 staining in aqueous suspension." Applied and environmental microbiology **72**(1): 87-95.
- 696 Martiny, A. C., M. L. Coleman and S. W. Chisholm (2006). "Phosphate acquisition genes in Prochlorococcus 697 ecotypes: evidence for genome-wide adaptation." <u>Proc Natl Acad Sci U S A</u> **103**(33): 12552-12557.
- Martiny, A. C., L. Ma, C. Mouginot, J. W. Chandler and E. R. Zinser (2016). "Interactions between Thermal
- Acclimation, Growth Rate, and Phylogeny Influence Prochlorococcus Elemental Stoichiometry." <u>PLOS ONE</u>
   **11**(12): e0168291.
- 701 Martiny, A. C., C. T. A. Pham, F. W. Primeau, J. A. Vrugt, J. K. Moore, S. A. Levin and M. W. Lomas (2013).
- "Strong latitudinal patterns in the elemental ratios of marine plankton and organic matter." <u>Nature</u>
   Geoscience 6: 279.
- Massana, R., A. E. Murray, C. M. Preston and E. F. DeLong (1997). "Vertical distribution and phylogenetic
- characterization of marine planktonic Archaea in the Santa Barbara Channel." <u>Applied and environmental</u>
   <u>microbiology</u> 63(1): 50-56.
- 707 McDonagh, B., M. A. Domínguez-Martín, G. Gómez-Baena, A. López-Lozano, J. Diez, J. A. Bárcena and J. M.
- García Fernández (2012). "Nitrogen starvation induces extensive changes in the redox proteome of Brochlerococcus on strain \$\$120 " Environmental Microhiology Peperts **4**(2): 257,267
- 709 Prochlorococcus sp. strain SS120." <u>Environmental Microbiology Reports</u> **4**(2): 257-267.
- 710 Metropolis, N., A. W. Rosenbluth, M. N. Rosenbluth, A. H. Teller and E. Teller (1953). "Equation of State
- 711 Calculations by Fast Computing Machines." <u>The Journal of Chemical Physics</u> **21**(6): 1087-1092.
- 712 Moore, L. R. (1997). <u>Physiological ecology of Prochlorococcus : a comparison of isolates from diverse</u>
- 713 <u>oceanographic regimes</u>. PhD, Massachusetts Institute of Technology.
- Moore, L. R. and S. W. Chisholm (1999). "Photophysiology of the marine cyanobacterium Prochlorococcus:
   Ecotypic differences among cultured isolates." <u>Limnology and Oceanography</u> 44(3): 628-638.
- Moore, L. R., A. Coe, E. R. Zinser, M. A. Saito, M. B. Sullivan, D. Lindell, K. Frois-Moniz, J. Waterbury and S. W.
- 717 Chisholm (2007). "Culturing the marine cyanobacterium Prochlorococcus." Limnology and Oceanography-
- 718 Methods **5**: 353-362.
- 719 Moore, L. R., R. Goericke and S. W. Chisholm (1995). "Comparative physiology of Synechococcus and
- 720 Prochlorococcus: influence of light and temperature on growth, pigments, fluorescence and absorptive
- 721 properties." <u>Marine Ecology Progress Series</u> **116**(1/3): 259-275.
- Moore, L. R., A. F. Post, G. Rocap and S. W. Chisholm (2002). "Utilization of different nitrogen sources by the
- marine cyanobacteria Prochlorococcus and Synechococcus." <u>Limnology and Oceanography</u> **47**(4): 989-996.
- 724 Neidhardt, F. C., J. L. Ingraham and M. Schaechter (1990). <u>Physiology of the Bacterial Cell: a Molecular</u>
- 725 <u>Approach</u>, Sinauer Associates, Sunderland, Mass.
- 726 Omta, A. W., J. Bruggeman, B. Kooijman and H. Dijkstra (2009). "The organic carbon pump in the Atlantic."
- 727 Journal of Sea Research **62**(2): 179-187.
- Omta, A. W., D. Talmy, D. Sher, Z. V. Finkel, A. J. Irwin and M. J. Follows (2017). "Extracting phytoplankton
- physiological traits from batch and chemostat culture data." <u>Limnology and Oceanography: Methods</u> **15**(5):
- 730 453-466.

