1 2 3 4 5	Responses of globally important phytoplankton groups to olivine dissolution products and implications for carbon dioxide removal via ocean alkalinity enhancement
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#### 36

### 37 Abstract

38

39 Anthropogenic greenhouse gas emissions are leading to global temperature increases, 40 ocean acidification, and significant ecosystem impacts. Given current emissions trajectories, the 41 IPCC calls for both the rapid abatement of CO<sub>2</sub> emissions and development of carbon dioxide 42 removal (CDR) strategies that can address legacy emissions and difficult to abate emissions 43 sources. These CDR methods must efficiently and safely sequester gigatons of atmospheric CO<sub>2</sub>. 44 Coastal Enhanced Weathering (CEW) via the addition of the common mineral olivine to coastal 45 waters is one promising approach to enhance ocean alkalinity for large-scale CDR. As olivine 46 weathers, it releases several biologically active dissolution products, including alkalinity, trace metals, and the nutrient silicate. Released trace metals can serve as micronutrients but may also 47 be toxic at high concentrations to marine biota including phytoplankton that lie at the base of 48 marine food webs. We grew several globally important phytoplankton functional groups under 49 50 elevated concentrations of olivine dissolution products using a synthetic olivine leachate (OL) based on olivine elemental composition, and monitored their physiological and biogeochemical 51 responses. This allowed us to determine physiological impacts and thresholds at elevated olivine 52 leachate concentrations, in addition to individual effects of specific constituents. We found both 53 54 positive and neutral responses but no evident toxic effects for two silicifying diatoms, a calcifving coccolithophore, and three cyanobacteria. In both single and competitive co-cultures, 55 56 silicifiers and calcifiers benefited from olivine dissolution products like iron and silicate or enhanced alkalinity, respectively. The non-N2-fixing picocyanobacterium could use synthetic 57 olivine-derived iron for growth, while N<sub>2</sub>-fixing cyanobacteria could not. However, other trace 58 59 metals like nickel and cobalt supported cyanobacterial growth across both groups. Growth 60 benefits to particular phytoplankton groups in situ will depend on species-specific responses and 61 ambient concentrations of other required nutrients. Results suggest olivine dissolution products 62 appear unlikely to cause negative effects for marine phytoplankton, even at high concentrations, 63 and may support growth of particular taxa under some conditions. Future studies can shed light on long-term evolutionary responses to olivine exposure, and on the potential effects that marine 64 65 microbes may in turn have on olivine dissolution rates and regional biogeochemistry. 66

## 67 Introduction

68

69 Excess anthropogenic greenhouse gas emissions are driving global changes to Earth systems and leading to simultaneous increases in sea surface temperatures, ocean acidification, 70 71 and regional shifts in nutrient supplies (Bach et al., 2019). To counteract these trends and limit 72 the average global temperature increase to 1.5-2°C, carbon dioxide removal (CDR) methods that 73 can collectively remove and permanently store gigatons of atmospheric CO<sub>2</sub> (GtCO<sub>2</sub>) must be 74 developed (Renforth and Henderson, 2017). Coastal Enhanced Weathering (CEW) with olivine 75 (Mg<sub>2-x</sub>Fe<sub>x</sub>SiO<sub>4</sub>) has been proposed as an economically scalable form of ocean alkalinity 76 enhancement (OAE), as it is a globally abundant, naturally occurring ultramafic silicate mineral 77 (Bach et al., 2019; Beerling et al., 2021). Olivine is considered to be one of the most favorable 78 minerals for CDR as it weathers quickly under Earth surface conditions (Renforth and 79 Henderson, 2017; Hartmann et al., 2013; Rimstidt et al., 2012). Similar to other silicate minerals, it dissolves in water to release cations (Mg<sup>2+</sup>, Fe<sup>2+</sup>) and generates alkalinity (principally HCO<sub>3</sub><sup>-</sup>), 80

- 81 with up to 4 mol of  $CO_2$  sequestered per mol of olivine [Eq. 1].
- 82
- 83 84

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86 87  $Mg_{2-x}Fe_{x}SiO_{4} + 4CO_{2} + 4H_{2}O - 2-xMg^{2+} + xFe^{2+} + 4HCO_{3} + H_{4}SiO_{4}$ [Eq. 1]

Forsteritic olivine is the magnesium-rich end-member of olivine and can contain various other trace constituents. For example, olivine used in this study contains ~92% magnesium (Mg<sup>2+</sup>) and ~8% ferrous iron (Fe<sup>2+</sup>) along with trace amounts (<1%) of other metals such as nickel (Ni),

chromium (Cr), and cobalt (Co). As olivine weathers, it releases several biologically important 88 89 dissolution products into the surrounding seawater: (I) bicarbonate ( $HCO_3^-$ ) and carbonate ion (CO<sub>3</sub><sup>2-</sup>), hereafter summarized as "alkalinity"; (II) silicic acid (Si(OH)<sub>4</sub>) hereafter termed 90 91 silicate; (III) and a variety of trace metals including iron ( $Fe^{2+}$ , or oxidized aqueous species), 92 nickel (Ni<sup>2+</sup>), cobalt (Co<sup>2+</sup>), and chromium (CrVI). These dissolution products have the potential 93 to affect important phytoplankton functional groups like silicifying algae (diatoms), calcifying 94 algae (coccolithophores), and cyanobacteria, which lie at the base of marine food webs and drive 95 the biological carbon pump (Moran, 2015). Hence, it is important to understand the specific

effects of these constituents on globally important phytoplankton groups, particularly at elevated
concentrations to simulate large-scale CEW applications.

98 Significant alkalinity additions from olivine weathering can consume  $CO_2$  from the 99 surrounding seawater, causing a  $CO_2$  deficit until air-sea equilibration. This shift in the carbonate 100 system from  $CO_2$  to  $HCO_3^{-}/CO_3^{2^-}$  by transient, non-equilibrated OAE will affect phytoplankton 101 functional groups differently, with some taxa being more sensitive than others. For example, it is 102 predicted that calcifying organisms like coccolithophores may benefit from CEW due to 103 decreases in proton concentrations (H<sup>+</sup>) and increases in the CaCO<sub>3</sub> saturation state.

