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5	differential domoic acid toxicity in Pseudo-nitzschia australis
6	
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# 21 <sup>1</sup>Abbreviations

CCMs carbon concentrating mechanisms

DA domoic acid

<sup>&</sup>lt;sup>1</sup>NUEs nitrogen use efficiencies

### 22 Abstract

Along the west coast of the United States, highly toxic Pseudo-nitzschia blooms have been 23 24 associated with two contrasting regional phenomena: seasonal upwelling and marine heatwaves. While upwelling delivers cool water rich in pCO<sub>2</sub> and an abundance of macronutrients to the upper 25 water column, marine heatwaves instead lead to warmer surface waters, low pCO<sub>2</sub>, and reduced 26 27 nutrient availability. Understanding Pseudo-nitzschia dynamics under these two conditions is 28 important for bloom forecasting and coastal management, yet the mechanisms driving toxic bloom 29 formation during contrasting upwelling vs. heatwave conditions remain poorly understood. To 30 gain a better understanding of what drives *Pseudo-nitzschia australis* growth and toxicity during these events, multiple-driver scenario or 'cluster' experiments were conducted using temperature, 31 pCO<sub>2</sub>, and nutrient levels reflecting conditions during upwelling ( $13^{\circ}C$ , 900 ppm pCO<sub>2</sub>, replete 32 33 nutrients) and two intensities of marine heatwaves (19°C or 20.5°C, 250 ppm pCO<sub>2</sub>, reduced 34 macronutrients). While P. australis grew equally well under both heatwave and upwelling 35 conditions, similar to what has been observed in the natural environment, cells were only toxic in the upwelling treatment. We also conducted single-driver experiments to gain a mechanistic 36 understanding of which drivers most impact P. australis growth and toxicity. These experiments 37 38 indicated that nitrogen concentration and N:P ratio were likely the drivers that most influenced 39 domoic acid production, while the impacts of temperature or  $pCO_2$  concentration were less 40 pronounced. Together, these experiments may help to provide both mechanistic and holistic

ENSO El Niño-Southern Oscillation

**pCO**<sub>2</sub> partial pressure of carbon dioxide

PDO Pacific Decadal Oscillation

**POC** particulate organic carbon

perspectives on toxic *P. australis* blooms in the dynamic and changing coastal ocean, where cells
interact simultaneously with multiple altered environmental variables.

43

## 44 Keywords

45 Harmful diatom blooms, *Pseudo-nitzschia*, domoic acid, upwelling, marine heatwaves, biotoxins46

# 47 1. Introduction

48 The biologically-rich California Current System is increasingly impacted by two contrasting 49 extreme phenomena: stronger upwelling events and marine heatwaves. These events differentially impact marine ecosystem function through distinct alteration of the bottom-up controls that 50 51 influence phytoplankton physiology and community structure. Seasonal upwelling-favorable wind 52 conditions vertically transport cold, carbon dioxide-  $(CO_2)$  and nutrient-rich deeper water to the surface, fueling high rates of primary productivity (Chavez and Messié 2009; Gruber et al. 2012; 53 54 Capone and Hutchins 2013). On the other hand, surface ocean warming during marine heatwaves 55 can enhance stratification, consequentially reducing the availability of inorganic nutrients and carbon available for primary producers (Cheung and Frölicher 2020; Gupta et al. 2020). 56

57

Climate change is predicted to amplify the intensity of both of these contrasting coastal processes in the California Current System, which may fundamentally alter the marine environment and conditions for phytoplankton (Du and Peterson 2013; Zhu et al. 2017; Smith et al. 2018; Barth et al. 2020; Trainer et al. 2020). Stronger and more frequent upwelling favorable winds could increase the influx of cold water and nutrients, counteracting the effects of warming and enhanced stratification that accompany rising global temperature (Bakun et al. 2015). These conditions may 64 favor dense and diverse diatom-dominated phytoplankton communities, as this group thrives with high inputs of nutrients (Lassiter et al. 2006; Du and Peterson 2018). Additionally, ocean 65 66 acidification is likely to intensify with the increased vertical transport of  $CO_2$ -rich bottom waters 67 (Gruber et al. 2012; Hauri et al. 2012; Capone and Hutchins 2013). For some phytoplankton groups, this combined exposure to low-pH upwelled waters and global ocean acidification could 68 69 exceed physiological tolerances and cause cellular stress (Hurd et al. 2009; Hutchins and Fu 2017). 70 On the other hand, upwelling could supplement the amount of inorganic carbon available for 71 photosynthesis, decreasing cellular energy expenditure on carbon concentrating mechanisms 72 (CCMs), especially in phytoplankton groups with low efficiency CCMs and weak Ribulose-1,5-73 bisphosphate carboxylase/oxygenase (RuBisCO) affinities for  $CO_2$  (Giordano et al. 2005). 74 Therefore, altered temperature, nutrient concentrations, and inorganic carbon availability with a 75 projected increase in upwelling may impact marine phytoplankton community structure and 76 function (Lassiter et al. 2006; Du and Peterson 2018).

77

Extreme heat events represent the other face of climate change in this coastal upwelling system. 78 79 Climate models and meta-analyses have attributed the increased frequency, duration, and intensity 80 (e.g., maximum temperature) of marine heatwaves to anthropogenic climate change (Di Lorenzo 81 and Mantua 2016; Oliver et al. 2018; Laufkötter et al. 2020). This increased heatwave severity 82 (extreme heatwaves) also plays a role in changing conditions that drive phytoplankton community 83 structure and system productivity (Soulié et al. 2022). For instance, during the 2013-2015 "blob" heatwave event in the North Pacific Ocean, surface warming and enhanced stratification caused 84 85 weaker upwelling that reduced nutrient input and therefore overall primary productivity of the 86 system (Cavole et al. 2016; Yang et al. 2018; Peña et al. 2019). In some regions of the North 87 Pacific, this combination of warming and upwelling relaxation led to a shift in phytoplankton community composition away from diatoms, and towards a high abundance and diversity of 88 dinoflagellates (Du and Peterson 2018). Lab experiments mimicking the temperature effects of 89 90 heatwaves on natural communities have also demonstrated shifts to low diversity, dinoflagellate-91 dominated assemblages (Remy et al. 2017). Furthermore, warm water phytoplankton species can 92 invade normally colder temperate regimes during extended surface ocean warming (Ajani et al. 93 2020). Additionally, during marine heatwaves  $CO_2$  concentrations may be lower due to thermally-94 reduced gas solubility or following drawdown by phytoplankton (Murata et al. 2002; Chavez and 95 Messié 2009). Decreased inorganic carbon availability could limit rates of carbon fixation by primary producers, or favor species with strong CCMs (Giordano et al. 2005). 96

97

98 Although upwelling and marine heatwaves differentially alter the environmental conditions in the 99 California Current System that control phytoplankton community structure and function, blooms 100 of the toxic diatom *Pseudo-nitzschia australis* have been associated with both of these climatic 101 extremes. This diatom often produces domoic acid, a potent mammalian and avian neurotoxin, 102 which can be harmful to human health, marine ecosystems, and commercial fisheries (Bates et al. 103 2018). High abundances of *Pseudo-nitzschia* spp. and domoic acid have frequently been observed 104 during springtime upwelling in the California Current System (Lange et al. 1994; Trainer et al. 105 2000; Brzezinski and Washburn 2011; Schnetzer, Burton H Jones, et al. 2013; Smith et al. 2018). 106 The correlation of toxic blooms and upwelling has led HAB researchers to categorize Pseudo*nitzschia* as a cold-water genus that blooms following large, pulsed inputs of nutrients. 107

