

1 **Shifts in dimethylated sulfur concentrations and microbiome composition in the red-**
2 **tide causing dinoflagellate *Alexandrium minutum* during a simulated marine heat wave**

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13
14
15 **Abstract**

16
17 The biogenic sulfur compounds dimethyl sulfide (DMS), dimethyl sulfoniopropionate (DMSP)
18 and dimethyl sulfoxide (DMSO) are produced and transformed by diverse populations of
19 marine microorganisms and have substantial physiological, ecological and biogeochemical
20 importance spanning organism to global scales. Understanding the production and
21 transformation dynamics of these compounds under shifting environmental conditions is
22 important for predicting their roles in a changing ocean. Here, we report the physiological and
23 biochemical response of a robust strain of *Alexandrium minutum*, a dinoflagellate with the
24 highest reported intracellular DMSP content, exposed to a 6-day increase in temperature
25 mimicking mild and extreme coastal marine heatwave conditions (+ 4°C and + 12°C). Under
26 mild temperature increases (+ 4°C), *A. minutum* growth was enhanced, with no measurable
27 physiological stress response. However, under a very acute increase in temperature (+ 12°C)
28 triggering thermal stress, *A. minutum* growth declined, photosynthetic efficiency (F_v/F_M) was
29 impaired, and enhanced oxidative stress was observed. These physiological responses
30 indicative of thermal stress were accompanied by increased DMS and DMSO concentrations
31 followed by decreased DMSP concentrations. At this temperature extreme, we observed a
32 cascading stress response in *A. minutum*, which was initiated 6h after the start of the experiment
33 by a spike in DMS and DMSO concentrations and a rapid decrease in F_v/F_M . This was followed
34 by an increase in reactive oxygen species (ROS) and an abrupt decline in DMS and DMSO on
35 day 2 of the experiment. A subsequent decrease in DMSP coupled with a decline in the growth
36 rate of both *A. minutum* and its associated total bacterial assemblage coincided with a shift in
37 the composition of the *A. minutum* microbiome. Specifically, an increase in the relative
38 abundance of OTUs matching the genus *Oceanicaulis* (17.0%), *Phycisphaeraceae* SM1A02
39 (8.8%) and *Balneola* (4.9%) as well as a decreased relative abundance of *Maribacter* (24.4%),

40 *Marinoscillum* (4.7%) and *Seohaecicola* (2.7%), were primarily responsible for differences in
41 microbiome structure observed between temperature treatments. These shifts in microbiome
42 structure are likely to have been driven by either temperature itself, the changing physiological
43 state of *A. minutum* cells, shifts in biogenic sulfur concentrations, the presence of other solutes,
44 or a combination of all. Nevertheless, we suggest that these results point to the significant effect
45 of extreme heatwaves on the physiology, growth and microbiome composition of the red-tide
46 causing dinoflagellate *A. minutum*, as well as potential implications for biogenic sulfur cycling
47 processes and marine DMS emissions.

48

49 **Keywords:** DMS, DMSP, DMSO, oxidative stress, thermal stress, marine heatwaves

50 **1. Introduction**

51

52

53 Many marine phytoplankton produce the organic sulfur compound dimethyl
54 sulfoniopropionate (DMSP) (Zhou et al., 2009; Berdalet et al., 2011; Caruana and Malin, 2014),
55 for which it can function as an antioxidant, osmolyte, chemoattractant and currency in
56 reciprocal chemical exchanges with heterotrophic bacteria (Stefels, 2000; Sunda et al., 2002;
57 Kiene et al., 2000; Seymour et al., 2010). Phytoplankton-derived DMSP is in fact a major source
58 of sulfur and carbon for marine heterotrophic bacteria (Kiene et al., 2000), which in turn play
59 a major role in the cycling and turnover of organosulfur compounds in the ocean (Todd et al.,
60 2007; Curson et al., 2011). The subsequent cycling of DMSP into other biogenic sulfur
61 molecules including dimethyl sulfide (DMS) and dimethyl sulfoxide (DMSO) by a suite of
62 microbial transformation pathways (Kiene et al., 2000; Sunda et al., 2002) and physical drivers
63 (Brimblecombe and Shooter, 1986) have important ecological and biogeochemical
64 implications spanning from cellular to global scales (Sunda et al., 2002; Charlson et al.,
65 1987; DeBose et al., 2008; Van Alstyne et al., 2001; Knight, 2012; Nevitt et al., 1995).

66

67 Among DMSP-producing phytoplankton, the dinoflagellate *Alexandrium minutum*, has the
68 highest recorded DMSP cell content, with an average concentration of 14.2 pmol cell⁻¹,
69 compared with less than 1 pmol cell⁻¹ in most other dinoflagellates (Caruana and Malin, 2014).
70 Blooms of *A. minutum* occur from the Mediterranean Sea to the South Pacific coast in sea
71 surface waters within temperature ranges of 12°C to 25°C (Laabir et al., 2011). Notably, some
72 strains of *Alexandrium*, including *A. minutum*, produce saxitoxins, which lead to paralytic
73 shellfish poisoning (PSP) and are responsible for the most harmful algal blooms in terms of
74 magnitude, distribution and consequences on human health (Anderson et al., 2012).

75

76 *A. minutum* commonly inhabits shallow coastal and estuarine waters (Anderson, 1998), which
77 are globally experiencing substantial shifts in environmental conditions, including increases in
78 sea surface temperature (SST) associated with climate change (Harley et al., 2006). Although
79 generally less studied than chronic temperature rises associated with global climate change
80 (Frölicher and Laufkötter, 2018), acute ephemeral temperature increases known as marine
81 heatwaves (MHWs) (Hobday et al., 2016) have recently been demonstrated to be becoming
82 more frequent and persistent as a consequence of climate change (Oliver et al., 2018). Increases
83 in MHW occurrence are anticipated to become particularly frequent within the shallow coastal

84 and estuarine waters, where *A. minutum* blooms occur (Ummenhofer and Meehl,
85 2017;Anderson, 1998).

86

87 Coastal MHW events have recently had dramatic impacts on coastal environments. MHWs of
88 up to 6°C increase in temperature in Western Australian (2011) and the Northeast Pacific
89 (2013-2015) resulted in significant ecosystem shifts with increases in novel species at the
90 expenses of others (Frölicher and Laufkötter, 2018). The 2016 MHW that was associated with
91 El Niño Southern Oscillations resulted in an 8°C increase in sea surface temperature leading to
92 the mass coral bleaching of more than 90% of the Great Barrier Reef (Hughes et al., 2017).
93 While it is clear that MHWs can have severe consequences on a variety of systems and
94 organisms, their effects on marine microbes and the biogeochemical processes that they
95 mediate have rarely been investigated (Joint and Smale, 2017).

96

97 While there is evidence that increases in seawater temperature can lead to increased DMSP
98 and/or DMS concentrations in phytoplankton (McLenon and DiTullio, 2012;Sunda et al.,
99 2002), it is not clear how a shift in DMSP net production by phytoplankton under acute
100 temperature stress will alter the composition and function of their associated microbiome and
101 how, in turn, this will influence biogenic sulfur cycling processes within marine habitats. There
102 is therefore a pressing need to understand the physiological and biogeochemical consequences
103 of thermal stress on phytoplankton-bacteria interactions within the context of events such as
104 MHWs. This is particularly important, given that a shift in the composition of the
105 phytoplankton microbiome could potentially dictate atmospheric DMS fluxes depending on
106 whether the bacterial community preferentially cleave or demethylate DMSP (Todd et al.,
107 2007;Kiene et al., 2000).

108

109 The aims of this study were to investigate how an acute increase in temperature (+12°C),
110 comparable to those associated with MHW events and leading to thermal stress in *A. minutum*
111 could alter the physiological state and biogenic sulfur cycling dynamics of *A. minutum* and
112 determine how these changes might influence the composition of the *Alexandrium* microbiome.
113 We hypothesized that an abrupt increase in temperature would lead to physiological
114 impairment (Falk et al., 1996;Robison and Warner, 2006;Iglesias-Prieto et al., 1992;Rajadurai
115 et al., 2005) and oxidative stress (Lesser, 2006) in *A. minutum*, leading to an up-regulation of
116 DMSP, DMS and DMSO production (McLenon and DiTullio, 2012;Sunda et al., 2002) in this

117 high DMSP producer, which could ultimately lead to a shift in the composition of the *A.*
118 *minutum* microbiome.