104 Additionally, dissolving one mole of olivine leads to a one mole increase in dissolved silicate,

105 which is an essential and often bio-limiting nutrient for silicifying organisms like diatoms, a

106 phytoplankton group estimated to contribute up to 40% of the marine primary production

107 (Bertrand et al., 2012). Hence, diatoms may especially benefit from CEW applications with
108 olivine. Additionally, diatoms are particularly noted for being dominant phytoplankton in the
109 coastal regimes where olivine deployments are likely to take place (Field et al., 1998). While
110 there are both planktonic and benthic species of diatoms, the latter will presumably be exposed to
111 especially sustained and elevated levels of dissolution products when olivine is deployed in
112 natural marine sediments. It is unknown if either group, calcifiers or silicifiers, may consistently

- 113 outcompete the other following CEW with olivine (Bach et al., 2019).
- 114 Trace metals like Fe and Ni are general micronutrients required by all classes of 115 phytoplankton, and could potentially support their growth upon fluxes into seawater from olivine weathering. In particular, dinitrogen (N<sub>2</sub>)-fixing cyanobacteria and diatoms both have elevated 116 Fe requirements (Hutchins and Sañudo-Wilhelmy, 2021; Hutchins and Boyd, 2016), and so may 117 stand to benefit from increases in Fe concentrations. Although a required micronutrient at low 118 119 levels, in high enough concentrations Ni may potentially negatively impact phytoplankton 120 growth, although one recent study showed limited to no toxic effects of very high Ni 121 concentrations (e.g. 50,000 nmol L<sup>-1</sup>) for several phytoplankton taxa (Guo et al., 2022). Cobalt 122 can also serve as a micronutrient for phytoplankton (Sunda and Huntsman, 1995; Hawco et al., 123 2020) but may also be toxic at high concentrations (Karthikeyan et al., 2019). However, other 124 trace metals found with olivine such as Cr are not nutrient elements and also need to be 125 considered in terms of their possible toxicity to phytoplankton (Flipkens et al., 2021; Frey et al., 126 1983).
- 127 Hence, it is important to understand the taxon-specific effects of these constituents to 128 determine thresholds at which key phytoplankton functional groups may experience positive or 129 negative effects. Furthermore, it is important to expose phytoplankton to elevated concentrations 130 of olivine dissolution products simultaneously to understand what impacts may occur for large 131 CEW applications. Exposures of organisms to concentrated olivine dissolution products also 132 provides an "worst case scenario" benchmark, which can be compared to lower actual 133 environmental exposures resulting from small CEW additions, slower olivine dissolution time 134 scales, and dilution from advection. While olivine weathers relatively quickly compared to other 135 silicate minerals (Hartmann et al., 2013), dissolution of olivine grains is gradual (i.e. years) relative to microbial physiological responses (hours), posing a challenge to test different 136 137 concentrations of olivine constituents on phytoplankton physiology. To address this, we prepared 138 a synthetic olivine leachate (OL) composed of olivine dissolution products with trace metal 139 concentrations well over those of seawater (17-12,000 times higher), in order to represent a 140 "worst case" scenario for a CEW project. This scenario was estimated based on the maximum 141 expected impact of olivine weathering on the chemistry of the overlying water column. 142 Assuming a 10 cm thick layer of pure olivine sand dissolves with a 100 year half-life dissolving 143 into 1 meter of overlying water with a 24 hour residence time, the anticipated steady state change 144 in the alkalinity of the overlying water column is 65 umol/kg (assuming 4 moles of alkalinity per 145 mole olivine (Meysman and Montserrat, 2017) and 100% release to the water column). The 146 concentrations of other components were chosen assuming stoichiometric congruent dissolution

- 147 and quantitative release to the water column as well -- a worst case scenario. Furthermore,
- 148 phytoplankton were exposed to OL within a small, enclosed batch culture. We cultured 6 species
- 149 representing three globally important phytoplankton functional groups: 2 diatoms (*Nitzschia*,
- 150 *Ditylum*), 1 coccolithophore (*Emiliana*), 2 dinitrogen (N<sub>2</sub>) fixing cyanobacteria (*Trichodesmium*,
- 151 *Crocosphaera*), and 1 non-N<sub>2</sub> fixing picocyanobacterium (*Synechococcus*). All of these species
- are planktonic, with the exception of the diatom *Nitzschia* which frequently forms benthic
- 153 biofilms (Yamamoto et al., 2008). Cultures were grown semi-continuously in natural seawater
- 154 based modified Aquil media (Sunda et al., 2005) with OL as the only available Fe source (and Si
- source for diatoms). For all experiments, cultures were sampled for a basic set of core
- biogeochemical and physiological parameters (Fu et al., 2005, 2008; Tovar-Sanchez et al., 2003;
- 157 Paasche et al., 1996). This approach allowed us to compare and contrast phytoplankton taxon-
- specific responses, including: 1) physiological impacts at extremely high OL concentrations, 2)
- 159 physiological thresholds and dose responses across a range of increasing concentrations of OL,
- and **3**) individual effects of specific OL constituents.
- 161