- Partensky, F. and L. Garczarek (2010). "Prochlorococcus: Advantages and Limits of Minimalism." <u>Annual</u>
   <u>Review of Marine Science</u> 2: 305-331.
- 733 Read, R. W., P. M. Berube, S. J. Biller, I. Neveux, A. Cubillos-Ruiz, S. W. Chisholm and J. J. Grzymski (2017).
- "Nitrogen cost minimization is promoted by structural changes in the transcriptome of N-deprived
   Prochlorococcus cells." The ISME journal **11**(10): 2267-2278.
- Redfield, A. C. (1934). On the Proportions of Organic Derivatives in Sea Water and Their Relation to the
- 737 Composition of Plankton. James Johnstone Memorial Volume, University Press of Liverpool: 176-192.
- Ribalet, F., J. Swalwell, S. Clayton, V. Jiménez, S. Sudek, Y. Lin, Z. I. Johnson, A. Z. Worden and E. V. Armbrust
- 739 (2015). "Light-driven synchrony of Prochlorococcus growth and mortality in the subtropical Pacific gyre."
- 740 <u>Proceedings of the National Academy of Sciences</u> **112**(26): 8008-8012.
- Richardson, T. L. and G. A. Jackson (2007). "Small phytoplankton and carbon export from the surface ocean."
  Science **315**(5813): 838-840.
- 743 Roshan, S. and T. DeVries (2017). "Efficient dissolved organic carbon production and export in the
- 744 oligotrophic ocean." <u>Nature Communications</u> **8**(1): 2036.
- Roth-Rosenberg, D., D. Aharonovich, T. Luzzatto-Knaan, A. Vogts, L. Zoccarato, F. Eigemann, N. Nago, H. P.
- Grossart, M. Voss and D. Sher (2019). "Prochlorococcus rely on microbial interactions rather than on
- 747 chlorotic resting stages to survive long-term stress." <u>BioRXiv</u> MS ID#: BIORXIV/2019/657627.
- Saito, M. A., M. R. McIlvin, D. M. Moran, T. J. Goepfert, G. R. DiTullio, A. F. Post and C. H. Lamborg (2014).
- "Multiple nutrient stresses at intersecting Pacific Ocean biomes detected by protein biomarkers." <u>Science</u>
   345(6201): 1173-1177.
- 751 Sauer, J., U. Schreiber, R. Schmid, U. Völker and K. Forchhammer (2001). "Nitrogen Starvation-Induced
- 752 Chlorosis in Synechococcus PCC 7942. Low-Level Photosynthesis As a Mechanism of Long-Term Survival."
   753 Plant Physiology 126(1): 233-243.
- 754 Sharma, A. K., J. W. Becker, E. A. Ottesen, J. A. Bryant, S. Duhamel, D. M. Karl, O. X. Cordero, D. J. Repeta and
- E. F. DeLong (2014). "Distinct dissolved organic matter sources induce rapid transcriptional responses in
- coexisting populations of Prochlorococcus, Pelagibacter and the OM60 clade." <u>Environmental Microbiology</u>
   **16**(9): 2815-2830.
- Shaw, S. L., S. W. Chisholm and R. G. Prinn (2003). "Isoprene production by Prochlorococcus, a marine
  cyanobacterium, and other phytoplankton." <u>Marine Chemistry</u> **80**(4): 227-245.
- 760 Steglich, C., M. Behrenfeld, M. Koblizek, H. Claustre, S. Penno, O. Prasil, F. Partensky and W. R. Hess (2001).
- 761 "Nitrogen deprivation strongly affects Photosystem II but not phycoerythrin level in the divinyl-chlorophyll b-
- 762 containing cyanobacterium Prochlorococcus marinus." <u>Biochimica et Biophysica Acta (BBA) Bioenergetics</u>
   763 **1503**(3): 341-349.
- 764 Steglich, C., C. W. Mullineaux, K. Teuchner, W. R. Hess and H. Lokstein (2003). "Photophysical properties of
- Prochlorococcus marinus SS120 divinyl chlorophylls and phycoerythrin in vitro and in vivo." <u>FEBS Letters</u>
   553(1–2): 79-84.
- 767 Szul, M. J., S. P. Dearth, S. R. Campagna and E. R. Zinser (2019). "Carbon Fate and Flux in
- 768 <em>Prochlorococcus</em> under Nitrogen Limitation." <u>mSystems</u> **4**(1): e00254-00218.