119 **2. Methods**

120

121 *2.1. Culturing and experimental design*

122 Cultures of *Alexandrium minutum* (CS-324), isolated from Southern Australian coastal waters
123 (Port River, Adelaide, 11/11/1988, CSIRO, ANACC's collection) were grown in GSe medium
124 at 18°C and 50 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$ under a 12:12 light:dark cycle. One month before the start
125 of each experiment, *A. minutum* cultures were acclimated over four generations to 20°C
126 (average summer temperature at Port River, IMOS) and 200 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$ using a
127 14:10 h light:dark cycle mimicking summer conditions. Light intensity was comparable to that
128 used in Berdalet et al. (2011) for *A. minutum* and conveniently allow to grow other algae
129 cultures using the same facilities. Cultures were grown to a cell concentration of ~60,000
130 mL^{-1} before cells were inoculated into fresh GSe medium. Six days prior to the start of each
131 experiment, 20 L of GSe medium was inoculated with a cell concentration of 1,140 mL^{-1}
132 (experiment 1, April 2016) and 680 mL^{-1} (experiment 2, June 2016) and aliquots of 500 mL
133 were transferred into 40 individual 750 mL sterile tissue culture flasks. Culture flasks were
134 incubated in four independent water baths (10 flasks in each) and maintained under control
135 conditions of 20°C and 200 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$. Temperature and light control was achieved
136 using circulating water heaters (Julabo, USA) and programmable LED lights (Hydra FiftyTwo,
137 Aqualllumination, USA). All cultures were mixed twice daily to keep cells in suspension by
138 gentle swirling.

139

140 On Day 1 (T_0), five culture flasks from each 20°C water bath were transferred to four new
141 water baths for exposure to experimental treatment temperatures (either 24°C experiment 1; or
142 32°C, experiment 2), so that each control and experimental water bath contained five flasks.
143 Experimental temperatures were carefully chosen based on preliminary experiments conducted
144 at 24°C, 28°C, 30°C and 32°C, where only a 12°C increase in temperature (32°C treatment)
145 led to a physiological stress response in this robust strain of *A. minutum* in culture. Although
146 an increase in temperature of this magnitude might be rare in coastal marine systems, this
147 presented a unique opportunity to investigate the consequences of MHW-induced thermal
148 stress on this relevant phytoplankton. One culture flask from each tank was immediately
149 sampled for baseline measurements of: DMS (2 mL), DMSP and DMSO (1 mL)
150 concentrations, photochemical efficiency (3 mL), algal and bacterial cell counts (1 mL), ROS
151 quantification (1 mL) and DNA extraction (~470 mL). The dissolved DMSP fraction was not
152 determined because preliminary investigations showed that gravity filtration was too time

153 consuming, potentially due to clogging of filters by the large *A. minutum* cells (30 μm
154 diameter), leading to filtration artefacts for DMSP analysis, as have previously been mentioned
155 by Berdalet et al. (2011). At 18:00 on Day 1 (T_6), 12:00 on Day 2 (T_{24}), 12:00 on Day 5 (T_{96})
156 and 12:00 on Day 6 (T_{120}), one flask from each of the eight water baths was removed from the
157 incubation conditions and sampled as described above.

158

159 *2.2. Photosynthetic efficiency measurements*

160 Subsamples for measurement of photosynthetic efficiency were dark adapted for 10 min under
161 aluminium foil and transferred to a quartz cuvette for Pulse Amplitude Modulated (PAM)
162 fluorometric analysis using a Water PAM (Walz GmbH, Effeltrich, Germany). Once the base
163 fluorescence (F_0) signal had stabilized (measuring light intensity 3, frequency 2s), a saturating
164 pulse (intensity 12, Width 0.8s) was used to measure the maximum quantum yield (F_v/F_M) of
165 photosystem II (PSII). As base fluorescence is dependent on cell density, the photomultiplier
166 gain was adjusted and recorded to maintain F_0 at a level of 0.2 a.u. before saturating the
167 photosystem. Samples were kept in suspension during measurements via continuous stirring at
168 minimal speed inside the quartz cuvette to avoid cells settling.

169

170 *2.3. Microalgal and bacterial cell counts*

171 Subsamples for bacterial cell counts were stained with SYBR Green at a final concentration of
172 1:10,000 and incubated in the dark for 15 min (Marie et al. 1997). Subsamples for microalgal
173 cell counts and stained subsamples for bacterial cell counts were diluted 1:10 and 1:100
174 respectively into sterile GSe medium prior to analysis with a BD Accuri C6 Flow Cytometer
175 (Becton Dickinson). Phytoplankton cells were discriminated using red auto-fluorescence and
176 side scatter (SSC), whereas bacterial populations were discriminated and quantified using
177 SYBR green fluorescence and SSC.

178

179 *2.4. Reactive oxygen species measurements*

180 The presence of reactive oxygen species (ROS) was detected within cultures using the
181 fluorescent probe 2,7-dichlorodihydrofluorescein-diacetate (CM-H2DCFDA; Molecular
182 Probes), which binds to ROS and other peroxides (Rastogi et al., 2010). The reagent was
183 thawed at room temperature for 10 min and activated using 86.5 μL of DMSO, with 5 μL of
184 activated reagent added to each sample (final concentration 5 μM). Samples were vortexed for
185 5 sec and incubated at room temperature for 30 min. Samples were then centrifuged at 2,000 g
186 for 2 min, the supernatant with reagent dye was discarded, and stained cells were resuspended

187 in 1 mL of PBS, prior to quantification of fluorescence by flow cytometry. Mean green
188 fluorescence was quantified from cytograms of forward light scatter (FSC) against green
189 fluorescence. A positive (+ 10 μ L of 30% H₂O₂, final concentration 97mM) and negative (no
190 ROS added) control of PBS were run to ensure that detected cell fluorescence was completely
191 attributable to the ROS probe.

192

193 *2.5.Sulfur analysis by gas chromatography*

194 The preparation of all blanks and samples used in the dilution steps described below were
195 prepared with sterile (0.2 μ M filtered and autoclaved) phosphate-buffered saline (PBS, salinity
196 35ppt) to avoid cell damage from altered osmolarity and to maintain similar physical properties
197 as seawater during headspace analysis by gas chromatography. Aliquots for DMS analysis were
198 transferred into 14 mL headspace vials that were immediately capped and crimped using butyl
199 rubber septa (Sigma Aldrich Pty 27232) and aluminum caps (Sigma Aldrich Pty 27227-U),
200 respectively. DMSP aliquots were 1:1 diluted with sterile PBS and DMSP was cleaved to DMS
201 by adding 1 pellet of NaOH to each vial, which was immediately capped and crimped. Samples
202 were incubated for a minimum of 30 min at room temperature to allow for the alkaline reaction
203 and equilibration to occur prior to analysis by gas chromatography (Kiene and Slezak, 2006).

204

205 DMS and DMSP samples were analyzed by 500 μ L direct headspace injections using a
206 Shimadzu Gas Chromatograph (GC-2010 Plus) coupled with a flame photometric detector
207 (FPD) set at 180°C with instrument grade air and hydrogen flow rates set at 60 mL min⁻¹ and
208 40 mL min⁻¹, respectively. DMS was eluted on a capillary column (30 m x 0.32 mm x 5 μ m)
209 set at 120°C using high purity Helium (He) as the carrier gas at a constant flow rate of 5 mL
210 min⁻¹ and a split ratio of five. A six-point calibration curve and PBS blanks were run by 500
211 μ L direct headspace injections prior to subsampling culture flasks using small volumes of
212 concentrated DMSP.HCl standard solutions (certified reference material WR002, purity 90.3
213 \pm 1.8% mass fraction, National Measurement Institute, Sydney, Australia) that were diluted in
214 sterile PBS to a final volume of 2 mL. Detection limit was 50 nM for 500 μ L headspace
215 injections. Concentrations obtained in vials treated with NaOH accounted for both DMS and
216 DMSP. Consequently, DMSP concentration in each sample was obtained by subtracting the
217 corresponding DMS concentration.