# 162 Materials and Methods

163

# 164 *Culture growth conditions and experimental set up*

165

166 Six species of phytoplankton were used in these experiments, including two diatoms, *Ditylum* 167 (centric, planktonic) and Nitzschia (pennate, benthic), one coccolithophore, Emiliania huxleyi, 168 one picoplanktonic cyanobacterium (Svnechococcus), and two marine dinitrogen (N<sub>2</sub>) fixing 169 cyanobacteria, Trichodesmium IMS 101 and Crocosphaera WH 0005. Cultures were grown in 170 500 mL polycarbonate flasks at 28°C for the three cyanobacteria, and 20°C for the diatoms and 171 Emiliania huxleyi. Cool-white fluorescent light was supplied following a 12:12 light:dark cycle at an irradiance level of 150 µEm<sup>-2</sup>s<sup>-1</sup>. Stock cultures were grown in natural offshore seawater 172 173 collected with trace metal clean methods (John et al., 2022), which was used to make modified 174 Aquil control medium (ACM) (Sunda et al., 2005). For experiments, cultures were inoculated 175 into the three treatments described below with the addition of  $4\mu$ M phosphate (PO<sub>4</sub><sup>3-</sup>) and 60  $\mu$ M nitrate (NO<sub>3</sub><sup>-</sup>). There was no nitrogen (N) added into the Aquil medium for the  $N_2$  fixers. Iron, 176 177 Cobalt, Nickel (Fe, Co and Ni, required by all species) and silicate (SiOH<sub>4</sub>, required by diatoms 178 only) were not added to the Aquil medium, except as components of the olivine leachate (see 179 below). The background nutrient concentrations in the collected natural seawater were 1µM

- 180 NO<sub>3</sub><sup>-</sup>,  $0.1\mu$ M PO<sub>4</sub><sup>3-</sup> and  $3\mu$ M SiOH<sub>4</sub>.
- 181
- 182 Synthetic olivine leachate preparation
- 183

184 To simulate acute exposure of phytoplankton to elevated levels of olivine dissolution products in

- 185 seawater, we prepared an artificial concentrated OL stock solution based on elemental analyses
  - 186 of commercial ground olivine rock (Sibelco. (2022) Technical Data Olivine Refractory Grade

187 Fine. Antwerp, Belgium). For experimental exposures, this concentrated OL stock was added to

- seawater growth medium to yield the final concentrations shown in **Table 1**, which will be
- referred to throughout as a "100%" concentration of OL. Experiments examining biological
- 190 effects across a dilution range (0-100%) used correspondingly lower additions of the
- 191 concentrated stock. The only other components added to the OL were the major nutrients nitrate
- 192 (60  $\mu$ M NO<sub>3</sub><sup>-</sup>) and phosphate (4  $\mu$ M PO<sub>4</sub><sup>3-</sup>). These are required nutrients for growth of
- 193 phytoplankton, and were added at the same concentration to the OL and the Aquil control
- medium. For experiments with N<sub>2</sub>-fixing cyanobacteria, nitrate was omitted and phosphate wasthe only nutrient added.
- 196

**Table 1.** Concentrations of added ions or compounds in serial dilutions of synthetic olivine leachate (OL, 0% to
 100%) used in the phytoplankton growth experiments; and concentrations of components in the three concentrated

stocks used to prepare experimental medium (1mL/L added for 100% OL). Stock C was prepared in 10 nM HCl to keep the trace metals in solution until addition.

	OL	$Mg^{2+}$	SiOH <sub>4</sub>	OH-	Fe(II)	Ni(II)	Cr(VI)	Co(II)
	Added	(µM)	(µM)	(µM)	(µM)	(µM)	(µM)	(µM)
Concentration	100%	44.9	25	100	3.36	0.13	0.12	0.006
added to	80%	35.9	20	80	2.7	0.10	0.10	0.005
growth medium	50%	22.5	12.3	50	1.7	0.07	0.06	0.003
	30%	13.5	7.5	30	1.0	0.04	0.04	0.002
	10%	4.5	2.5	10	0.34	0.01	0.01	0.001
	0%	0	0	0	0	0	0	0
Concentrated		MgCl <sub>2</sub>	NaSiO <sub>2</sub>	NaOH	FeCl <sub>2</sub>	NiCl <sub>2</sub>	$K_2CrO_4$	CoCl <sub>2</sub>
stock solutions								
(mM)								
Stock A		44.9						
Stock B			25	100				
Stock C					3.36	0.13	0.12	0.006

201

# 202 *Experimental methods*

203

204 Semi-continuous culturing methods were used to achieve nearly steady-state growth. Cultures 205 were diluted with fresh medium every 2 or 3 days, using in vivo fluorescence as a real time 206 biomass indicator. Dilutions were calculated to bring the cultures back down to the biomass 207 levels that were recorded after the previous day's dilution. In this way, cultures were allowed to 208 determine their own growth rates under each set of experimental conditions, without ever nearing 209 stationary phase, significantly depleting nutrients or self-shading (Fu et al., 2022). For all 210 experiments, cultures were sampled for a basic set of core biomass and physiological parameters, including cell counts, CO<sub>2</sub> fixation, particulate organic carbon (POC), particulate organic 211

- 212 nitrogen (PON), particulate organic phosphorus (POP) and biogenic silica (BSi, diatoms only)
- 213 once steady-state growth was obtained for each growth condition (typically after 8–10

generations). Steady-state growth status was defined as no significant difference in cell- or in
 vivo-specific growth rates for at least 3 consecutive transfers.

- 216
- 217 There were four sets of experiments in this project:
- 218

*1) Acute responses to elevated olivine leachate levels.* The goal of this set of experiments was to
investigate the responses of the diatoms *Nitzschia* and *Ditylum* to relatively high concentrations
of olivine leachate, in order to determine acute exposure responses. To see if the leachate may
have a positive or negative effect on their physiology, they were compared to their respective
control cultures. There were a total of three treatments consisting of: OL (100%), ACM, and

- ACM with low Fe/Si (with 2 nM Fe EDTA added, and no added SiOH<sub>4</sub>).
- 225

226 2) *Responses to a broad range of olivine leachate levels*. In these experiments, *Synechococcus*,

227 Crocosphaera, Ditylum, and Emiliania huxleyi were grown in culture medium across a series of

228 OL dilutions (Table 1) to determine their responses across a range of leachate concentrations,

- from high to very low-level exposures.
- 230

3) Fe bioavailability and Cr toxicity from olivine leachate to N<sub>2</sub>-fixing cyanobacteria. The goal
of this set of experiments was to investigate OL-derived Fe bioavailability to N<sub>2</sub>-fixing
cyanobacteria, *Trichodesmium* and *Crocosphaera*. An additional experiment was conducted to
investigate potential Cr(VI) toxicity.