109 In contrast, a marine heatwave in 2015 induced a persistent, highly toxic *Pseudo-nitzschia* bloom 110 of unprecedented size. In the northern portion of the California Current System, several species, including P. australis, P. fraudulenta, and P. pungens thrived in these warmer, low-nutrient, low-111 112 CO<sub>2</sub> conditions, resulting in record concentrations of domoic acid in shellfish and marine 113 mammals (McCabe et al. 2016). Over longer time periods, increased concentrations of domoic 114 acid in the Northern California Current System have been correlated with the warm phases of the 115 El Niño-Southern Oscillation (ENSO) and the Pacific Decadal Oscillation (PDO; McKibben et al. 116 2017, Sandoval-Belmar et al. 2023). This indicates that in some regions heatwaves (and extreme 117 heatwaves) are another trigger of toxic Pseudo-nitzschia bloom formation, in addition to 118 upwelling. The cellular strategies that support the remarkable ability of *P. australis* to bloom under 119 these vastly different environmental conditions remain unknown. However, it should be noted 120 that other studies have found little association between heatwaves and *Pseudo-nitzschia* blooms in 121 the Southern California Current system, thus considerations of regional differences are important 122 (Barron et al. 2013; Ryan et al. 2017).

123

The physiological responses of *P. australis* to single environmental drivers have been well 124 125 documented (reviewed in Lelong et al. 2012; Trainer et al. 2012; Bates et al. 2018). For instance, 126 warming can increase the growth rate and toxicity of several species of *Pseudo-nitzschia* (Lewis 127 et al. 1993; Zhu et al. 2017). However, one study measured more domoic acid per cell in *P. seriata* 128 at 4°C, compared to cells grown at 15°C (Lundholm et al. 1994). Furthermore, in the absence of other triggering factors such as nutrient limitation, decreases in pH associated with increases in 129 130 pCO<sub>2</sub> did not themselves greatly alter the amount of total domoic acid produced by *P. multiseries* 131 and P. fraudulenta (Sun et al. 2011; Tatters et al. 2012). However, more domoic acid per Pseudo-

132 *nitzschia spp.* cell was measured in a natural phytoplankton community exposed to 800 ppm pCO<sub>2</sub>, 133 relative to 380 ppm (Tatters et al. 2018). Furthermore, growth phase plays an important role, as 134 more domoic acid per *P. australis* cell was measured at higher  $pCO_2$  concentrations in early 135 stationary phase, but not during exponential growth (Wingert and Cochlan, 2021). Lastly, 136 phosphate and silicate limitation have been implicated in increasing both dissolved and cellular 137 domoic acid (Pan et al. 1998; Sun et al. 2011; Tatters et al. 2012). However, sufficient quantities 138 of nitrogen are necessary for domoic acid biosynthesis, as the amino acid glutamate is a building 139 block in its biosynthesis (Bates et al. 1991).

140

Despite this extensive body of knowledge about the individual impacts of temperature,  $pCO_2$ , and 141 142 nutrients on *Pseudo-nitzschia* spp., little is known about the synergistic effects between multiple 143 variables that may contribute to toxic bloom formation during upwelling and marine heatwaves. 144 Previous multiple-driver studies with *Pseudo-nitzschia* spp. have demonstrated synergistic effects 145 between multiple variables that can exacerbate growth and/or toxicity (Sun et al. 2011; Tatters et 146 al. 2012; Kelly et al. 2021). This suggests that single stressor experiments cannot fully elucidate bloom dynamics in the natural environment where multiple drivers are at play; however, these 147 148 single stressor experiments are critical to understanding the mechanisms driving observed 149 responses to multiple drivers. To our knowledge, multiple driver experiments have not been used 150 to directly compare the impacts of upwelling and marine heatwaves of *P. australis* toxic bloom 151 formation.

152

153 The objective of our experiment was to examine the physiological changes that allow *P. australis*154 to thrive under these disparate conditions. We studied the impacts of simulated upwelling, marine

heatwaves, and extreme marine heatwaves on *P. australis* physiology and toxicity to gain a better understanding of how blooms may be impacted by these events. Our in-depth approach to this question involved (a) examining relevant variables (temperature, pCO<sub>2</sub>, and nutrients) clustered together into a scenario approach for a holistic understanding, and (b) single-driver experiments for a mechanistic understanding of how individual variables may be impacting the responses we see. We hypothesized that similar to what has been observed in the field, *P. australis* would be able to grow and produce toxins under both heatwave and upwelling conditions.

162

### 163 **2. Materials and Methods**

### 164 2.1 Culture maintenance

165 Pseudo-nitzschia australis (strain NWFSC 731) was isolated from Long Beach, Washington State, 166 USA on November 3, 2020. The temperature and salinity were 14°C and 27 ppt, respectively at the time of collection. In the lab, cultures were maintained in modified f/2 medium under 120 µmol 167 photons m<sup>-2</sup> s<sup>-1</sup> LED light on a 12:12 light:dark cycle. The salinity of waters off the Washington 168 169 coast can fluctuate between 19.7 and 33 ppt dependent on riverine input (Aguilar-Islas and Bruland 170 2006). The salinity of the natural seawater used to make maintenance media was 33 ppt and 171 cultures were maintained at this salinity for all experiments. To make modified f/2 medium, natural seawater was collected from the San Pedro Ocean Time Series site (33° 33' N, 118° 24' W). The 172 173 natural seawater was then filter-sterilized (0.2 µm Whatman Polycap 150 TC cartridge filter) and 174 microwaved. A Panasonic Genius Sensor microwave was used to sterilize 2L bottles of seawater 175 for approximately 16 minutes (until boiling began) at 1250 W, mixing halfway through to ensure 176 homogeneity in temperature. The salinity of the natural seawater collected from the California 177 Bight was reduced from 36 to 33 ppt by diluting with microwave sterilized Milli-Q water. Lastly,

178 the sterilized seawater was amended with nutrients (100  $\mu$ M nitrate, 100  $\mu$ M silicate, 6  $\mu$ M 179 phosphate; standard trace metals and vitamins; Guillard and Ryther, 1962; Guillard, 1975). The seawater used in this study was collected at the surface in the summer. The background nutriment 180 181 concentrations of phosphate and silicate were 0.3 µM and 1.62 µM respectively. Our values were 182 very close to the concentrations (0.2-0.4  $\mu$ M phosphate; 2-3  $\mu$ M silicate) at the surface in the study 183 of Caron et al. (2017), who found that the nitrate concentrations in the surface seawater at this site 184 were almost undetectable (<1  $\mu$ M). Although the nitrate concentration in the seawater was not 185 measured, it can be assumed to be almost zero since our other two nutrient measurements were 186 close to the values in the study of Caron et al. (2017).