218

219 Following DMS and DMSP analysis, alkaline samples used for DMSP analysis were uncapped
220 and left to vent overnight under a fume hood. On the next day, samples were purged for 10 min

221 with high purity N₂ at an approximate flow rate of 60 mL min⁻¹ to remove any remaining DMS
222 produced from the alkaline treatment. Samples were then neutralized by adding 80 µL of 32 %
223 HCl and DMSO was converted to DMS by adding 350 µL of 12 % TiCl₃ solution to each vial,
224 which was then immediately capped and crimped (Kiene and Gerard, 1994; Deschaseaux et al.,
225 2014b). Vials were then heated in a water bath at 50°C for 1h and cooled down to room
226 temperature prior to analysis by 500 µL direct headspace injections on the GC-FPD as
227 described above. A 5-point calibration curve was run prior to DMSO analysis using DMSO
228 standard solutions (Sigma Aldrich Pty, D2650) diluted in PBS to a final volume of 2 mL and
229 converted to DMS with TiCl₃ in the same manner as the experimental samples. PBS blanks
230 treated with NaOH and TiCl₃ were also run along with the calibration curves. All dimethylated
231 sulfur compounds were normalised to cell density, which best reflects biogenic production.

232

233 *2.6. DNA extraction*

234 Following sub-sampling for the physiological and biogenic sulfur measurements described
235 above, the remaining 400 mL within each culture flask was filtered onto a 47 mm diameter,
236 0.22 µm polycarbonate filter (Millipore) with a peristaltic pump at a rate of 80 rpm to retain
237 cells for DNA analysis. The filters were subsequently stored in cryovials, snap frozen with
238 liquid nitrogen and stored at -80°C until extraction. DNA extraction was performed using a
239 bead-beating and chemical lysis based DNA extraction kit (PowerWater DNA Isolation Kit,
240 MoBio Laboratories) following the manufacturer's instructions. DNA quantity and purity were
241 checked for each sample using a Nanodrop 2000 (Thermo Fisher Scientific, Wilmington, DE,
242 USA). Three replicate samples with the highest DNA quantity and purity from the control and
243 treatment tanks, collected at the beginning (T₀) and end (T₁₂₀) of the experiment, were
244 subsequently sequenced.

245

246 *2.7. 16S rRNA amplicon sequencing and bioinformatics*

247 To characterize the bacterial assemblage structure (microbiome) of *A. minutum* cultures, we
248 employed 16S rRNA amplicon sequencing. We amplified the V1-V3 variable regions of the
249 16S rRNA gene using the 27F (AGAGTTTGATCMTGGCTCAG, Lane, 1991) and 519R
250 (GWATTACCGCGGCKGCTG, Turner et al., 1999) primer pairing, with amplicons
251 subsequently sequenced using the Illumina MiSeq platform (Ramaciotti Centre for Genomics;
252 Sydney, NSW, Australia) following the manufacturer's guidelines. Raw data files in FASTQ
253 format were deposited in the National Center for Biotechnology Information (NCBI) Sequence
254 Read Archive (SRA) under the study accession number PRJNA486692.

255 Bacterial 16S rRNA gene sequencing reads were analysed using the QIIME pipeline (Caporaso
256 et al., 2010;Kuczynski et al., 2012). Briefly, paired-end DNA sequences were joined, de novo
257 Operational Taxonomic Units (OTUs) were defined at 97% sequence identity using UCLUST
258 (Edgar, 2010) and taxonomy was assigned against the SILVA v128 database (Quast et al.,
259 2012;Yilmaz et al., 2013). Chimeric sequences were detected using usearch61 (Edgar, 2010)
260 and together with chloroplast OTUs were filtered from the dataset. Sequences were then
261 aligned, filtered and rarefied to the same depth to remove the effect of sampling effort upon
262 analysis.

263

264 *2.8. Statistical analysis*

265 Repeated measures analysis of variance (rmANOVA) models were fitted to the data to quantify
266 the effects of temperature and time (fixed factors) on all response variables measured in this
267 experiment (cell density, F_V/F_M , ROS, DMS, DMSP and DMSO concentrations) using IBM
268 SPSS Statistics 20. Assumptions of sphericity were tested using Mauchly's test. In cases where
269 this assumption was violated, the degrees of freedom were adjusted using the Greenhouse-
270 Geisser correction factor. Bonferroni adjustments were used for pairwise comparisons. Each
271 variable was tested for the assumption of normality and log, ln or sqrt transformations were
272 applied when necessary.

273

274 For sequencing data, alpha diversity parameters of the rarefied sequences and Jackknife
275 Comparison of the weighted sequence data (beta diversity) were calculated in
276 QIIME (Caporaso et al., 2010). A two-way PERMANOVA with Bray-Curtis similarity
277 measurements was performed on abundance data of taxonomic groups that contained more
278 than 1% of total generated OTUs (represent 90.23% of the data) using PAST (Hammer et al.,
279 2008). In addition, PAST was used to perform non-metric multidimensional scaling (nMDS)
280 analysis and isolate the environmental parameters (normalised as follows: $(x-\text{mean})/\text{stdev}$) that
281 contributed the most to the differences between groups using the Bray-Curtis similarity
282 measure. SIMPER analysis performed with the White *t*-test was used to identify the taxonomic
283 groups that significantly contributed the most to the shift in bacterial composition in *A.*
284 *minutum* cultures over time and between temperature treatments.

285

286 **3. Results**

287

288 *3.1. Algal growth and physiological response*

289 *A. minutum* cell abundance exponentially increased over time in both the control (20°C) and
290 24°C temperature treatment, but a significantly faster growth rate ($p = 0.001$, t -test) occurred
291 at 24°C ($2.66 \pm 0.01 \text{ d}^{-1}$; average \pm SE) compared to the 20°C control ($2.57 \pm 0.01 \text{ d}^{-1}$), resulting
292 in significantly greater cell abundance at 96h ($p = 0.007$) and 120h ($p < 0.001$) (rmANOVA,
293 **Table 1, Fig. 1a**). On the other hand, the 32°C treatment resulted in decreased growth rates
294 ($2.40 \pm 0.02 \text{ d}^{-1}$ versus $2.58 \pm 0.02 \text{ d}^{-1}$; t -test) and significantly lower cell abundance, relative
295 to the 20°C control, at all time points from 6h after the start of the experiment ($p \leq 0.03$;
296 rmANOVA, **Table 1, Fig. 1b**). *A. minutum* abundance demonstrated a marked decline on day
297 5 in the 32°C treatment.

298

299 No significant difference in the maximum quantum yield (F_V/F_M) of *A. minutum* cultures
300 occurred between 20°C and 24°C until 120h after the start of the experiment, where a
301 significantly lower F_V/F_M occurred in the 24°C treatment ($p = 0.01$; rmANOVA, **Table 1, Fig.**
302 **2a**). In contrast, F_V/F_M was significantly lower in *A. minutum* cultures maintained at 32°C
303 compared to the 20°C control at all time points from 6h after the start of the experiment ($p \leq$
304 0.01 ; rmANOVA, **Table 1, Fig. 2b**). However, on days 5 and 6, the F_V/F_M of cultures kept at
305 32°C recovered to values (0.72 ± 0.008) close to those of the control (0.75 ± 0.004) (**Fig. 2B**),
306 although it remained significantly lower than at 20°C ($p < 0.01$ and $p < 0.001$ on day 5 and 6,
307 respectively).

308

309 3.2. Reactive oxygen species (ROS)

310 Significantly lower concentrations of ROS were measured at 24°C than at 20°C at 96h ($p =$
311 0.003) and 120h ($p = 0.03$) (rmANOVA, **Table 1, Fig. 2c**). In contrast, significantly greater
312 concentrations of ROS were measured at 32°C than at 20°C 24h ($p < 0.001$), 96h ($p = 0.001$)
313 and 120h ($p = 0.01$) after the start of the experiment (rmANOVA, **Table 1, Fig. 2d**). In-line
314 with the recovery in measured F_V/F_M , ROS concentrations in cultures kept at 32°C started to
315 decline to values closer to those of the control on days 5 and 6 of the experiment (**Fig. 2d**). A
316 significant negative correlation between F_V/F_M levels and ROS concentrations was observed
317 under the 32°C temperature treatment ($R^2 = 0.623$; $p = 0.02$, $n = 18$; **Fig. 3**).