235

236 4) Two species co-culture competition experiments during olivine leachate exposure. In order to 237 test how OL may affect co-existence and competition between the diatom Ditylum and the 238 coccolithophore *Emiliania huxleyi*, a simple batch co-culture competition experiment was carried 239 out in which the 2 species were inoculated at a 1:1 ratio (based on equivalent levels of cellular 240 Chlorophyll a due to the large differences in their cell sizes) into 100% OL and regular ACM, 241 and grown for 10 days until early stationary phase. In vivo fluorescence and cell counts were 242 monitored daily. Relative abundance and growth rates of the two species were determined based 243 on microscopic cell counts during the exponential growth phase of the mixed cultures. Biogenic 244 silica (BSi, an indicator of diatom abundance) and particulate inorganic carbon (PIC or calcite, 245 an indicator of coccolithophore abundance) were collected every other day in order to further 246 determine how these two species responded to co-culture with and without leachate additions.

- 247
- 248 Analytical methods
- 249

<u>Determination of growth rates and chlorophyll a</u> Growth rates were determined based on both
 microscopic cell counts and chlorophyll a. For chlorophyll a determination, subsamples of 30 ml
 from each triplicate bottle were GF/F filtered, extracted in 6 ml of 90% acetone, stored overnight
 in the dark at -20°C, and chlorophyll a concentrations were measured fluorometrically using a

254 Turner 10-AU fluorometer (Welschmeyer, 1994). Specific growth rates were determined using 255 the equation:

256

257

 $\mu = \frac{\ln\left(\frac{N_{Tfinal}}{N_{Tinitial}}\right)}{T_{final} - T_{initial}}$ 258

where  $\mu$  is the specific growth rate (per day) and N is the chlorophyll a concentration at T<sub>initial</sub> 259 260 and T<sub>final</sub> (Kling et al., 2021).

261

262 Particulate C, N, P and Si. Particulate organic carbon and nitrogen (CHN) samples from all 263 experiments were filtered (pre-combusted GF/F) and frozen for analysis using a Costech 264 Elemental Analyzer (Hutchins et al., 2007). Samples for biogenic silica (BSi) were analyzed according to (Brzezinski, 1985). POP (particulate organic phosphorus) samples were collected 265 266 onto pre-combusted 25 mm GF/F filters and analyzed as in Fu et al. 2005 (Fu et al., 2005). 267

268 *Primary productivity*. For all species other than the coccolithophore (see below), primary

269 production was measured in triplicate using 24h incubations (approximating net PP) with 270 H<sup>14</sup>CO<sub>3</sub> under the appropriate experimental growth conditions for each treatment (Fu et al.,

- 271 2008). CO<sub>2</sub> fixation rates were calculated using measured final experimental DIC concentrations 272 and biomass. All samples for primary production were counted using a Wallac System 1400 273 liquid scintillation counter.
- 274

275 Photosynthetic and calcification rates of Emiliania huxlevi. For the coccolithophore, two 40 mL 276 subsamples from each triplicate bottle were spiked with 0.5 µCi NaH<sup>14</sup>CO<sub>3</sub>. One subsample was 277 incubated in the light and the other in the dark for 24 h. Then two sets of 20 mL aliquots from 278 each sub- sample were filtered onto Whatman GF/F filters. The filters for photosynthetic rate 279 determination were fumed with saturated HCl before adding scintillation cocktail fluid. 280 Photosynthetic rate and calcification rate were calculated as described in Paasche et al. 1996 281 (Paasche et al., 1996).

282

283 Fe quota. Intracellular Fe content was determined by filtering culture samples onto acid-washed 284 0.2-µm polycarbonate filters (Millipore), and rinsing with oxalate reagent to remove extracellular 285 trace metals (Tovar-Sanchez et al., 2003). Fe was determined with a magnetic sector-field high-286 resolution inductively coupled plasma mass spectrometer (ICPMS) (Element 2, Thermo) (Jiang 287 et al., 2018; John et al., 2022).

288

289 Statistical methods. A one-way ANOVA analysis of variance was used to analyze differences 290 between treatments using Prism 8. Differences between treatments were considered significant at 291 p<0.05. Post-hoc comparisons were conducted using the Tukey's multiple comparison test to

292 determine any pairwise differences.

293

#### 294 **Results**

# 295

#### 296 Diatoms

297 We hypothesized that diatoms may benefit from the OL products Si and Fe, as they are 298 both required for growth and can be limiting for this group (Tréguer et al., 2018). Hence, we 299 grew the benthic diatom Nitzschia across three treatments: OL alone, Aquil control medium 300 (ACM), and ACM but with low, limiting Si and Fe concentrations (ACM-low-SF). Nitzschia grew and fixed carbon just as well in the OL as the ACM (p=0.35; p=0.21), while showing 301 302 reduced rates in the ACM-low-SF treatment (p=0.02; p< 0.0001; Fig. 1A, B). Likewise, the 303 particulate Si:C ratios demonstrated OL to be just as good a source of Si to Nitzschia as the ACM 304 (p=0.98), and considerably better than the ACM-low-SF (p=0.0012; Fig. 1C). 305

Growth and CO<sub>2</sub> fixation rates of the planktonic diatom *Ditylum* were significantly higher



Fig 1. Effects of olivine leachate versus culture medium controls on growth and physiology of the benthic diatom Nitzschia. A) Cell-specific growth rates (d<sup>-1</sup>), **B**) Carbon-specific fixation rates (hr<sup>-1</sup>) and **C**) Si:C ratios (mol:mol) of the benthic diatom Nitzschia sp. Abbreviations: OL is olivine leachate, ACM is Aquil control medium, ACM-low-SF is Aquil control medium with lowered Si and Fe concentrations. Values represent the means and error bars are the standard deviations of triplicate cultures for each treatment.