187

#### 188 2.2 Experimental conditions

189 In each experiment, cultures were acclimated to the respective conditions (described below and in Tables 1 & 2) for at least eight generations before sampling. All cultures were maintained semi-190 191 continuously in 1L polycarbonate bottles with experimental base medium (sterilized natural 192 seawater + standard f/2 trace metals and vitamins; Guillard and Ryther, 1962; Guillard, 1975). 193 Culture volume was also maintained at 1L. Nitrate, silicate, and phosphate were added directly 194 into culture flasks after dilution for final concentrations specific to each treatment (Tables 1 & 2). 195 These semi-continuous culturing methods allowed *P. australis* to remain in the exponential phase 196 of growth by diluting with fresh medium every other day. This repetitive reduction in the number 197 of cells not only prevents cultures from ever reaching the stationary phase of growth, which would 198 lead to self-shading and depletion of nutrients and CO<sub>2</sub>, but also allows cultures to determine their 199 own growth rates, based on the conditions to which they are exposed (Kelly et al. 2021; Chen and 200 Gao, 2021). The percentage of the total volume replaced was dependent on the biomass increase

201 over the dilution cycle; cultures were diluted down to approximately the same initial biomass with202 each dilution (estimated with in vivo fluorescence).

203

204 In the cluster experiments, cultures were simultaneously exposed to a combination of temperature, 205  $pCO_2$ , and nutrient concentrations representative of upwelling, heatwave, and extreme heatwave 206 conditions (Table 1). Values were chosen according to nutrient concentrations measured during 207 the blob heatwave (Du et al., 2016; Gentemann et al. 2017; Gómez-Ocampo et al. 2018; Bif et al. 208 2019; Jiménez-Quiroz et al. 2019) and upwelling events (Feely et al. 2008; Schnetzer et al. 2013; 209 Siedlecki et al. 2016; Messié et al. 2017; Larkin et al. 2020) off the West Coast of the United 210 States. An additional treatment with low temperature, pCO<sub>2</sub>, and nutrient concentrations (LTCN), 211 although less environmentally relevant, was included to tease apart the potential interactive effects 212 due to temperature. In order to change the carbonate chemistry of these cluster experiments, 213 commercial pre-mixed gas was gently bubbled directly into cultures. pH measurements were used 214 during the acclimation phase to characterize the carbonate system (see Carbonate buffer system 215 measurements and Table 3).

216

For the thermal gradient response curve, a temperature gradient was established using a thermal block connected to a VWR 1180-S recirculating chiller and a VWR 1130-2S recirculating heater (Qu et al. 2022). Cultures were grown in modified f/2 replete medium (100  $\mu$ M total nitrate, 100  $\mu$ M silicate, 6  $\mu$ M phosphate; standard trace metals and vitamins; Guillard and Ryther, 1962; Guillard, 1975) in triplicate 100 mL polystyrene vials at the following temperatures: 8.4, 11.0, 12.5, 14.1, 15.2, 16.6, 17.7, 19.3, 20.2, 22.7°C. For simplification, each temperature was rounded to the nearest whole number when referenced in the text: 8, 11, 12, 14, 15, 17, 18, 19, 20, 23°C.

225 In the pCO<sub>2</sub> single-factor experiments, all cultures were grown at 19°C with modified f/2 replete 226 medium (described above). Triplicate bottles for each treatment were gently bubbled with 227 commercially pre-mixed gas at concentrations of 200 ppm (pre-industrial), 600 ppm (ocean 228 acidification), and 1040 ppm (extreme ocean acidification). The extreme ocean acidification 229 treatment only had two replicates due to an accidental culture loss the day of sampling. pH was 230 measured daily to characterize the carbonate system (see Carbonate buffer system measurements 231 and Table 3). The extreme ocean acidification treatment only had two replicates for physiological 232 measurements as the third replicate was lost during final sampling. A replicate was also removed 233 from the pre-industrial  $pCO_2$  DA quota and production measurements due to a sample processing 234 error.

235

Two-factor nutrient-temperature experiments were conducted to examine interactive effects
between temperature and nutrients, as well as the impacts of N:P ratios on growth and domoic acid
biosynthesis. Cultures were grown across a matrix of three nitrogen to phosphorus ratios (N:P=5,
10, 50), two temperatures (13 vs. 19°C), and low vs. high total nutrient concentrations (Table 2).
Cultures were diluted semi-continuously with experimental base medium (described above). Postdilution, cultures were directly spiked with nitrate, phosphate, and silicate for final concentrations
specific to each treatment (Table 2).

243

# 244 2.3 Sample collection and analysis

245 <u>2.3.1 Growth rates</u>

Cultures were sampled once fully acclimated to the experimental conditions (as determined by
steady-state growth ±10% for at least 8 generations). Fresh medium in 1L polycarbonate bottles
was inoculated with approximately 650 cells/mL and incubated under respective experimental
conditions for 48 h.

250

For the single factor temperature and pCO<sub>2</sub> experiments, specific growth rates were calculated using in vivo chlorophyll a fluorescence measurements. Fluorescence at  $T_{initial}$  and  $T_{final}$  was measured on a Turner Designs 10-AU fluorometer. For all other experiments, growth rates were calculated using extracted chlorophyll a measurements collected at  $T_{initial}$  and  $T_{final}$  (see *Pigment analysis*). Specific growth rates were calculated using the following equation:

256

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

258

where  $\mu$  is the specific growth rate (per day) and *N* is the in vivo fluorescence or chlorophyll a concentration (for temperature-only; or cluster, nutrient, and pCO<sub>2</sub> experiments, respectively) at T<sub>initial</sub> and T<sub>final</sub> (Kling et al. 2021)

262

### 263 <u>2.3.2 Elemental analysis</u>

For particulate organic carbon (POC) and nitrogen measurements, cells were filtered onto precombusted glass microfiber (GF/F) filters (500°C for 3 h) and placed in the drying oven at 60°C for at least 48 h. Filters were pelleted in foil capsules for analysis on a Costech 4010 Elemental Analyzer (Fu et al. 2007).

# 269 <u>2.3.3 Domoic acid analysis</u>

Samples for particulate domoic acid were filtered onto Supor 0.2 µm 47 mm PES filters and stored
in 15 mL centrifuge tubes at -20°C for 1-4 months (Smith et al. 2017, Harðardóttir et al. 2018,
Jennings et al. 2020). The filtrate was also collected and frozen in 100 mL amber HDPE bottles
for dissolved domoic acid analysis.

274

275 For particulate domoic acid analyses, filters were extracted for four hours in 90% methanol and 276 10% water, vigorously vortexing each hour following the methods described in Wang et al. (2012) 277 where DA recoveries exceeded 90% in cultured phytoplankton samples. Extracts were passed 278 through a 0.2 µm syringe filter directly into a 1.5 mL LC-MS vial for LC-MS/MS analysis on 279 a Prominence UFLC system (Shimadzu, Kyoto, Japan) coupled to a SCIEX 4500 QTRAP mass 280 spectrometer (AB Sciex, Framingham, MA, USA). The mass spectrometry methods followed 281 those of Sterling et al. (2022) with some minor modifications (the eluent was sent to waste during 282 the first 5 min of the run rather than 2 min and the CRM for DA was obtained from the National 283 Research Council of Canada rather than Sigma-Aldrich). The peak of DA eluted at 11.00 min. LC-284 MS/MS with multiple reaction monitoring (MRM) was employed for detection and quantification. Analysis was carried out in positive mode, and three transitions from the protonated DA molecule 285 were used: m/z 312  $\rightarrow$  266 (quantitation transition), m/z 312  $\rightarrow$  248, and m/z 312  $\rightarrow$ 286 193. Plankton-associated domoic acid was quantified to ng particulate domoic acid L<sup>-1</sup> of filtered 287 seawater using an external calibration curve created from pure domoic acid standards of increasing 288 289 concentrations (CRM, National Research Council of Canada), included in each analysis.

One DA sample was removed from analysis in the pre-industrial treatment in the CO<sub>2</sub> single factor
experiment, and from the NP=50, high nutrient, 19°C treatment in the N:P ratio experiment, due
to a sampling and/or analytical error.