- 769 Talmy, D., A. C. Martiny, C. Hill, A. E. Hickman and M. J. Follows (2016). "Microzooplankton regulation of
- 570 surface ocean POC:PON ratios." <u>Global Biogeochemical Cycles</u> **30**(2): 311-332.
- 771 Thompson, A. W., K. Huang, M. A. Saito and S. W. Chisholm (2011). "Transcriptome response of high- and
- low-light-adapted Prochlorococcus strains to changing iron availability." <u>Isme J</u> **5**(10): 1580-1594.
- Thornton, D. C. O. (2014). "Dissolved organic matter (DOM) release by phytoplankton in the contemporary and future ocean." European Journal of Phycology **49**(1): 20-46.
- Ting, C. S., G. Rocap, J. King and S. W. Chisholm (2002). "Cyanobacterial photosynthesis in the oceans: the
- origins and significance of divergent light-harvesting strategies." <u>Trends in Microbiology</u> **10**(3): 134-142.
- Tolonen, A. C., J. Aach, D. Lindell, Z. I. Johnson, T. Rector, R. Steen, G. M. Church and S. W. Chisholm (2006).
- 778 "Global gene expression of Prochlorococcus ecotypes in response to changes in nitrogen availability." <u>Mol</u>
- 779 <u>Syst Biol</u> **2**: 53.

- 780 Van Mooy, B. A., G. Rocap, H. F. Fredricks, C. T. Evans and A. H. Devol (2006). "Sulfolipids dramatically
- decrease phosphorus demand by picocyanobacteria in oligotrophic marine environments." <u>Proc Natl Acad Sci</u>
  U S A **103**(23): 8607-8612.
- 783 Vargas, M. A., H. Rodríguez, J. Moreno, H. Olivares, J. A. D. Campo, J. Rivas and M. G. Guerrero (1998).
- 784 "BIOCHEMICAL COMPOSITION AND FATTY ACID CONTENT OF FILAMENTOUS NITROGEN-FIXING
- 785 CYANOBACTERIA." Journal of Phycology **34**(5): 812-817.
- 786 Veldhuis, M. and G. W. Kraay (1990). "Vertical distribution and pigment composition of a picoplanktonic
- prochlorophyte in the subtropical North Atlantic:a combined study of HPLC-analysis of pigments and flow
   cytometry." Marine Ecology Progress Series 68: 121-127.
- Zavřel, T., M. Faizi, C. Loureiro, G. Poschmann, K. Stühler, M. Sinetova, A. Zorina, R. Steuer and J. Červený
   (2019). "Quantitative insights into the cyanobacterial cell economy." <u>eLife</u> 8: e42508.
- 791 Zhao, Z., M. Gonsior, J. Luek, S. Timko, H. Janiri, N. Hertkorn, P. Schmitt-Kopplin, X. Fang, Q. Zeng, N. Jiao and
- F. Chen (2017). "Picocyanobacteria and deep-ocean fluorescent dissolved organic matter share similar optical
- 793 properties." Nature communications **8**: 15284-15284.
- 794 Zhu, Y., J. E. Graham, M. Ludwig, W. Xiong, R. M. Alvey, G. Shen and D. A. Bryant (2010). "Roles of
- xanthophyll carotenoids in protection against photoinhibition and oxidative stress in the cyanobacterium
- 796 Synechococcus sp. strain PCC 7002." <u>Archives of Biochemistry and Biophysics</u> **504**(1): 86-99.
- 797 Zimmerman, A. E., S. D. Allison and A. C. Martiny (2014). "Phylogenetic constraints on elemental
- stoichiometry and resource allocation in heterotrophic marine bacteria." <u>Environmental Microbiology</u> 16(5):
  1398-1410.
- Zinser, E. R., D. Lindell, Z. I. Johnson, M. E. Futschik, C. Steglich, M. L. Coleman, M. A. Wright, T. Rector, R.
- 801 Steen, N. McNulty, L. R. Thompson and S. W. Chisholm (2009). "Choreography of the transcriptome,
- 802 photophysiology, and cell cycle of a minimal photoautotroph, prochlorococcus." <u>PLoS ONE</u> **4**(4): e5135.

803