in the OL treatment compared to either the ACM (p=0.002; p=0.0001) or ACM-low-Si/Fe treatments (p=0.0002; p<0.0001; Supp. Fig. 1 A, **B**), while Si:C ratios were the same (p=0.93; Supp. Fig 1C). When *Ditylum* was grown across a range of OL concentrations (i.e., a dilution series from 0% to 100% additions, where 100% corresponds to the OL treatment used in the Nitzschia experiment above), we observed increasing growth and CO<sub>2</sub>

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324 fixation rates with increasing OL concentrations, with maximum rates observed at and above 325 50% of the original OL (Fig. 2A, B). Ditvlum particulate Si:C ratios also reached levels similar 326 to those seen in the ACM medium in the 80-100% additions (Fig. 2C). Likewise, *Ditylum* 327 cellular Fe:P ratios measured by ICP-MS were not significantly different between 100% OL and 328 ACM treatments, suggesting the diatom could access the same amount of Fe from the 329 precipitated Fe(III) in the OL as from the soluble (EDTA-chelated) Fe(III) in the culture medium 330 (p=0.56; Supp. Fig 2A). These data demonstrate that even at extremely high concentrations, 331 olivine dissolution products including trace metals are not toxic to diatoms, but instead may 332 provide sources of the essential nutrients iron and silicate to support their growth in nutrient

333 replete conditions.

#### 334 *Coccolithophores*

It has been hypothesized that calcifying coccolithophores may benefit from an increase in alkalinity from olivine dissolution (Bach et al., 2019). In the OL dilution series, maximum growth rates for the coccolithophore *Emiliana* equivalent to those recorded in the ACM positive



control medium were achieved at all added OL concentrations from 10% to 100% (Fig. 2D). POC production (CO<sub>2</sub>) fixation) rates were similar at all OL levels from 30% to 100%, and were not significantly different than in the ACM treatment (p >0.05, Fig. 2E). Particulate inorganic carbon to particulate organic carbon fixation ratios (PIC:POC production ratios) were higher at OL levels of 50-100% than in the ACM positive controls (**Fig. 2F**), possibly due to enhanced alkalinity in the high OL concentration treatments. An independent

set of basic twotreatment experiments

- 366 with the coccolithophore (ACM versus 100% OL, Supp. Fig 3) supported the results of the
- dilution series experiments shown in **Fig. 2**. *Emiliana* specific growth rates (p=0.05), and
- 368 cellular particulate inorganic:particulate organic carbon ratios (p= 0.07; PIC:POC, mol:mol)
   369 were very similar in the ACM and OL treatments (Supp. Fig. 3A). Likewise, POC-specific
- fixation rates (p=0.04; TC  $h^{-1}$ ) were slightly higher in the OL than in the ACM treatments, while
- 371 PIC:POC fixation ratios were the same (p=0.07, Supp. Fig. 3B). Similar to diatoms, these data
- 372 demonstrate that olivine dissolution products are also not toxic to coccolithophores and that
- and alkalinity may support growth in nutrient replete conditions.

### 374

# 375 Cyanobacteria

Like diatoms and coccolithophores, cyanobacteria could benefit from olivine dissolution 376 377 due to their relatively high Fe (Hutchins and Boyd, 2016) and Ni requirements (Dupont et al., 378 2008). The OL dilution series experiments using the widely distributed picocyanobacterium 379 Synechococcus showed positive responses in growth rates (Fig. 3A) and CO<sub>2</sub> fixation rates (Fig. 380 **3B**) across the range of OL levels, similar to those of the eukaryotic algae. Both growth rates and carbon fixation rates were the same in the 100% OL treatment as in the ACM positive 381 382 control treatment (p=0.94; p=0.46). ICP-MS measurements of Synechococcus cellular Fe:P 383 ratios across a range of OL levels (0-100%) showed that this isolate accumulated much less Fe in 384 the 0% OL than in the ACM treatment (p=0.02), but in all treatments with added OL, Fe:P ratios 385 were the same as or higher than the ACM values (Supp. Fig. 2B). As with the eukaryotic 386 phytoplankton, the synthetic OL provided a good source of Fe to support the growth of the 387 picocyanobacterium.



**Fig 3. Dilution series of olivine leachate on growth and CO<sub>2</sub> fixation of a marine cyanobacterium.** The unicellular picocyanobacterium *Synechococcus* was grown across a range of dilutions of the olivine leachate (OL, 0-100%), and in the Aquil control medium (ACM). Shown are: **A)** Cell-specific growth rates (d<sup>-1</sup>) and **B)** Carbon-specific fixation rates (hr<sup>-1</sup>). Values represent the means and error bars are the standard deviations of triplicate cultures for each treatment.

In striking contrast to the eukaryotic algae and the nondiazotrophic (i.e., non-N2-fixing) picocyanobacterium Synechococcus, the N<sub>2</sub>-fixing cyanobacterium *Trichodesmium* could not grow at any concentration of OL tested (Fig. 4A). One possible explanation for this lack of growth is toxic effects by one of the trace metal components of the OL. We hypothesized that Ni and Co are less likely to be toxic, as these nutrient metals have been found to be relatively non-toxic to phytoplankton at similar environmental concentrations (Guo et al., 2022; Karthikeyan et al., 2019; Panneerselvam et al., 2018). Hence, we hypothesized that Cr toxicity should be considered as a likely

- 408 possible scenario (Frey et al., 1983; Kiran et al., 2016).
- 409 Another possibility is that *Trichodesmium* did not experience toxic effects but instead
- 410 was unable to access Fe from OL. This N<sub>2</sub>-fixer requires more Fe than virtually any other
- 411 phytoplankton species (Hutchins and Sañudo-Wilhelmy, 2021), and the OL was the only source
- 412 of Fe provided in our experiments. Fe(II) released into seawater from olivine dissolution likely
- 413 quickly oxidizes to Fe(III), which then precipitates and becomes insoluble at the elevated