294

For environmental relevance and to make inferences about trophic transfer of toxins, domoic acid was normalized to cells per L. Domoic acid was normalized also to moles of POC per liter; because *Pseudo-nitzschia spp.* cell size and volume can change with nutrient limitation, normalizing only to cell counts may result in skewed results (Tatters et al. 2012). In the text, discussion of "DA quotas" refers to this POC normalized value, unless otherwise stated.

300

301 Dissolved DA was measured from acidified filtered seawater (50 mL, 0.2 µm) passed over a C18
302 solid phase extraction column and eluted with methanol based on the methods of Wang et al. 2012
303 with some modification. The eluent was placed directly into a 1.5 mL LC-MS vial and analyzed
304 using the LC-MS method detailed above.

305

Domoic acid production rates were calculated by multiplying specific growth rates by DA quotas.
This value provides an estimate of how toxic a bloom might be, based on the ability of *Pseudo-nitzschia* to increase cell abundances and produce high DA quotas (per mol POC). For instance, if *Pseudo-nitzschia* is present in high abundances but not toxic, the bloom might not have negative
ecosystem implications (Kelly et al. 2021).

311

312 <u>2.3.4 Chlorophyll a analysis</u>

313 Samples for chlorophyll a were filtered on GF/F filters and stored in scintillation vials at -20°C for

~24-48 h. Samples were extracted in 6 mL of 90 % acetone at -20°C for 24 h, then analyzed using

a Turner 10AU field fluorometer (Welschmeyer 1994; Fu et al. 2007).

316

## 317 <u>2.3.5 Net carbon fixation and elemental use efficiencies</u>

318 Primary production was determined by measuring the uptake of radiolabeled bicarbonate (Fu et al. 2008).  $^{14}\text{C}\text{-bicarbonate}$  was added to 45 mL sub-cultures at  $T_{24\ h}$  and incubated for 24 h 319 320 (approximating net carbon fixation) under the respective experimental conditions. After the 321 incubation period, cells were collected on GF/F filters and placed in a scintillation vial containing 322 scintillation cocktail. Samples were stored for 24 h before being read on a Wallac System 1400 323 liquid scintillation counter. Carbon fixation rates were calculated by converting raw counts of 324 disintegrations per minute to µmol of carbon (based on total activity of the radiolabeled 325 bicarbonate), then normalizing to the incubation time. Dissolved inorganic carbon (DIC) values 326 used in the calculation were adjusted for each treatment based on measured values of DIC (see 327 below). These rate measurements were then normalized to POC. In the cluster experiment, one 328 sample from the extreme heatwave treatment was removed from primary production analyses due 329 to an error made during the assay.

330

Nitrogen use efficiencies (NUEs) were calculated by normalizing net carbon fixation rates to
particulate organic nitrate (mols C fixed hour<sup>-1</sup> mol N<sup>-1</sup>; Yang et al. 2022).

333

334 <u>2.3.6 Carbonate buffer system measurements</u>

335 pH measurements were made on a Mettler Toledo SevenCompact pH meter using a three-point 336 calibration curve and total pH scale (Cooley and Yager 2006). Samples for total DIC analysis were 337 collected at T<sub>final</sub>. Seawater from undisturbed culture bottles was removed with a sterile syringe, 338 ejected into pre-evacuated borosilicate Exetainers, and poisoned with 5% MgCl<sub>2</sub>. Samples were stored at 4°C until analysis. Total DIC was measured using a Picarro cavity ring-down 339 340 spectrophotometer according to Subhas et al. (2015). Experimental seawater  $pCO_2$  and total 341 alkalinity were calculated from measured DIC and pH using CO2SYS version 2.1 software (Table 342 3; Lewis and Wallace, 1998).

343

344 <u>2.3.7 Cell counts</u>

For cell count samples, 1 mL of the final experimental culture was preserved with 40 ul
glutaraldehyde and stored at 4°C in the dark. Cells were counted on a Olympus BX51 microscope
using a Sedgewick Rafter Chamber.

348

### 349 <u>2.3.8 Statistical methods</u>

Multivariate analyses were conducted in R version 1.3.1093 (http://www.r-project.org) using statistical tools in Rallfun-v38 (https://dornsife.usc.edu/labs/rwilcox/software/). For global analyses, percentile bootstrap methods for comparing 20% trimmed means were used to detect significant differences between treatments. t1way was used in the single-factor and cluster experiments, while t3way was used in the nutrient-temperature experiments. To test for significant differences between any two treatments, pairwise analyses using percentile bootstrapping of 20% trimmed means (trimpb2) were conducted. These statistical tools are analogous to ANOVAs and post-hoc and Welch's t-test pairwise analyses but are more robust as they do not assume normalityor homoscedasticity (Wilcox, 2003).

359

360 **3. Results** 

### 361 *3.1 Multiple driver cluster experiments*

Specific growth rates were greatest in the heatwave treatment, but only 17% lower in the upwelling treatment (p-value < 0.01), averaging 0.44 and 0.37 day<sup>-1</sup> (respectively; Figure 1A). In contrast, *P. australis* growth rates in the extreme heatwave treatment were 61% lower than in the heatwave treatment, although these experimental conditions only differed by  $1.5^{\circ}$ C in temperature. Furthermore, when *P. australis* was grown with lower heatwave concentrations of nutrients and pCO<sub>2</sub>, but also at a reduced temperature (i.e., low temperature, CO<sub>2</sub>, and nutrients; LTCN), growth rates were not significantly different from those measured in the extreme heatwave treatments.

369

370 Dissolved DA (dDA) concentrations were a negligible fraction of the total DA in all treatments 371 (data not shown), likely due to the constant exponential phase growth and regular culture medium 372 dilutions used in our semi-continuous incubations. POC normalized pDA quotas were highest in the upwelling treatment, averaging 0.9 ng DA  $\mu$ mol C<sup>-1</sup> (Figure 1B). In contrast, toxins were not 373 374 detected in the heatwave treatment, and pDA concentrations were 93% lower than upwelling 375 values under extreme heatwave conditions. Furthermore, pDA concentrations for the LTCN 376 treatment were 41% lower than upwelling concentrations. Trends in DA quotas were the same 377 when normalized to cell abundance and to volume (mL), with the greatest quotas measured in the 378 upwelling treatment Table 4). DA production rates demonstrated a similar trend, with the highest production rates measured under upwelling conditions (0.33 ng DA µmolC<sup>-1</sup> day<sup>-1</sup>) and zero or 379

near zero for the heatwave and extreme heatwave treatments (Figure 1C). DA production rates in
the LTCN treatment were 92% higher than in the extreme heatwave treatment (p < 0.01).</li>

382

Net C-specific primary productivity trends were similar to those observed for growth rates (Figure 2A). The highest rates were measured in upwelling and heatwave treatments (0.012 and 0.013 day<sup>-</sup> 1, respectively), which were not significantly different from one another (p = 0.221). Rates for the LTCN and extreme heatwave conditions were 54% lower that the heatwave treatment (p < 0.01).

387

Nitrogen use efficiencies (NUEs) measure how efficiently the cells use their cellular nitrogen quotas to fix carbon under different experimental conditions. NUEs were highest under heatwave conditions and decreased by 17% in the upwelling treatment (p = 0.028 Figure 2B). For the LTCN treatment, NUEs were slightly lower than those for the upwelling treatment (p < 0.01). Finally, NUEs for the extreme heatwave treatment declined 61% relative to the heatwave treatments (p < 0.01).