318

319 3.3. Biogenic sulfur dynamics

320 Biogenic concentrations of DMSP, DMS and DMSO ranged from 424 ± 35 to $1629 \pm 170 \text{ fmol}$
321 cell^{-1} , from 13 ± 1.02 to $87 \pm 5 \text{ fmol cell}^{-1}$ and from 9 ± 1.41 to $94 \pm 24 \text{ fmol cell}^{-1}$, respectively,
322 over both experiments (**Fig. 4**). Concentrations of all three sulfur compounds slowly decreased

323 over time in all *A. minutum* cultures regardless of the temperature treatment. No significant
324 difference in DMSP concentration was recorded between 20°C and 24°C throughout the
325 experiment ($p > 0.05$; rmANOVA, **Table 1, Fig. 4a**), whereas significantly less DMSP was
326 measured in cells at 32°C than in the 20°C control at 96h ($p = 0.02$; rmANOVA, **Table 1, Fig.**
327 **4b**).

328

329 Significantly lower DMS concentrations were measured at 24°C compared to 20°C at 24h (p
330 < 0.001) and 120h ($p = 0.002$) (rmANOVA, **Table 1, Fig. 4c**). In contrast, DMS was
331 significantly higher at 32°C than 20°C 6h after the start of the experiment ($p = 0.008$;
332 rmANOVA, **Table 1, Fig. 4d**). A similar pattern was observed for DMSO, where relative to
333 the controls, it was significantly lower at 24°C 24h after the start of the experiment ($p = 0.001$;
334 rmANOVA, **Table 1, Fig. 4e**) and significantly greater at 32°C after 6h and 24h ($p < 0.05$, **Fig.**
335 **4f**).

336

337 3.4. Bacterial abundance and composition

338 Bacterial cell abundance exponentially increased over time at both 20°C and 24°C (**Fig. 5a**).
339 Bacterial abundance was significantly greater at 24°C than at 20°C 120 h after the start of the
340 experiment ($p = 0.05$; rmANOVA, **Table 1, Fig. 5a**). However, no significant difference ($p >$
341 0.05 , t -test) in bacterial growth rate was observed between 20°C ($4.15 \pm 0.05 \text{ d}^{-1}$) and 24°C
342 ($4.18 \pm 0.01 \text{ d}^{-1}$). In contrast, bacterial growth rate was significantly lower at 32°C than in the
343 20°C control ($4.05 \pm 0.01 \text{ d}^{-1}$ versus $4.23 \pm 0.02 \text{ d}^{-1}$; $p < 0.001$, t -test) (**Fig. 5b**), resulting in
344 significantly lower bacterial cell densities at 24h ($p = 0.002$), 96h ($p = 0.002$) and 120h ($p <$
345 0.001) relative to the control (rmANOVA, **Table 1, Fig. 5b**).

346

347 The composition of the initial (T_0) *A. minutum* microbiome was consistent across all samples,
348 but then diverged significantly with time and between temperature treatments (**Fig. 6a-b**; Bray-
349 Curtis similarity measurement, Shepard plot stress = 0.0587). A significant temporal shift in
350 bacterial composition occurred at both 20°C and 32°C, with dissimilarities in community
351 composition between T_0 and T_{120} of 27% and 42% occurring respectively (SIMPER analysis).
352 Notably, bacterial communities at 32°C differed significantly (two-way PERMANOVA; $p <$
353 0.05) to 20°C at T_{120} , with 32% dissimilarity in community composition. These differences
354 were primarily driven by increased relative abundance of bacterial Operational Taxonomic
355 units (OTUs) within the *Oceanicaulis* (17%), *Phycisphaeraceae SMIA02* (8.8%) and *Balneola*
356 (4.9%) genus along with a decline in the relative abundance of OTUs matching *Maribacter*

357 (24%), *Marinoscillum* (4.7%) and *Seohaecicola* (2.7%) (*Rhodobacter* family) in the 32°C
358 treatment (White test, **Fig. 6c**), with all taxa cumulatively contributing to 63% of the OTU
359 differences between temperature treatments at T₁₂₀ (SIMPER analysis). In the 32°C treatment,
360 differences in microbiome composition between T₀ and T₁₂₀ were aligned with the elevated
361 levels of ROS, while in the control (20°C) the community shift was principally aligned with
362 differences in bacterial and algal cell abundance (**Fig. 6a**; MDS analysis). Similarly, the
363 elevated concentration of ROS as well as the lower F_v/F_M, lower algal and bacterial cell
364 abundance and lower DMSP, DMS and DMSO concentrations at 32°C were aligned with the
365 differences in microbiome composition between the temperature treatments (**Fig. 6b**; MDS
366 analysis)

367

368 **4. Discussion**

369

370 Climate change induced shifts within marine ecosystems are predicted to fundamentally alter
371 the physiology of planktonic organisms and the biogeochemical transformations that they
372 mediate (Finkel et al., 2009; Tortell et al., 2008; Hallegraeff, 2010). Rising seawater
373 temperatures are one of the major impacts of climate change on marine ecosystems (Harley et
374 al., 2006), and can be manifested both as long-term gradual increases (IPCC, 2007, 2013) or
375 intense episodic marine heatwaves (Frölicher and Laufkötter, 2018; Hobday et al., 2016).
376 Although less examined to date than chronic temperature increases, MHWs are predicted to
377 become more frequent and severe (Oliver et al., 2018) and have been proposed as a mechanism
378 for triggering toxic algal blooms (Ummenhofer and Meehl, 2017). Against this backdrop of
379 changing environmental conditions, microbial production and cycling of dimethylated sulfur
380 compounds could be particularly relevant because they simultaneously play a role in the stress
381 response of marine phytoplankton (Berdalet et al., 2011; Deschaseaux et al., 2014a; Sunda et
382 al., 2002; Wolfe et al., 2002; Stefels and van Leeuwe, 1998) and have been predicted to have
383 biogeochemical feed-back effects that are relevant for local climatic processes (Charlson et al.,
384 1987).

385

386 This study investigated the biogenic sulfur cycling dynamics of *A. minutum*, and its
387 microbiome, in response to an extreme, short-term thermal stress event, akin to the marine
388 heat-wave events occurring with increasing frequency within coastal habitats (Oliver et al.,
389 2018). Indeed, MHWs have been defined as an abrupt and ephemeral increase in temperature
390 of at least 3 to 5°C above climatological average that lasts for at least 3 to 5 days (Hobday et
391 al., 2016). Large increases in temperature of about 8°C above the monthly climatological

392 average led to red-tides of exceptional density in San Francisco Bay (Cloern et al., 2005). While
393 a 12°C increase in temperature constitutes an extreme scenario of MHWs, even for coastal
394 habitats, this experimental temperature was selected after preliminary investigations with the
395 intention to induce thermal stress in this particularly robust strain of *A. minutum* in culture.

396

397 *A. minutum* has been targeted in this study as 1) an ecologically relevant phytoplankton
398 responsible for some of the most harmful algal blooms (Anderson et al., 2012) and 2) as
399 biochemically relevant for containing the highest DMSP concentrations ever reported in marine
400 dinoflagellates (Caruana and Malin, 2014). However, it is to be noted that DMSP
401 concentrations reported in this study were a degree of magnitude lower (0.42 ± 0.04 to $1.63 \pm$
402 1.70 pmol cell⁻¹) than that previously reported for *A. minutum* (14.2 pmol cell⁻¹; Caruana and
403 Malin, 2014; Jean et al., 2005). This is potentially because this culture of *A. minutum* had been
404 isolated from free-living *A. minutum* for a long time (1988) or because culturing conditions
405 failed to mimic the exact same biochemical conditions in which this strain of *A. minutum*
406 usually grow. This biochemical difference could potentially reflect that this strain of *A.*
407 *minutum* in culture was more robust than free-living dinoflagellates of the same species,
408 thereby potentially justifying the need of a 12°C increase in temperature to induce thermal-
409 stress.