**Fig 4. Effects of olivine leachate versus culture medium controls on growth of the marine N<sub>2</sub>fixing cyanobacterium** *Trichodesmium*. Shown are **A)** Cell-specific growth rates (d<sup>-1</sup>) of the colonial cyanobacterium *Trichodesmium* across a range of dilutions of the olivine leachate (OL, 0-100%) and in the Aquil control medium (ACM), and **B)** Cell-specific growth rates (d<sup>-1</sup>) of *Trichodesmium* in two concentrations of OL (30% and 100%) with the synthetic metal chelator EDTA, and in OL without Cr, or OL without Cr but plus EDTA, versus ACM. concentrations in our OL (Manck et al., 2022). This could render it biologically unavailable to the cellular Fe uptake systems of some species. We deliberately designed our OL to replicate this oxidation/precipitation process, and as expected observed visible reddish-brown amorphous Fe precipitates on the bottom of the growth flasks for all synthetic OL treatments.

Accordingly, we designed another set of experiments to test for both lack of Fe bioavailability and specific sensitivity to Cr, as has been done in previous cyanobacterial studies (Kiran et al., 2016). To do this, we formulated several variants of the olivine leachate: 1) normal OL (100% concentration), 2) OL (100% concentration) with a synthetic ligand (EDTA) that solubilizes Fe(III), and thus makes it broadly bioavailable (OL-EDTA), 3) OL (100% concentration) but with no Cr (OL-noCr), 4) and OL (100% concentration) but with no Cr and with

EDTA (OL-EDTA-noCr) (Fig. 4B). *Trichodesmium* also could not grow in the OL medium
without added Cr (OL-noCR, Fig. 4B), demonstrating that the lack of growth observed in OL
was not due to Cr toxicity. However, growth recovered to the same levels as in the ACM when
EDTA was added (OL-EDTA) to the leachate (p=0.16; Fig. 4B). This suggests that poor
bioavailability of the precipitated Fe(III) was the likely cause for *Trichodesmium*'s inability to
grow in the unmodified OL.

440 OL also inhibited the growth of the unicellular N<sub>2</sub>-fixing cyanobacterium Crocosphaera, 441 although not to the same extent as for Trichodesmium. Crocosphaera exhibited growth rates that 442 were 22-44% of those in ACM across the full range of OL concentrations, but growth recovered 443 in OL-EDTA to 76% of rates in ACM (Fig. 5A). Results were very similar for CO<sub>2</sub> fixation rates 444 and N<sub>2</sub>-fixation rates in OL, which were severely reduced by 64-100% (carbon fixation) and 69-445 88% (N<sub>2</sub> fixation) relative to ACM in all OL treatments, but reached maximum values of 80% 446 and 63% of ACM treatment rates, respectively, when EDTA was added to the OL (Fig. 5B and C). This demonstrates that oxidized Fe from OL was not effectively utilized to support growth 447 448 for either of the two  $N_2$ -fixing cyanobacteria tested, in contrast to diatoms, coccolithophores, and 449 Synechococcus. However, their growth recovery after EDTA additions indicates that the other 450 trace metals Cr, Ni, and Co in the olivine leachate were not toxic, even at extremely high 451 concentrations. Interestingly, unlike *Trichodesmium* which either could not grow at all on OL 452 alone or recovered fully upon EDTA additions, Crocosphaera could still grow at lower growth 453 rates on OL but could not grow as fast upon EDTA additions as in ACM. Future experiments are





A) Cell-specific growth rates (d<sup>-1</sup>), B) Carbon-specific fixation rates (hr<sup>-1</sup>), and C) N-specific fixation rates (day<sup>-1</sup>) of the unicellular cyanobacterium *Crocosphaera* grown across a range of dilutions of the olivine leachate (OL, 0-100%), in 100% OL plus EDTA (OL-EDTA), and in the Aquil control medium (ACM). Values represent the means and error bars are the standard deviations of triplicate cultures for each treatment.

needed to understand these differences in speciesspecific responses between these two N<sub>2</sub>-fixers. Taken together, these data suggest that when olivine dissolves in seawater, it will likely have negligible or no effect (positive or negative) on these cyanobacteria.

Diatom/Coccolithophore Competitive Co-culture

Results of the coculture, or competition, experiment with the diatom *Ditylum* and the coccolithophore *Emiliania* 

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472 *huxleyi* are shown in **Fig. 6**. Unlike the semi-continuous experiments shown in the previous figures, this experiment used closed-system "batch" culturing methods in order to assess and 473 474 compare effects on relative biomass accumulation by each species over time. OL (100% 475 concentration) supported growth of both the diatom (Fig. 6A) and the coccolithophore (Fig. 6B) 476 in mixed culture, and biomass was very similar for both species between the OL and ACM 477 treatments throughout most of the experiment. However, cell yields were higher in the ACM at 478 the final timepoint for the diatom (p = 0.009, Fig. 6A). Final cell counts were also higher in the 479 ACM for the coccolithophore, but this difference was not significant (p=0.31; Fig. 6B). Similar 480 trends were observed when diatom biomass was estimated as biogenic silica (BSi, Fig. 6C, p =0.002) and when coccolithophore biomass was assessed as calcite or particulate inorganic 481 482 carbon (PIC, Fig. 6D, p = 0.04). For the diatom, OL supported growth rates similar to those in the ACM treatment during the first half of the experiment (Fig. 6A.C; Supp. Fig. 4A). Growth 483 484 rates were also similar in the OL and ACM mediums for the coccolithophore (Fig. 6B,D; Supp. 485 Fig. 4B). Hence, both phytoplankton groups were able to grow similarly well in co-culture where 486 each did not exhibit any strong competitive advantage over the other.