394

### 395 *3.2 Single factor experiments*

When temperature alone was considered, the fastest specific growth rates were measured between 14 and 19°C, reaching a maximum of 0.35 day<sup>-1</sup> at 18°C (t1way p-value <0.01; Figure 3A). At temperatures above 19°C or below 13°C, growth rates declined rapidly. Furthermore, *P. australis* grew extremely slowly (0.02 day<sup>-1</sup>) and was unable to persist stably at temperatures  $\leq$ 9°C and could not survive when temperatures exceeded 23°C.

Particulate domoic acid cellular quotas (pDA, ng DA µmol C<sup>-1</sup>) had an inverse relationship with 402 403 temperature, whereby pDA decreased as temperature increased, except at 8°C where cold-stressed cultures did not produce any detectable toxin (Figure 3B). The  $R^2$  value was 0.88 for a linear 404 405 regression that excluded the 8°C values, indicating a strong linear relationship between temperature and pDA. The highest concentrations of pDA were measured at 11°C, averaging 3.52 406 ng DA µmol C<sup>-1</sup>. Between 13 and 18°C, pDA quotas were slightly lower than the maximum, 407 408 though similar to each other. Relative to the maximum, pDA declined by 72% with warming to 409 19°C and 20°C, and no pDA was measured at 23°C. DA production rates (ng DA µmol C<sup>-1</sup> d<sup>-1</sup>) 410 were highest and not significantly different from one another between 11 and 18°C, averaging 0.68 ng DA µmolC<sup>-1</sup> d<sup>-1</sup> and decreased by 62% to 19°C. At 8 and 23°C, DA production rates were at or 411 near zero (Figure 3C). 412

413

414 In the single factor  $CO_2$  experiment, the highest growth rates were measured in the 600 µatm 415 treatment (Figure 4A). Growth rates were 23% and 11% lower in the 200 and 1040 µatm 416 treatments, although this difference was only significant for the 200 µatm treatment (pairwise p-417 value < 0.01). On the other hand, cellular DA content was greatest at the extreme ends of the CO<sub>2</sub> 418 concentration gradient (Figure 4B). In the 600 µatm treatment, DA quotas were 66% and 56% 419 lower relative to 200 µatm and 1040 µatm (respectively, p-values <0.01 and 0.021). Lastly, DA 420 production rates followed a trend similar to DA quotas; the lowest rates were observed at 600 421 µatm, with a 54% and 50% increase compared to 200 and 1040 µatm (although differences were only statistically significant between 200 and 600  $\mu$ atm; p < 0.01). 422

423

424 3.3 N:P ratio experiments

Within each combination of temperature and total nutrient concentration (i.e., low vs. high nutrients), growth rates were greatest for the treatments with an N:P ratio of 50 (Figure 5A). Within each temperature for the N:P=5 and N:P=10 treatments, differences in growth rates between high and low total nutrient treatments were very minor, yet statistically significant (t3way N:P ratio affects p-value = 0.0001).

430

There were no other significant 2- or 3-way interactive effects between variables for growth rates
in this experiment. Furthermore, growth rates increased with both temperature and higher total
nutrient concentrations (high nutrient treatments), except for the 19°C high nutrient N:P=50
treatment.

435

436 Particulate domoic acid was not detected in the N:P=5 treatments. The most pDA was produced in the N:P=50 treatments, which were 71 to 90% higher than the N:P=10 treatments (t3way single 437 438 variable N:P ratio effects p-value = 0.0001; Figure 5B). Furthermore, within the N:P=50 treatments 439 lower concentrations of pDA were measured in the treatments with lower total nutrient concentrations (t3way interactive effects between N:P ratio and total nutrient concentration p-440 441 value = 0.014). Differences between N:P=10 treatments were minor at  $13^{\circ}$ C, yet at  $19^{\circ}$ C the high 442 total nutrient concentration treatment was 96% higher than the treatment with a low total nutrient 443 concentration (t3way interactive effects between all variables p-value = 0.046).

444

445 **4. Discussion** 

446 <u>4.1 Single factor experiments and potential interactions reveal the influence of each environmental</u>

447 <u>variable the cluster experiment</u>

The single and dual factor experiments provided a mechanistic understanding of how the drivers examined contributed to the responses in the cluster experiment. We found that the trends observed in the single-factor treatments could not fully explain the results of the multiple-driver experiments. This provides evidence that non-linear, interactive effects may be influencing trends observed in the multivariate scenario experiments.

453

### 454 *4.1.1 Growth rates*

455 Other studies have shown that optimal growth temperatures are highly species and strain dependent 456 (Lelong et al. 2012; Bates et al. 2018). The heatwave temperature was within the broad optimal 457 temperature range for growth for this P. australis strain, while the upwelling temperature was 458 suboptimal. This aligns with results from the cluster experiment, where growth rates for the 459 heatwave treatment were slightly higher than those in the upwelling cluster, indicating that 460 temperature did influence growth rates in the cluster experiment. However, temperature was only 461 partially responsible for the decreased growth rates in the extreme heatwave treatment, as growth 462 rates declined more than expected from temperature alone.

463

On the other hand, pCO<sub>2</sub> alone had only a slight impact on *P. australis* growth rates. However, inorganic carbon concentration did not influence growth rates in the cluster experiments as expected based on the single factor experiment; growth increased slightly with pCO<sub>2</sub> concentration alone, yet growth was higher in the heatwave treatment with less pCO<sub>2</sub>. Similar to the trends observed in the present single-factor study with *P. australis*, increases in growth rates have been observed from 200 to 765 ppm for *P. multiseries* and from 220 to 730 ppm for *P. fraudulenta*. However, when *P. multiseries* and *P. fraudulenta* were both exposed to pCO<sub>2</sub> and limited for either 471 phosphate or silicate, growth rates further declined, suggesting that there were interactive effects 472 between nutrient limitation and pCO<sub>2</sub> concentration (Sun et al. 2011; Tatters et al. 2012). Zhu et 473 al. (2017) also demonstrated that there were interactive effects of pCO<sub>2</sub> and temperature on *P*. 474 *subcurvata*; growth rates under suboptimal temperature and high pCO<sub>2</sub> conditions were lower than 475 growth rates at optimal temperatures combined with low pCO<sub>2</sub>.

476

477 Single-factor nutrient experiments were not conducted in the present study, as individual nutrient 478 effects on *Pseudo-nitzschia* growth have been studied extensively. As with many other diatom 479 species, nutrient limitation typically reduces Pseudo-nitzschia growth (Fehling et al. 2004; Hagström et al. 2011; Sun et al. 2011; Tatters et al. 2012; Auro and Cochlan 2013). In the nutrient-480 481 temperature matrix experiment, growth rates were higher for treatments with higher total nutrient 482 concentrations. However, in the cluster experiments *P. australis* growth rates were slightly higher 483 in the heatwave treatment, despite having lower nutrient concentrations relative to the upwelling 484 treatment. Furthermore, regardless of equal nutrient concentrations in both heatwave treatments, 485 growth rates in the extreme heatwave treatment were lower. This indicates that there was a temperature-nutrient interaction in the cluster experiments (see below). 486

487

### 488 4.1.2 Domoic acid quotas

Temperature alone had a significant impact on DA quotas in the present study. In contrast to our observation that DA quotas decreased with warming, most published *Pseudo-nitzschia* studies on temperature have observed increases in DA with warming (Lelong, Hégaret, et al. 2012; Thorel et al. 2014; Zhu et al. 2017; Kelly et al. 2021); however, one found that *P. seriata* produced more DA at relatively colder temperatures, similar to observations in the present study (Lundholm et al.