410

411 *4.1. Effects of thermal stress on A. minutum growth, physiology and ROS production*

412 A mild increase in temperature (4°C) resulted in faster algal growth and lower oxidative stress,
413 indicating that 24°C was close to a temperature optimum for this strain of *Alexandrium*. This
414 is perhaps not surprising considering that *Alexandrium* species are capable of growing under a
415 wide range of temperatures from 12°C to 25°C (Laabir et al., 2011). In contrast, an extreme
416 increase in temperature (12°C) resulted in a rapid and clear cascade of physiological responses,
417 indicative of an acute thermal stress response in *A. minutum*. Overall, *A. minutum* cells exposed
418 to 32°C immediately exhibited slower growth relative to the 20°C control, suggesting that a
419 12°C increase in temperature rapidly led to either an increase in cell death rate or a decrease in
420 cell division (Rajadurai et al., 2005; Veldhuis et al., 2001). The slower growth rate at 32°C was
421 coupled with a drop in photosynthetic efficiency and an increase in ROS concentrations, which
422 are both common stress responses to thermal stress in marine algae (Lesser, 2006; Falk et al.,
423 1996; Robison and Warner, 2006; Iglesias-Prieto et al., 1992). In fact, these two physiological
424 responses are often interconnected as increased ROS production generally occurs in both the
425 chloroplast and mitochondria of marine algae exposed to thermal stress, causing lipid

426 peroxidation and ultimately leading to a loss in thylakoid membrane integrity (Falk et al., 1996)
427 and a decrease in the quantum yield of PSII (Lesser, 2006). This was reflected in the negative
428 correlation observed between the maximum quantum yield of PSII and ROS concentrations.

429

430 Although photosynthetic efficiency remained impaired and ROS concentrations remained high
431 under 32°C until the end of the experiment, both biomarkers of stress started to return to values
432 closer to those of the 20°C control by day 5 and 6 of the experiment. This was most likely at
433 the expense of a decline in algal abundance since slow growth often coincides with concurrent
434 cellular repair and photosystem activity recovery (Robison and Warner, 2006). The differential
435 physiological response between 24°C and 32°C indicates that although cultures of this strain
436 of *A. minutum* appear to be highly resistant to temperature changes, an abrupt increase in
437 temperature of 12°C simulating an extreme case of marine heatwave led to a clear stress
438 response. The physiological pattern at 32°C also suggested an acclimation period necessary for
439 such an abrupt shift in temperature, especially since recovery (in F_v/F_m and ROS levels) was
440 observed towards the end of the experiment.

441

442 *4.2. Biogenic sulfur cycling as a response to thermal stress in A. minutum*

443 Biogenic organic compounds are key compounds in the stress response of phytoplankton, with
444 evidence they can be used in response to changes in temperature (Van Rijssel and Gieskes,
445 2002;Stefels, 2000). An up-regulation of the biogenic sulfur yield was expected as a stress
446 response to increased temperature in *A. minutum*, through either an increase in cellular DMSP
447 concentrations, or an increase in DMS via the cleavage of DMSP (McLenon and DiTullio,
448 2012;Berdalet et al., 2011;Wolfe et al., 2002;Sunda et al., 2002). No significant change in
449 DMSP concentrations was observed between the control and 24°C treatment, where, as
450 described above, physiological responses converged to indicate that 24°C was in fact a more
451 optimal growth temperature for this organism. This temperature optimum was generally
452 associated with lower DMS and DMSO concentrations than in the 20°C control, although this
453 was only evident 24h after the start of the experiment. Since algal stress responses often result
454 in increased cellular sulfur concentrations in dinoflagellates (McLenon and DiTullio,
455 2012;Berdalet et al., 2011), it is perhaps not surprising that DMS and DMSO concentrations
456 were lower under what appear to have been more optimal growth temperature conditions.

457

458 In contrast to the lower DMS and DMSO concentrations observed at 24°C compared to the
459 20°C control, exposure to 32°C resulted in spikes in DMS and DMSO 6h after the start of the

460 experiment, which accompanied decreased algal growth and impaired photosystem II.
461 Although sporadic, the increases in DMS and DMSO observed in the 32°C treatment may have
462 resulted from enhanced intracellular DMSP cleavage by phytoplankton (Del Valle et al., 2011)
463 or enhanced DMSP exudation from phytoplankton cells during cell lysis (Simó, 2001),
464 resulting in an increasing pool of dissolved DMSP made readily available to both bacteria and
465 phytoplankton DMSP-lyases (Riedel et al., 2015;Alcolombri et al., 2015;Todd et al.,
466 2009;Todd et al., 2007). However, it is notable that lower DMSP concentrations in the 32°C
467 treatment than in the control only occurred on day 4, whereas the spike in DMS and DMSO
468 were evident at the outset of the experiment (6h). Since this decrease in DMSP at 96h was not
469 coupled with an increase in DMS, this could alternatively be indicative of a decrease in
470 methionine synthase activity (McLenon and DiTullio, 2012) or assimilation of DMSP-sulfur
471 by bacterioplankton for *de novo* protein synthesis (Kiene et al., 2000), with this demethylation
472 pathway often accounting for more than 80% of DMSP turnover in marine surface waters. The
473 spike in DMSO measured 6h after the increase in temperature to 32°C most likely indicated
474 rapid DMS oxidation by ROS under thermal stress (Sunda et al., 2002;Niki et al., 2000). At
475 that time however, we found no evidence for ROS build up in *A. minutum* cultures, possibly
476 because ROS concentrations were kept in check by sufficient DMS synthesis and active DMS-
477 mediated ROS scavenging (Lesser, 2006;Sunda et al., 2002). In contrast, 24h after the start of
478 the experiment, increased ROS coincided with an abrupt decline in DMS and DMSO, perhaps
479 suggestive of serial oxidation via active ROS scavenging of both DMS to DMSO and DMSO
480 to methane sulfinic acid (MSNA) (Sunda et al., 2002), although it is always difficult to
481 confidently link DMS(O) and ROS dynamics unless using tracing techniques. It is also to be
482 noted that measuring standing stocks may constitute a limitation to capture subtle changes in
483 DMS, DMSP and DMSO over time.

484

485 The only previous study that has examined sulfur responses to stress exposure in *A. minutum*
486 examined the effect of physical turbulence by shaking *A. minutum* cultures for up to four days
487 (Berdalet et al., 2011). While the authors of that study also observed slower cell growth as a
488 response to stress exposure, in contrast to our study, cellular DMSP concentrations increased
489 by 20%. Here, a drop in DMSP concentration was observed at 96h between the control and
490 temperature treatment. Therefore, even though DMSP concentrations were quantified with a
491 similar approach as in Berdalet et al. (2011) (no filtration of the samples with assuming that
492 particulate DMSP concentrations overrule dissolved DMSP and DMS concentrations), it seems
493 that heat stress and turbulence triggered a dissimilar sulfur response to stress in *A. minutum*.

494

495 Overall, a 12°C increase in temperature led to lower photosynthetic efficiency, increased
496 oxidative stress and slower cell growth in this robust strain of the red-tide mediating
497 dinoflagellate *A. minutum*. This physiological stress response was coupled with a differential
498 biogenic sulfur cycling as shown by spikes in DMS and DMSO as well as lower DMSP
499 concentrations, most likely translating ROS scavenging and DMSP uptake by
500 bacterioplankton, respectively. Because the turnover of DMS, DMSP and DMSO in biological
501 systems can occur very quickly (Simo et al 2000), DMS and DMSO concentrations can change
502 rapidly, which sometimes makes it difficult to clearly establish cause-effect relationships
503 between physiological stress and the biogenic sulfur response.

504

505 4.3. A shift in *A. minutum* associated-bacteria composition triggered by thermal stress

506 In light of DMSP and related biogenic sulfur compounds constituting an important source of
507 carbon and sulfur to phytoplankton-associated bacteria (Kiene et al., 2000), it follows that any
508 shift in biogenic sulfur concentrations could influence the microbiome composition of *A.*
509 *minutum*. However, it is undeniable that a shift in the microbial community could also be driven
510 by a range of physiological and biochemical parameters that were not measured in this study.
511 Nevertheless, the most pronounced temporal shift in the composition of the bacterial
512 community associated with *A. minutum* occurred in the 32°C treatment. This shift was
513 primarily characterized by a statistically significant increase in the relative abundance of OTUs
514 classified as members of the *Oceanicaulis*, *Phycisphaeraceae* and *Balneola* and a significant
515 decrease in OTUs classified as members of the *Maribacter*, *Marinoscillum* and *Seohaecicola*.