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## 489 Discussion

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In general, simulated olivine dissolution products did not show toxicity even at extremely
high concentrations across all phytoplankton groups, consistent with other recent observations
(Guo et al., 2022). Guo et al. (2022) particularly focused on exposing 11 phytoplankton groups



Fig. 6. Effects of olivine leachate versus culture medium controls on growth competition and biomineralization during co-culture of a diatom and a coccolithophore. Shown are 5 day growth curves (cells mL<sup>-1</sup>) for A) the coccolithophore *Emiliana huxleyi* and B) the diatom *Ditylum* in mixed cultures grown in olivine leachate (OL, red symbols) and Aquil control medium (ACM, blue symbols). Also shown are C) Biogenic silica (BSi,  $\mu$ mol L<sup>-1</sup>), a proxy for diatom biomass, and D) Calcite or particulate inorganic carbon (PIC,  $\mu$ mol L<sup>-1</sup>), a proxy for coccolithophore biomass, in the OL and ACM treatments in the same growth competition experiment. Values represent the means and error bars are the standard deviations of triplicate cultures for each treatment. to elevated Ni concentrations and did not observe strong effects across these taxa. Although it is unknown what chemical species of dissolved Ni primarily influence phytoplankton physiology, most studies indicate that phytoplankton primarily interact with free Ni<sup>2+</sup> ions but are not particularly sensitive to the total dissolved Ni concentration (Guo et al., 2022). Guo et al. (2022) and our study used the same Ni-containing compound, NiCl<sub>2</sub>, as a source of Ni<sup>2+</sup>. Guo et al. also used the same base Aquil control medium as our ACM. ACM contains EDTA that binds with metal ions like Ni to improve their dissolution, which subsequently lowers the free Ni ion concentrations (e.g.,  $Ni^{2+}$ ) relative to the total dissolved Ni pool. Although broad negative effects of enhanced Ni concentrations were not observed across taxa, Guo observed some

521 species-specific responses across variations in  $(0-100 \ \mu\text{M})$  EDTA and Ni  $(0-50 \ \mu\text{M})$ 522 concentrations, indicating that specific phytoplankton groups are impacted differently depending on the chemical species in the total dissolved Ni pools and/or the concentration and type of 523 524 organic ligands in seawater. Our results are generally consistent with their overall findings, as the 525 phytoplankton groups tested here did not exhibit negative effects upon elevated exposure to Ni 526 with (e.g., ACM) and without added EDTA (e.g., OL), suggesting that Ni was not toxic irrespective of the concentration of different Ni species in the dissolved pool or that of the Ni<sup>2+</sup> 527 528 ion. However, our experiments were not designed to test for taxon-specific differences in 529 responses to specific Ni species or variations in EDTA concentrations in particular.

530 Diatoms were able to use synthetic OL-derived Si and Fe to support near-maximum 531 growth rates and carbon fixation rates, as well as robust silica frustule development; both of 532 these nutrients can frequently limit diatom growth in various parts of the ocean (Tréguer et al., 533 2018; Hutchins and Boyd, 2016). OL-derived alkalinity and iron increases also supported

coccolithophore growth, consistent with previous observations in calcifying corals (Albright et 534 535 al., 2016). Similarly, the globally-distributed picocyanobacterium Synechococcus increased its 536 growth and carbon fixation rates as OL concentrations increased. Although OL could not support 537 N<sub>2</sub>-fixing cyanobacteria due to their inability to use Fe(III), olivine dissolution products were not 538 observed to be toxic. Their inability to use Fe(III) is a neutral effect due to other sources of 539 bioavailable Fe in the water column (Hutchins and Boyd, 2016). Thus, these results suggest that 540 many phytoplankton will not be negatively impacted by even high levels of elements derived 541 from olivine dissolution, and that some olivine dissolution products may support their growth, 542 primary productivity, and biomineralization when OL is available at high enough concentrations 543 in certain environmental conditions. For example, it is important to note that potential growth 544 benefits to phytoplankton in situ will also depend on ambient concentrations of other important nutrients, such as nitrogen (N) and phosphorus (P). Our cultures contained an abundance of other 545 546 required nutrients, thus enabling phytoplankton to take advantage of particular dissolution 547 products for growth (e.g., Si, Fe, alkalinity). However, if nutrients like N and P are primarily 548 limiting in natural environments, then olivine dissolution products are not expected to have any 549 growth effect. In addition, these cultures represent closed systems that do not allow olivine 550 products to be diluted with fresh seawater. In natural settings, advection in both sediment 551 porewaters (Reimers et al., 2004) and the water column (He and Tyka, 2022) will lead to short 552 residence times, thereby rapidly diluting olivine dissolution products. Hence, these physical 553 dynamics will prevent high concentrations of olivine dissolution products from accumulating in 554 seawater in coastal systems. Thus, even the most dilute leachate treatment in this study is likely 555 more concentrated than the anticipated concentrations of olivine dissolution products expected 556 under field conditions.

557 Bach et al. (2019) (Bach et al., 2019) hypothesized that silicate, iron and nickel releases 558 from marine applications of silicate minerals like olivine might particularly benefit diatoms and 559 cyanobacteria, as these groups have especially high requirements for one or more of these 560 nutrients. Thus, they expected that olivine applications might produce a "Greener" ocean. They 561 also suggested that adding minerals derived from CaCO<sub>3</sub> (such as quicklime applications) would 562 particularly favor coccolithophores, due to rapidly enhanced seawater alkalinity. This outcome would produce a "Whiter" ocean (the color of coccolithophore calcite). Although we did not test 563 564 CaCO<sub>3</sub><sup>-</sup> derivatives, our results with synthetic OL seem to point instead to a both "Green and 565 White" ocean, since in individual experiments diatoms, picocyanobacteria, and coccolithophores 566 all responded positively to OL at the relatively elevated levels applied in our experiments. This 567 conclusion is further supported by the results of our diatom/coccolithophore co-culture 568 experiment, which showed that OL stimulated both species simultaneously rather than conferring 569 a competitive advantage on one or the other.