494 1994). Furthermore, *P. australis* has been associated with relatively cold temperatures in 495 Narraganset Bay, RI (Roche et al. 2022). In March 2017, a highly toxic *P. australis* bloom occurred 496 at temperatures ranging from 1 to 5°C in Narraganset Bay, RI (Sterling et al. 2022). The diversity 497 of thermal tolerances documented in the literature demonstrate that different species and strains of 498 *Pseudo-nitzschia* have distinct thermal triggers for toxin production, though the data in the present 499 study indicates that this strain of *P. australis* specializes in being more toxic at colder temperatures.

500

501 Because temperature alone (under nutrient replete conditions) had a strong inverse relationship 502 with DA concentrations in the single factor experiments, it likely contributed to the differences in DA between cluster treatments where more DA was measured in the cold, upwelling treatment 503 504 compared to the warm, heatwave treatments. However, in the heatwave and extreme heatwave 505 treatments the decrease in DA quotas were greater than expected based on the temperature single 506 factor experiments. Furthermore, it should be noted that in the upwelling treatment and 507 temperature experiment, nitrogen availability was replete and ample to support DA biosynthesis, 508 which may not have been the case in the heatwave and extreme heatwave cluster treatments (the 509 interactive effects are discussed more below).

510

Similar to our observation that more DA was measured at the extreme ends of the pCO<sub>2</sub> concentration gradient, another strain of *P. australis* was highly toxic at low and extremely high pCO<sub>2</sub> concentrations (371 and 1849  $\mu$ atm), and only moderately toxic at an intermediate pCO<sub>2</sub> concentration (785  $\mu$ atm; Ayache et al. 2021). In contrast, other strains of *P. australis* have demonstrated decreases in DA along the same pCO<sub>2</sub> gradient described above, or no change in DA across pCO<sub>2</sub> gradients during the exponential phase of growth (Sun et al. 2011; Tatters et al. 2012; 517 Ayache et al. 2021; Wingert and Cochlan 2021). While there are species and strain differences 518 with regard to  $pCO_2$  concentration, toxicity of the *P. australis* strain used in the present study is 519 more sensitive to changes in  $CO_2$  alone.

520

Inorganic carbon may have influenced the upwelling cluster DA quotas, as the extreme ocean acidification  $pCO_2$  concentration in the single factor experiment was similar to the concentration used in the upwelling experiment. However, it is likely that  $pCO_2$  alone did not trigger DA biosynthesis in the heatwave and extreme heatwave cluster treatments. Although the pre-industrial  $pCO_2$  level, similar to levels used in the heatwave and extreme heatwave treatments, did trigger DA biosynthesis, very little to no DA was observed in these clusters.

527

528 Previous investigations have robustly demonstrated that phosphate and/or silicate limitation trigger 529 DA biosynthesis in toxic strains of *Pseudo-nitzschia* sp. (Pan et al. 1996; Fehling et al. 2004; Sun 530 et al. 2011; Tatters et al. 2012). This suggests that silicate and phosphate limited conditions should 531 have triggered DA biosynthesis in the heatwave and extreme heatwave treatments, yet little to no 532 DA was measured. However, these treatments were also nitrate limited, and sufficient nitrogen is 533 required for synthesis of DA, a nitrogen-containing molecule (Lelong et al. 2012). Therefore, the 534 total amount of nitrogen in these treatments may have strongly negatively impacted DA quotas 535 (see more below).

536

537 <u>4.2 Mechanistic studies and potential interactive effects inform dynamics in the cluster experiment</u>
 538 The three drivers interacted in the cluster experiment to alter growth rates more than expected from
 539 individual variables or additive effects. We hypothesize that there was a tradeoff between

temperature, nutrients, and carbon dioxide: at cooler temperatures, high nutrients and inorganic
carbon offset the growth limitation of low temperature. On the other hand, warming supported
higher growth rates, even if limited by nutrients and inorganic carbon. This indicates that there
was a synergistic interactive effect between these three variables.

544

545 In terms of bloom toxicity, temperature and inorganic carbon may have partially contributed to the 546 increased DA measured in the upwelling cluster. However, it is likely that the main driver of 547 differences in toxicity between treatments was nitrogen. Both the heatwave treatments were 548 supplied with less nitrogen, while the upwelling treatment had ample nitrate, which is necessary 549 for the biosynthesis of DA molecules (Lelong et al. 2012). Therefore, it is possible that the lack of 550 nitrogen in the heatwave and extreme heatwave treatments had an antagonistic effect, inhibiting 551 DA biosynthesis regardless of the  $CO_2$  conditions that may trigger toxin production. Therefore, 552 we surmise that nitrate concentration was a key factor influencing DA production in the cluster 553 experiments.

554

555 DA is a specialized metabolite and biosynthesis is an energetically expensive process. Therefore, 556 DA is typically produced when there is sufficient energy for both processes, or when silicate or 557 phosphate availability limit or reduce cell division (Pan et al. 1998). However, this tradeoff does 558 not always seem to be the case in the present study. In the temperature experiment, slower growth 559 rates and high DA quotas at 11°C were associated with high DA production rates, while higher 560 growth rates at slightly warmer temperatures were associated with lower DA quotas yet near equal 561 DA production rates. For the pCO<sub>2</sub> experiments, differences in growth rates were marginal and 562 DA production rates closely followed DA quota trends. In this case, growth rates were decoupled

from bloom toxicity. Together, these experiments demonstrate that high biomass does not
necessarily indicate high toxicity, and low biomass does not indicate a low threat of DA poisoning.
Similarly, observations of high *Pseudo-nitzschia* sp. biomass yet low DA concentrations have been
detected in the natural environment (Roche et al., 2022).

567

568 DA concentrations in the cluster treatments can also be examined through the lens of cellular 569 energetics and the tradeoff between growth and toxicity. In addition to the need for sufficient 570 cellular energy, nitrogen is also required for both growth and DA synthesis. This tradeoff between 571 growth and toxicity is not evident in the cluster experiments: upwelling and heatwave treatments 572 grew equally well yet differed in terms of toxicity. Instead, this may be explained by the total 573 amount of nitrogen available in each treatment.

574

575 Considering growth rate results, DA measurements, and previous studies with nitrogen, we 576 hypothesize that the upwelling treatment had more than enough nitrogen for growth, and the excess 577 was shunted to DA biosynthesis. P. australis cells in the heatwave and extreme heatwave 578 treatments had less total nitrogen available. In the heatwave treatment, nitrogen did not limit 579 growth, carbon fixation, or NUEs, but there might not have been enough for concurrent DA 580 synthesis. This indicates that the absolute amount of nitrogen plays a role in the tradeoff between 581 growth and toxicity. Additionally, in the extreme heatwave treatment, both growth rates and DA 582 concentrations were reduced. Previous studies have observed declines in growth rates due to 583 warming, concurrent with increases in toxicity, indicating that the energy and nitrogen not used 584 for growth was instead shunted to DA biosynthesis (Zhu et al. 2017). However, for this strain of *P. australis* the low DA quotas in the extreme heatwave treatment may be due to a lack of sufficient
nitrogen and a warming-induced reduction in DA biosynthesis.