516

517 To predict any potential role of these key OTUs in biogenic sulfur cycling processes, we
518 screened the genomes of members of these groups using BLAST for four genes commonly
519 involved in DMSP catabolism: *dmdA*, CP000031.2 (Howard et al., 2006); *dddP*, KP639186
520 (Todd et al., 2009); *tmm*, JN797862 (Chen et al., 2011); and *dsyB*, KT989543 (Curson et al.,
521 2017). A BLAST query of the sequences in the NCBI nucleotide collection (nr/nt) database
522 revealed that previously sequenced members of the genera *Maribacter* (taxid:252356, 357
523 sequences), *Oceanicaulis* (taxid:153232, 36 sequences), *Marinoscillum* (taxid:643701, 23
524 sequences), *Seohaecicola* (taxid:481178, 18 sequences) and *Balneola* (taxid:455358, 44
525 sequences) did not possess any homologs of these sulfur cycling genes. While no homologs
526 were found in the genus *SMIA02*, perhaps because very little genomic information is available
527 for this genus, a close phylogenetic relative to *SMIA02* (99% query cover, 80% identical, E-

528 value = 0.0), and also a member of the *Phycisphaeraceae* family (*P. mikurensis* 10266;
529 genbank accession numbers AP012338.1), possessed significant homologues to all four query
530 genes involved in DMSP metabolism: *dmdA* (92% identical, E-value < 0.001), *dddP* (87%
531 identical, E-value = 0.003), *tmm* (82% identical, E-value = 0.002) and *dsyB* (92% identical, E-
532 value < 0.001). It is thus possible that the spike in DMS and DMSO concentrations in the early
533 stage of the 32°C heat treatment was a consequence of (or contributed to) the preferential
534 recruitment of *Phycisphaeraceae* SM1A02.

535

536 Some members of the *Rhodobacter* family such as several members of the *Roseobacter* genus
537 and *Rhodobacter sphaeroides* are known to possess homologues of either or both *dmdA* and
538 *ddd* genes, which are responsible for DMSP demethylation and DMSP-to-DMS cleavage,
539 respectively (Howard et al., 2006;Curson et al., 2008). However, none of the available
540 reference genomes for *Seohaecicola*, a member of the *Rhodobacteraceae*, possessed any
541 homologs of targeted biogenic sulfur cycling. Similarly, members of the *Maribacter*, which
542 was the main contributor to the difference in microbiome structure between the control and
543 thermal stress treatment, are known not to possess DMSP/DMS transformation pathways
544 (Kessler et al., 2018). Hence, the decline of this taxa in the heat stress treatments, where an
545 upshift in biogenic sulfur availability occurred, is perhaps not surprising. However, this change
546 in microbial abundance could have also been triggered by a range of other parameters that were
547 not measured in this study.

548

549 Ultimately, the rapid changes in DMS and DMSO concentrations were potentially caused by
550 (or led to) a shift in microbiome composition towards the preferential growth of sulfur-
551 consuming bacteria (e.g. *Phycisphaeraceae* SM1A02) at the expense of other types of bacteria
552 (e.g. *Seohaecicola*). Alternatively, the observed shifts in microbiome structure may have
553 occurred independently to the biogenic sulfur cycling processes and was instead related to
554 either temperature itself or other metabolic shifts in the heat-stressed *A. minutum*. Notably, the
555 temporal shift in bacterial composition under thermal stress was associated with increased
556 cellular ROS at the end of the experiment, indicating a potential link to oxidative stress.

557

558 **5. Conclusion**

559

560 Abrupt and intense increases in seawater temperatures associated with MHWs are predicted to
561 become more frequent and intense (Oliver et al., 2018) and have the potential to influence the

562 structure of coastal microbial assemblages and the nature of the important biogeochemical
563 processes that they mediate. Here, we hypothesized that a very acute increase in temperature,
564 mimicking an extreme scenario of coastal MHWs, would trigger both a physiological and
565 biochemical stress response in the DMSP-producing dinoflagellate *A. minutum*. This response
566 was indeed observed following a 12°C-increase in temperature, with evidence for impaired
567 photosynthetic efficiency, oxidative stress, spikes in DMS and DMSO concentrations, a drop
568 in DMSP concentration and a shift in the composition of the *A. minutum* microbiome. These
569 patterns are indicative of a profound shift in the physiological state and biochemical function
570 of this ecologically relevant dinoflagellate in the context of MHWs and suggest that extreme
571 thermal stress has the potential to not only influence the composition and interactions of coastal
572 microbial food-webs, but re-shape sulfur budgets in coastal waters.

Author contribution

ED designed the experiment with KP and JS. ED conducted the experiment as well as the sample analysis, interpreted the data, conducted the statistical analysis and took the lead on writing the manuscript. JOB assisted with conducting the experiment, sample analysis and data interpretation. NS helped with the statistical analysis and interpretation of the sequencing data. KP and JS financed the experiment.

Competing interests:

The authors declare that they have no conflict of interest.

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Author contribution:

ED, KP and JS devised the experimental design. ED and JOB conducted the thermal stress experiments, including sampling and sample analysis. NS and JOB processed sequencing data while ED processed the physiological and sulfur data. ED wrote the manuscript with significant contributions from all co-authors.

Competing interests:

The authors declare that they have no conflict of interest.

Figure captions

Figure 1 – Algal cell abundance in *A. minutum* cultures in experiment 1 (20°C and 24°C) (A) and experiment 2 (20°C and 32°C) (B); average \pm SE, $n = 4$.

Figure 2 – Photosynthetic efficiency (A, B) and reactive oxygen species (ROS) (C, D) in *A. minutum* cultures in experiment 1 (20°C and 24°C) (A, C) and experiment 2 (20°C and 32°C) (B, D); average \pm SE, $n = 4$.

Figure 3 – Correlation between the photosynthetic efficiency and reactive oxygen species (ROS) in *Alexandrium minutum* under the 32°C thermal stress treatment; $n = 18$.

Figure 4 – DMSP (A, B), DMS (C, D) and DMSO (E, F) concentrations in *A. minutum* cultures in experiment 1 (20°C and 24°C) (A, C, E) and experiment 2 (20°C and 32°C) (B, D, F); average \pm SE, $n = 4$. Variability in between the two 20°C control is probably a consequence of experiments 1 and 2 being conducted at a different times (April and June), whereby changes in the physiological state of the culture at each time led to different dimethylated sulfur profiles.

Figure 5 – Bacterial cell abundance in *A. minutum* cultures in experiment 1 (20°C and 24°C) (A) and experiment 2 (20°C and 32°C) (B); average \pm SE, $n = 4$.

Figure 6 – Multi-dimensional scaling (MDS) of the three phylogenetic groups defined by 16s sequencing of the bacteria population associated with *A. minutum* cultures grown under control conditions (20°C) and acute thermal stress (32°C) at T₀ and T₁₂₀ (A) and MDS excluding the T₀ control (B). Vectors represent the factors that most likely drove the shift in bacterial composition between groups. The taxonomic groups that significantly contributed to the difference in bacterial composition between T₀ and T₁₂₀ at 32°C ⁽¹⁾, between T₀ and T₁₂₀ at 20°C ⁽²⁾ and between 32°C and 20°C at T₁₂₀ ⁽³⁾ appear in bold next to the heatmap (C), with scaling being based on relative abundance.

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Table 1. Output of repeated measures analysis of variance (rmANOVA) for algal (CELLS_A) and bacterial (CELLS_B) cell abundance, photosynthetic efficiency (F_V/F_M), oxidative stress (ROS), dimethylsulfoniopropionate (DMSP), dimethylsulfide (DMS) and dimethylsulfoxide (DMSO) concentrations as a function of temperature (24°C or 32°C) and time. Numbers in bold indicate significant data based on the level of significance $p < 0.05$. df1 = numerator df; df2= denominator df.