570 Iron in olivine minerals is present as reduced Fe(II), and we added it in this form to our
571 synthetic OL. However, when Fe(II) dissolves in oxic seawater, it quickly (within minutes)
572 oxidizes to highly insoluble Fe(III), which almost entirely precipitates out as amorphous iron
573 oxyhydroxides (Millero et al., 1987). Clearly, in our experiments this oxidized particulate iron

must have been available to the species that showed growth enhancement with OL, since no
other iron source was provided in the seawater growth medium. Diatoms and some other
eukaryotes can access precipitated Fe(III) oxyhydroxides using their well-studied reductive
uptake systems (Morrissey and Bowler, 2012), or even potentially through endocytosis in some

578 cases (Kazamia et al., 2018).

579 The responses of the two N<sub>2</sub>-fixing cyanobacteria were in striking contrast to those of the 580 other three phytoplankton groups tested. These diazotrophs were either unable to grow in our 581 artificial OL at all (*Trichodesmium*), or could only grow to a very limited degree 582 (Crocosphaera). Since olivine dissolution products support photoautotrophic growth by other phytoplankton but not diazotrophic growth, this could be taken to indicate that olivine 583 584 applications might tend to drive the marine ecosystem towards nitrogen limitation, which would limit the upper bound of phytoplankton proliferation. However, our results with experimental 585 additions of the artificial iron chelator EDTA (ethylene diamine tetra acetic acid) suggest that 586 587 other mechanisms may enable iron bioavailability. For example, previous research has suggested 588 that Trichodesmium cannot directly access particulate Fe(III) forms, but likely relies on bacteria 589 residing on and in natural colonies to produce siderophores, which then solubilize particulate 590 Fe(III) sources and make them bioavailable (Rubin et al., 2011; Lee et al., 2018). Since cultured 591 Trichodesmium such as ours typically do not produce colonies, but grow instead as individual 592 filaments of cells, cultures of this diazotroph are likely deficient in many of these iron-acquiring 593 microbial symbionts (Rubin et al., 2011). The iron uptake systems of Crocosphaera have been 594 less well-characterized, but like Trichodesmium, molecular studies suggest this unicellular 595 diazotroph lacks the genetic capacity to produce endogenous siderophores (Shi et al., 2010; Yang 596 et al., 2022). Our results show that when we add the artificial iron chelator EDTA (which 597 substitutes for ligands produced by the missing bacteria in cultures), the synthetic OL supports 598 near-maximum growth of both of these diazotrophs. Thus, reduced growth rates of these 599 cyanobacteria in OL without EDTA appear to be due to severe iron limitation, not toxicity of any 600 OL component. In our experiments the cells were forced to grow on OL as a sole source of iron, 601 but in coastal ecosystems where olivine deployments would occur, there are typically many other 602 natural sources of iron to support algal growth (Capone and Hutchins, 2013; Hutchins and Boyd, 603 2016). In nature, *Trichodesmium* is also likely to occur mostly as colonies, and so may have 604 access to additional microbiome-provided iron, including from both naturally-occurring supplies 605 as well as potentially from any shallow-water olivine applications. Thus, iron limitation of N<sub>2</sub> 606 fixation due to olivine additions in coastal waters may not occur in the real ocean.

Growth inhibition of diazotrophs by our synthetic OL appears to be due to iron limitation,
but our experiments also shed light on potential effects of other trace metals present in the
formulation. Of the metals found in our synthetic OL, Ni and Co are considered nutrient
elements with relatively low toxicity; in fact, the concentrations added even in our maximum
dosage experiments were well below those that have been reported to be toxic to phytoplankton
(Karthikeyan et al., 2019; Vink and Knops, 2023). However, Cr has the potential to be
biologically problematic. Cr(III) found in olivine is relatively insoluble, so in this form it is

probably not a major source of exposure for planktonic organisms. However, if it oxidizes to 614 615 Cr(VI), it becomes much more soluble, and thus more bioavailable and potentially toxic. Cr(III) 616 oxidation is thermodynamically unfavorable, but can be facilitated by borate ions always present 617 in seawater, or by the presence of biologically or photochemically-produced oxidants like H<sub>2</sub>O<sub>2</sub> 618 (Pettine et al., 1991), and by naturally occurring manganese oxides (Weijden and Reith, 1982). For these reasons, following the principle of "worst case scenario", we used a soluble Cr(VI) salt 619 620 in our synthetic OL formulation. Despite this, we found that the presence or absence of the 621 relatively elevated levels of dissolved Cr(VI) present in our regular synthetic OL did not make 622 any difference to the growth of *Trichodesmium* or the other tested phytoplankton species. 623 Particularly, because synthetic OL stimulated near-maximum growth rates in the diatoms, 624 coccolithophore and picocyanobacterium, we presume that the Cr(VI) additions did not

625 adversely affect these groups either.

626 The goal of this work was to test both extreme levels and simultaneous exposure of 627 multiple, biologically important olivine dissolution products that could influence microbial 628 physiology in order to identify thresholds and response curves. Accordingly, our experiments 629 focused on determining acute effects of high concentrations of olivine dissolution products. In 630 general, they suggest that negative impacts may be few even for large olivine deployments, given 631 the high concentrations of tested olivine dissolution products. Because these microplankton serve 632 as important links to higher trophic levels, these data suggest minimal long-term impacts from 633 olivine dissolution on ecosystem services like fisheries. Future research directions may include 634 longer term experiments with prokaryotes and natural microbial communities to expand our 635 understanding of olivine exposure on important taxa that help drive biogeochemical cycling in 636 the oceans. Similar experiments can also be conducted except with other OAE feedstocks 637 harboring different chemical compositions and more rapid dissolution timescales (Renforth and 638 Henderson, 2017). Future studies can also focus on determining how biological processes like 639 photosynthesis, respiration, and organic ligand production could influence olivine dissolution 640 kinetics and their impacts on carbon dioxide removal.

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analyzed the data. D.A.H., F.-X.F., S.-C.Y, N.G.W., S.J.R., M.G.A., and S.G.J. wrote the paper.

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647 N.G.W., M.G.A., and S.J.R. are full time employees at Vesta, PBC.

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