587

588 The LTCN cluster treatment, though not environmentally realistic, further aids in teasing apart the 589 nutrient-temperature interactive effects on DA quotas. These LTCN cells were more toxic than the 590 heatwave and extreme heatwave treatments, though not as toxic as the upwelling treatment. Low 591 temperature and  $pCO_2$  are triggers of DA for this strain of *P. australis*, which could partially 592 explain the increased toxicity relative to the heatwave treatments. However, growth rates for the 593 LTCN treatment were lower. Therefore, if all of the available nitrogen was not used for growth in 594 this treatment, there may have been enough left over for DA synthesis. Furthermore, both the 595 upwelling and LTCN treatments efficiently used the available nitrate to support carbon fixation, 596 as NUEs were similar. This indicates that temperature does indeed affect the ability of *P. australis* 597 to use nutrients and synthesize DA.

598

599 Results from the N:P ratio experiments also demonstrate this temperature-nutrient interaction. 600 While growth rates changed minimally across the N:P=50 treatments, there were significant 601 differences in DA quotas within each temperature. This indicates a decoupling between growth 602 rates and DA biosynthesis. Furthermore, when total nutrient concentrations were low, warming 603 constrained DA quotas, yet under high total nutrient concentrations differences in DA quotas were 604 not considerable. This suggests that temperature is more influential on DA quotas when total 605 nutrient concentrations are high. Consequently, under these high total nutrient conditions with 606 excess nitrogen (relative to phosphorus), cells can both double their population size approximately 607 once per day and produce substantial DA, generating a large toxic bloom.

## 609 <u>4.3 Cluster experiments: a holistic view and ecosystem implications</u>

610 Examining the data empirically using cluster experiments can provide a holistic view of how P. 611 australis bloom formation and toxicity may be impacted by complex events like upwelling and 612 marine heatwaves. In the present study's simulated upwelling and heatwave conditions, similarly 613 high growth rates for both treatments indicate that both conditions could be triggers for bloom 614 formation. These results are consistent with what has been observed in the natural environment: 615 blooms occurring during both upwelling and marine heatwave events (Schnetzer et al. 2013; 616 McCabe et al. 2016; McKibben et al. 2017; Smith et al. 2018). With more frequent heatwave and 617 upwelling events expected as climate change progresses, rapid bloom-forming growth may be 618 triggered more frequently (Bakun et al. 2015). However, if extreme heatwaves exceed the broad 619 thermal optimum for *P. australis* growth, climate change may impede bloom formation. Therefore, 620 the severity of future heatwaves will determine whether *P. australis* will be able to rapidly grow.

621

622 Despite similar growth rates between upwelling and heatwave treatments, only the upwelling 623 conditions led to substantial DA biosynthesis. This suggests that blooms triggered by upwelling 624 may be more toxic than blooms occurring during a heatwave or extreme heatwave event. 625 Furthermore, DA production rates were only substantial in the upwelling treatment. This parameter 626 combines growth rates and toxin quotas to estimate the impact of the bloom (Kelly et al. 2021). A 627 high DA production rate suggests that upwelling events may promote both rapid increases in 628 population size and high concentrations of toxins per cell. Therefore, *P. australis* may be able to 629 form a large, toxic bloom rapidly under these conditions, supporting predictions that blooms will 630 worsen with climate change (Fu et al. 2012; Smith et al. 2018; Gobler 2020; Trainer et al. 2020).

631 Consequentially, blooms triggered by upwelling may be especially harmful to the marine 632 ecosystem, with severe implications for the coastal California system. More frequent, highly toxic 633 blooms could have devastating consequences for ecosystem health and fisheries (McCabe et al. 634 2016). However, in the present experiment pDA quotas normalized to volume were very low 635 compared to maximum concentrations of pDA observed during the Blob bloom (McCabe et al. 636 2016). These values are also well below the regulatory limit for DA (20 ppm). Additionally, DA 637 quotas normalized to cell abundance were a few orders of magnitude lower than a highly toxic P. 638 australis bloom triggered by upwelling in 2006 (Schnetzer et al., 2013). Therefore, the risk of 639 trophic transfer of domoic acid may not be high for this strain of *P. australis* under upwelling 640 conditions.

641

642 In contrast, although cells in the heatwave treatment grew faster, DA was not detected and thus 643 DA production rates were inconsequential. P. australis in the extreme heatwave treatment had 644 neither high growth rates nor high DA quotas. This indicates that despite the ability of *P. australis* 645 to multiply rapidly and form a bloom, toxin production may not be triggered by heatwave 646 conditions. The rapid growth of *Pseudo-nitzschia* spp. under heatwave conditions was also 647 observed during the Blob bloom, with cells persisting in the warm waters, reducing nutrient 648 concentrations in the surface layer of these stratified waters (McCabe et al. 2016). Similar to our 649 experiment, the large population of cells in the northern region of the California Current System 650 did not produce toxins without the availability of nitrogen. However, upwelling occurred in the 651 spring of 2015, fueling the eastern edge of the Blob with nutrients, including nitrogen, which likely 652 allowed the Pseudo-nitzschia spp. present to produce DA (McCabe et al. 2016). These seed 653 populations of highly abundant and persistent *Pseudo-nitzschia* sp. populations, triggered by

warming, pose a looming threat ecosystem health, as their toxicity may be triggered by episodic
nitrogen inputs. Though our study did not mimic both the heatwave and upwelling aspects of the
Blob bloom, future studies should consider heatwave conditions along with a pulse of nutrients.

657

658 In contrast, Ryan et al. (2017) found that in Monterey Bay, the high Pseudo-nitzschia cell 659 abundance and toxicity was not due the warming anomaly (as temperatures were near normal), but 660 instead correlated with the cold water, upwelling phases. Similarly, Barron et al. (2013) found no 661 correlation between *P. australis* presence in the sediment record and warm anomalies in the Santa 662 Barbara Channel. These studies in the Southern Portion of the California Current System contrast 663 studies that occurred in the Northern portion of the California Current System (McCabe et al. 2016, 664 McKibben et al. 2017). Furthermore, Sandoval-Belmar et al. (2023) found that toxic Pseudo-665 nitzschia blooms in Northern California have been associated with the warm phases of ENSO and 666 PDO, while blooms in Southern California were associated with the cool phases. The strain of P. 667 australis used in the present study was isolated from the Northern California Current System 668 (Washington State) and thrived under heatwave conditions. This indicates that regional differences 669 exist regarding the ability of heatwaves to trigger bloom formation.

670

On the other hand, although heatwaves may trigger a lot of growth, those cells are not expected to be notably toxic without nitrogen input from simultaneous upwelling or an anthropogenic source. While high nutrient concentrations occur in the natural environment during upwelling, dissolved N:P ratios of this water are typically slightly below the Redfield Ratio (N:P=  $\sim$ 12; Feely et al. 2008; Schnetzer et al. 2013; Siedlecki et al. 2016; Larkin et al. 2020). However, increased flow of anthropogenic nitrogen from land to sea (fertilizers and wastewater treatment facility discharge) 677 may increase the N:P ratio of coastal surface waters (Howard et al. 2014). Increased growth and 678 DA quotas in the N:P=50 treatments of the temperature-nutrient matrix experiment suggest that upwelling combined with anthropogenic nutrient inputs might trigger especially toxic P. australis 679 680 blooms, more than upwelling alone. If anthropogenic nutrient inputs occurred during a heatwave, 681 a highly toxic bloom could be triggered due to the excess nitrogen, relative to phosphorus. 682 Furthermore, considerable species and strain specificity exists with regard to the ability of *Pseudo*-683 nitzschia to utilize different nitrogen sources for growth and toxin production (Howard et al. 2007; 684 Kudela et al. 2008; Thessen et al. 2009). Therefore, it is possible that DA biosynthesis for this 685 strain of P. australis is not optimized for nitrate, and other sources of nitrogen (e.g., urea and 686 ammonia) should be tested with this strain in the future to further explore the implications of 687 anthropogenic nutrient inputs and their potential interactions with bloom formation and toxicity.