Parameters	24°C – mild thermal stress			32°C – mild thermal stress			
	temperature	time	temperature × time	temperature	time	temperature × time	
CELLS _A	<i>F</i>	4.04	335	4.16	27.47	237.62	8.28
	<i>df1</i>	1	4	4	1	2.04	2.04
	<i>df2</i>	6	24	24	6	12.26	12.26
	<i>p</i>	0.91	< 0.001	0.01	< 0.001	< 0.001	0.005
CELLS _B	<i>F</i>	2.13	52.2	1.35	32.56	199.8	22.26
	<i>df1</i>	1	1.29	1.29	1	4	4
	<i>df2</i>	6	7.74	7.74	6	24	24
	<i>p</i>	0.2	< 0.001	0.3	0.001	< 0.001	< 0.001
F _V /F _M	<i>F</i>	0.42	33.43	6.90	48.79	12.58	13.11
	<i>df1</i>	1	4	4	1	1.19	1.19
	<i>df2</i>	6	24	24	5	5.93	5.93
	<i>p</i>	0.54	< 0.001	0.001	0.001	0.01	0.01
ROS	<i>F</i>	37.26	6.30	5.88	33.23	8.85	8.41
	<i>df1</i>	1	4	4	1	2.32	2.32
	<i>df2</i>	6	24	24	6	13.9	13.9
	<i>p</i>	0.001	0.001	0.002	0.001	0.003	0.003
DMSP	<i>F</i>	0.79	31.16	0.95	3.03	15.18	3.17
	<i>df1</i>	1	1.56	1.56	1	4	4
	<i>df2</i>	6	9.35	9.35	6	24	24
	<i>p</i>	0.41	<0.001	0.4	0.13	< 0.001	0.03
DMS	<i>F</i>	51.5	38.73	2.01	5.08	30.77	5.23
	<i>df1</i>	1	2.14	2.14	1	4	4
	<i>df2</i>	6	12.87	12.87	6	24	24
	<i>p</i>	< 0.001	< 0.001	0.17	0.07	< 0.001	0.004
DMSO	<i>F</i>	36.56	26.64	7.21	4.68	14.74	7.14
	<i>df1</i>	1	4	4	1	4	4
	<i>df2</i>	6	24	24	6	24	24
	<i>p</i>	0.001	< 0.001	0.001	0.07	< 0.001	0.001