688

### 689 **5.** Conclusions

690 This study is among the few to examine unialgal *Pseudo-nitzschia* cultures in a multiple driver 691 context, and the first to examine both upwelling and heatwave scenarios in holistic laboratory 692 cluster experiments. These experiments are important for improving our understanding of P. 693 *australis* specific responses to these conditions in order to make better predictions about future 694 bloom dynamics in the California Current System. Furthermore, the single factor experiments here 695 provided a unique mechanistic understanding of specific triggers of toxicity and bloom formation 696 for this strain of *P. australis*. This has the potential to improve our ability to predict the occurrence of toxic blooms in the natural environment. 697

699 This experiment only examined one strain of *Pseudo-nitzschia australis*, and these responses may 700 be strain specific. Similar experiments should be conducted with different species and strains isolated from different areas in the California Current System. Furthermore, while we chose to 701 702 focus on only temperature, nutrients, and pCO<sub>2</sub>, the clusters are not truly complete without other 703 factors that have been shown to impact domoic acid synthesis, including biological factors like 704 grazing, competition with other phytoplankton, and the presence of associated bacteria and fungi. 705 Other physical and chemical factors like light intensity and quality, iron concentration, and 706 alternative nitrogen sources are also clearly important (Howard et al. 2007; Thessen et al. 2009; 707 Lelong et al. 2012; Bates et al. 2018; Radan and Cochlan 2018). Considering these factors as well 708 could make these experiments more realistic. Nevertheless, these cluster experiments provide a 709 valuable start towards obtaining a more holistic picture of *P. australis* dynamics during these 710 contrasting events in the future rapidly changing coastal ocean.

711

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725	
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727	F-XF, KJK, and DAH conceived of and designed the experiments with help from BDJ and MJB.
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730	quantification. Data analyses were performed by KK, with help from F-XF and DAH. KK wrote
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733	
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**Figure 1.** Average growth rates (a), particulate domoic acid quotas (b), and domoic acid production rates for scenario experiments with combined CO<sub>2</sub>, temperature, and nutrients representative of upwelling, heatwaves, extreme heatwave, and low temperature, CO<sub>2</sub>, and nutrients (LTCN) conditions. Different letters indicate that differences between treatments are statistically significant, while the same letters indicate that differences between treatments are not statistically significant. Error bars represent standard deviation of the mean.



**Figure 2.** Net primary productivity (A), nitrogen use efficiency (NUE; B) of *P. australis* under upwelling, LTCN, heatwave, and extreme heatwave conditions. Different letters indicate that differences between treatments are statistically significant, while the same letters indicate that differences between treatments are not statistically significant. Error bars represent standard deviation of the mean.



**Figure 3.** N Average *P. australis* growth rates (A), particulate domoic acid (pDA) cellular quotas (ng DA  $\mu$ mol C<sup>-1</sup>) (B), and domoic acid production rates (ng DA  $\mu$ mol C<sup>-1</sup> d<sup>-1</sup>) (C) across a range of temperatures. Different letters indicate different statistically significant differences between temperatures, and error bars represent standard deviations of the mean (n=3).



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1008 **Figure 4.** Growth rates (A), particulate domoic acid concentrations (B), and domoic acid 1009 production rates (C) measured in the single factor pCO<sub>2</sub> experiments. Different letters indicate 1010 statistically significant differences between temperatures, and error bars represent standard 1011 deviations of the mean.



1013 Figure 5. Specific growth rates for all N:P ratio experiments (A) and particulate domoic acid 1014 quotas for the N:P=10 and N:P=50 experiments (B). Domoic acid was not detected in the N:P=5 1015 experiments. Different letters indicate different statistically significant differences between 1016 temperatures, and error bars represent standard deviations of the mean.

Treatment	Upwelling	LTCN	Heatwave	Extreme heatwave
Temperature	13°C	13°C	19°C	20.5°C
pCO <sub>2</sub>	900 µatm	230 µatm	255 µatm	240 µatm
Nutrients	3 μM PO <sub>4</sub> 40 μM Si(OH) <sub>4</sub> 30 μM NO <sub>3</sub>	0.5 μM PO <sub>4</sub> 10 μM Si(OH) <sub>4</sub> 5 μM NO <sub>3</sub>	0.5 μΜ ΡΟ4 10 μΜ Si(OH)4 5 μΜ NO3	0.5 μM PO4 10 μM Si(OH)4 5 μM NO3

1018 Table 1: Levels of treatment for cluster experiments, with altered temperature,  $pCO_2$ , and 1019 <u>nutrients<sup>1</sup></u>.

1020  $\overline{}^{1}$  LTCN = low temperature, carbon dioxide, and nutrients.

N:P ratio	Total nutrient concentration	Nitrate (µM)	Phosphate (µM)	Silicate (µM)
5	Low	5	1	10
	High	30	6	40
10	Low	5	0.5	10
10	High	30	3	40
50	Low	5	0.1	10
	High	30	0.6	40

Table 2: Experimental treatments in the nutrient-temperature matrix experiment. Each treatment
below was grown under two temperature conditions: 13 and 19°C.

		Measured pH	Measured DIC (µmol/kg)	Calculated bulk alkalinity (µmol/kg)	Calculated pCO <sub>2</sub> (µatm)
Cluster	Upwelling	7.7 (0.01)	2098.0 (17.8)	32.9 (0.7)	889.1 (29.0)
experiments	LTCN	8.2 (0.01)	1868 (16.9)	90.1 (0.8)	229.3 (4.6)
	Heatwave	8.24 (0.02)	1785 (4.6)	93.5 (2.5)	254.3 (9.4)
	Extreme heatwave	8.24 (0.04)	1787.1 (24.8)	95.2 (3.9)	240.2 (23.2)
Single-	<b>Pre-industrial</b>	8.29 (0.08)	1884 (23.2)	117.7 (14.6)	201.8 (36.5)
factor experiments	Ocean acidification	7.89 (0.01)	2058 (13.1)	53.5 (1.1)	592.2 (14.3)
	Extreme ocean acidification	7.68 (0.01)	2166 (26.2)	34.4 (0.5)	1038.9 (4.7)

Table 3: Calculated carbonate buffer system based on pH and DIC measurements. Values in
 parentheses represent the standard deviation of the mean<sup>2</sup>.

 $^{2}$ LTCN = low temperature, carbon dioxide, and nutrients.

1028	Table 4: Average domoic acid (DA) quotas for the cluster experiment, normalized to cell
1029	abundance.

Treatment	Average ng DA/cell	Average ng DA/mL
	(standard deviation)	(standard deviation)
Upwelling	7 x 10 <sup>-5</sup> (4.3 x 10 <sup>-5</sup> )	0.07 (0.01)
Heatwave	0 (0)	0 (0)
Extreme heatwave	6.9 x 10 <sup>-6</sup> (1.2 x 10 <sup>-5</sup> )	0.01 (0.01)
LTCN	2.61 x 10 <sup>-5</sup> (8.65 x 10 <sup>-6</sup> )	0.04 (0.04)