Figure 1

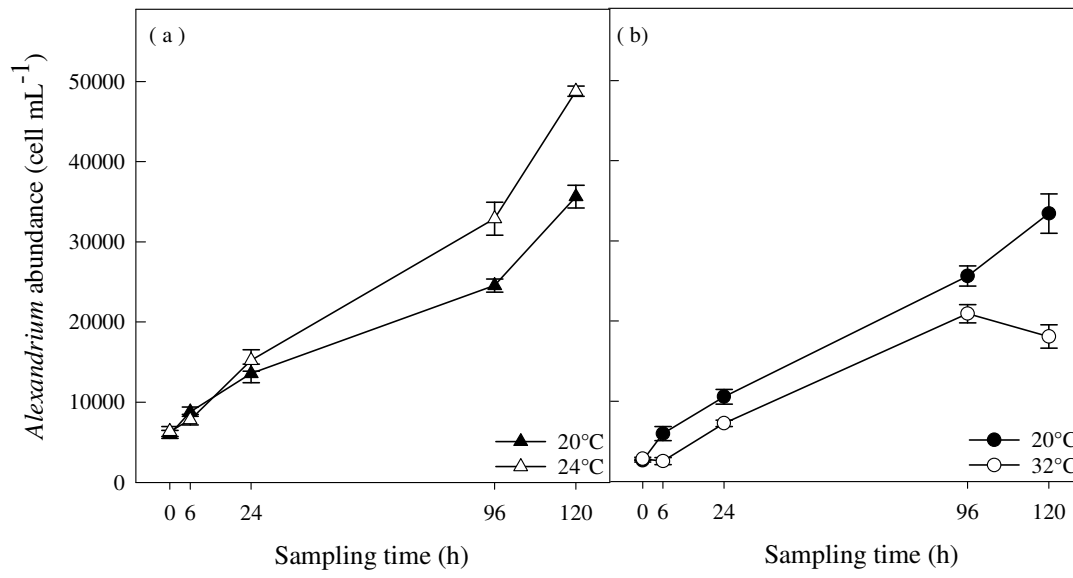


Figure 2

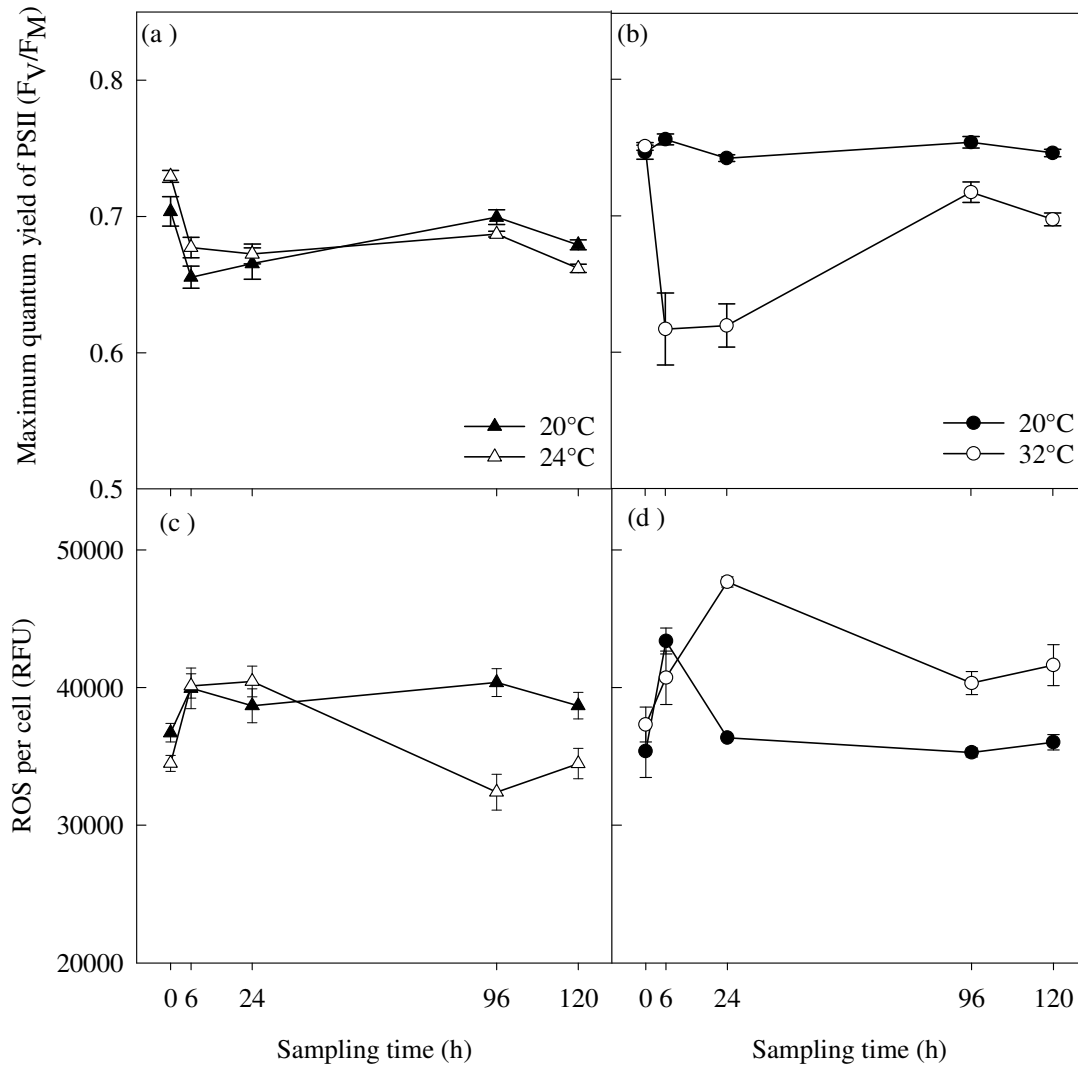


Figure 3

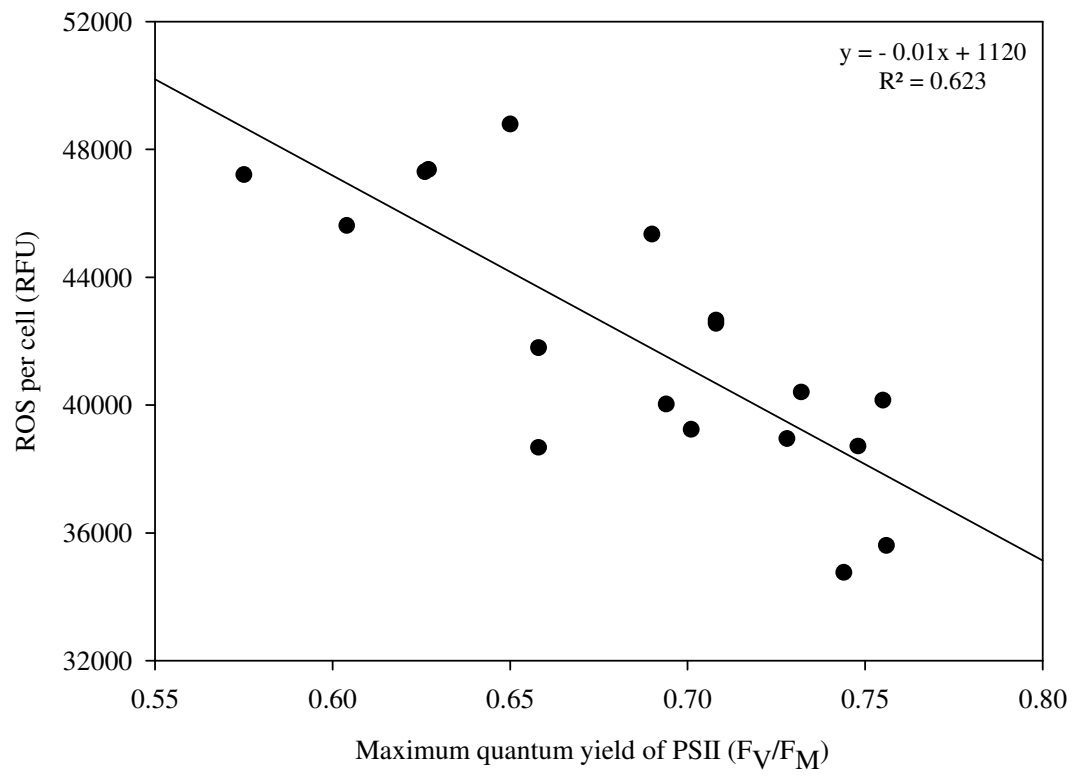


Figure 4

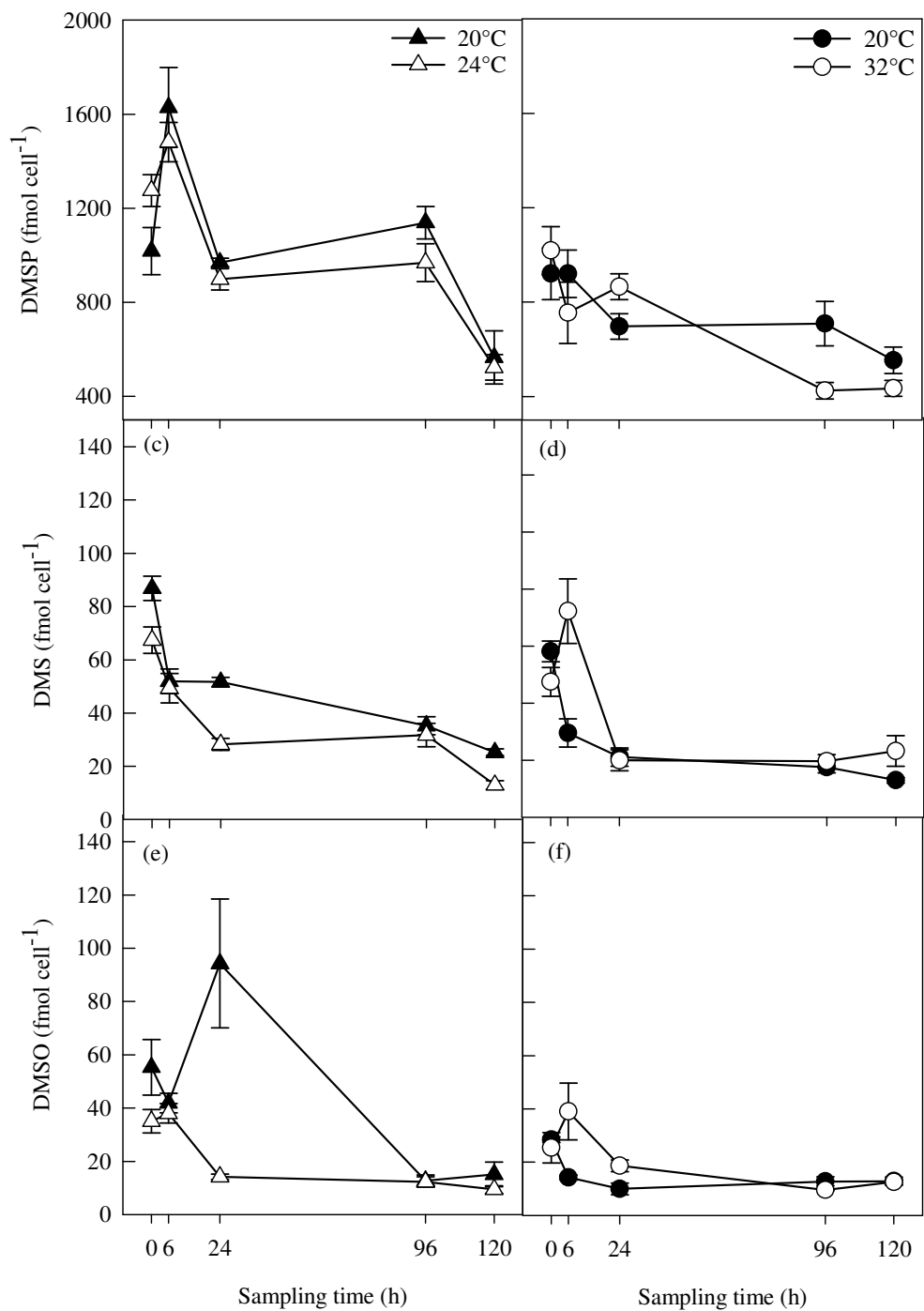


Figure 5

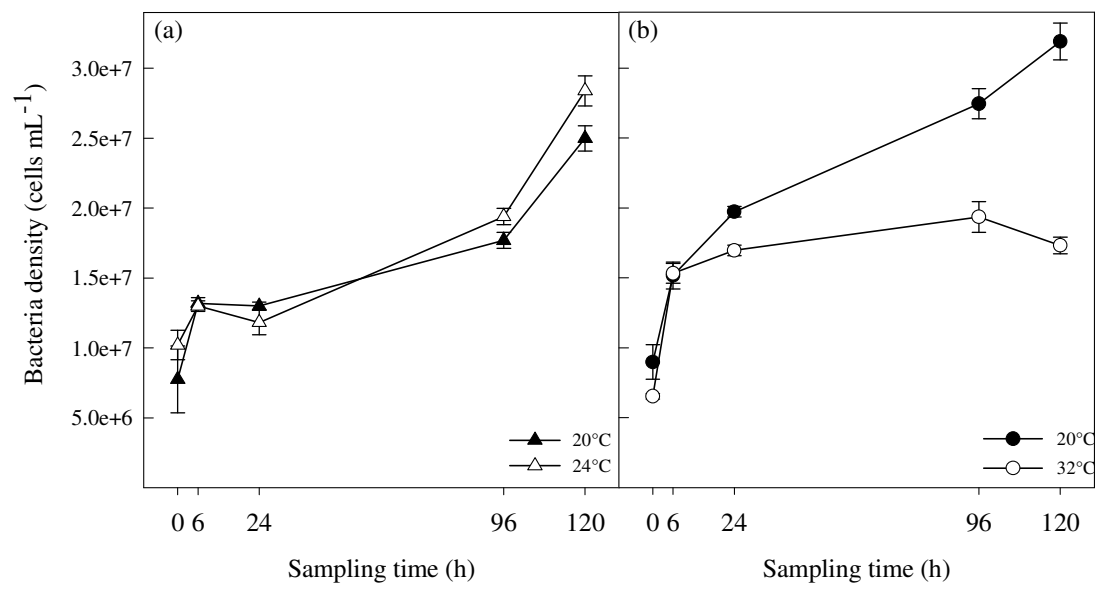


Figure 6

