1	Effect of CO ₂ enrichment on phytoplankton photosynthesis in the
2	North Atlantic sub-tropical gyre
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22 Abstract

23 The effects of changes in CO₂ concentration in seawater on phytoplankton community structure and photosynthesis were studied in the North Atlantic sub-tropical gyre. Three 24 25 shipboard incubations were conducted for 48 h at ~760 ppm CO₂ and control (360 ppm CO₂) from 49°N to 7°N during October and November 2010. Elevated CO₂ caused a decrease in pH 26 27 to ~7.94 compared to ~8.27 in the control. During one experiment, the biomass of nano- and 28 picoeukaryotes increased under CO₂ enrichment, but primary production decreased relative to 29 the control. In two of the experiments the biomass was dominated by dinoflagellates, and there was a significant increase in the maximum photosynthetic rate (P_m^B) and light-limited 30 slope of photosynthesis (α^{B}) at CO₂ concentrations of 750 ppm relative to the controls. 77 K 31 32 emission spectroscopy showed that the higher photosynthetic rates measured under CO₂ 33 enrichment increased the connection of reversible photosystem antennae, which resulted in an 34 increase in light harvesting efficiency and carbon fixation.

35

37 **1. Introduction**

38

39 ocean acidification on the functionality and productivity of marine ecosystems (Riebesell, 40 2008). Increases in atmospheric CO_2 from 280 to 370 ppm since the industrial revolution have 41 decreased surface ocean pH by 0.12 units (Riebesell, 2004). The "business as usual" scenario predicts that CO₂ will rise to 700 ppm over the next 100 years (Houghton, 2001), which will 42 43 decrease seawater pH by a further 0.3 to 0.6 units (Riebesell, Schulz, Bellerby, Botros, 44 Fritsche et al., 2007), bicarbonate ion (HCO₃⁻) concentration by 50% (Riebesell, 2004) and 45 raise the sea surface temperature (SST) by 2 to 6°C (Alley, Berntsen, Bindoff, Chen & others, 46 2007; Bopp, Monfray, Aumont, Dufresne, Le Treut et al., 2001). The projected pH shift from 47 8.2 to 7.7 covers the entire range of variation in pH currently observed in open ocean surface 48 waters. The increase in SST is predicted to increase stratification and light availability of the 49 surface ocean due to shoaling of the upper mixed layer (Rost, Zondervan & Wolf-Gladrow, 2008; Sarmiento, Slater, Barber, Bopp, Doney et al., 2004), which could directly impact the 50 51 physiology of phytoplankton. These changes in pH, SST and light regime are predicted to 52 enhance primary production (PP) (Hein & SandJensen, 1997; Riebesell, Wolfgladrow & 53 Smetacek, 1993), especially in the North Atlantic (Bopp et al., 2001; Doney, Lindsay, Fung & 54 John, 2006; Feng, Hare, Leblanc, Rose, Zhang et al., 2009), though this will depend on the 55 associated dominant species and nutrient regime (Rost et al., 2008). 56 Information on the effects of CO₂ enrichment on phytoplankton photosynthesis in a range of organisms and different trophic systems is essential to understand potential shifts in 57 58 the carbon cycle due to ocean acidification (Riebesell, 2004). Within the usual seawater pH 59 range (8.0-8.3), 90% of the total DIC is HCO_3^- , and CO_2 is less than 1 % when the system is 60 in equilibrium with atmospheric CO_2 (Skirrow, 1975). The total inorganic carbon

There is global concern over increases in the CO₂ concentration of seawater and the effects of

61 concentration in seawater is 2 μ mol L⁻¹ and the CO₂ content is 10 μ mol L⁻¹. This is not

62 sufficient to saturate carbon fixation by the algal photosynthetic enzyme ribulose bisphosphate carboxylase oxygenase (RuBisCO), which has half-saturation constants of 20-63 40 μ mol L⁻¹ CO₂ for eukaryotic microalgae and up to 750 μ mol L⁻¹ CO₂ for marine 64 cyanobacteria (Badger, Andrews, Whitney, Ludwig, Yellowlees et al., 1998; Hopkinson, 65 Dupont, Allen & Morel, 2011; Raven, 2011a; Raven & Johnston, 1991). Different 66 67 phytoplankton groups or species have therefore evolved a preference for different forms of 68 DIC, with some taking up CO_2 directly, whereas others draw on the pool of HCO_3 present 69 (Elzenga, Prins & Stefels, 2000) and or mechanisms to concentrate CO₂ or HCO₃. Sensitivity 70 to CO_2 therefore varies in relation to the HCO_3^- : CO_2 preference and the affinity of 71 phytoplankton for carbon fixation. Some species can rapidly acclimate to changes in the 72 concentration of dissolved CO₂, or total DIC (Nimer & Merrett, 1996). From an ecosystem 73 perspective, it has been suggested that the responses of phytoplankton to reduced pH in a high 74 CO₂ ocean are likely to be species-specific, with potential 'winners' and 'losers' (Hinga, 2002). 75

76 There is a growing body of literature on changes in photosynthesis due to increases in 77 CO₂ and related effects (Brading, Warner, Davey, Smith, Achterberg et al., 2011; Feng et al., 78 2009; Feng, Warner, Zhang, Sun, Fu et al., 2008; Flynn, Clark, Mitra, Fabian, Hansen et al., 79 2015). The majority of studies have been conducted on phytoplankton in laboratory culture 80 rather than on natural samples (Feng et al., 2008; Fu, Warner, Zhang, Feng & Hutchins, 2007; 81 King, Jenkins, Wallace, Liu, Wikfors et al., 2015; Shi, Li, Hopkinson, Hong, Li et al., 2015; 82 Wu, Gao & Riebesell, 2010). In diatoms, some cyanobacteria and coccolithophorids, elevated 83 CO₂ can lead to an increase in photosynthesis especially in large chain forming diatoms 84 (Tortell & Morel, 2002; Tortell, Payne, Li, Trimborn, Rost et al., 2008), Synechococcus spp. 85 (Fu et al., 2007) and *Emiliania huxleyi* (Leonardos & Geider, 2005). In the diatom 86 Chaetoceros muelleri for example, it has been shown that under saturating irradiance,

87 maximal photosynthetic rates are stimulated by increasing CO₂ availability (Ihnken, Roberts

88 & Beardall, 2011). For *Phaeodactylum tricornutum* grown at elevated CO₂ (1000 ppm)

89 corresponding to 7.8 pH, there was greater photoinhibition of the electron transport rate from

- 90 photosystem II (PSII) under high irradiance, whereas non-photochemical quenching was
- 91 reduced compared to low CO_2 grown cells (Wu et al., 2010).

92 Most studies on natural samples in the Atlantic Ocean have been conducted in 93 eutrophic and mesotrophic environments of the North Atlantic during natural or simulated 94 blooms. Some of these report an increase in the abundance of diatoms compared to 95 Phaeocystis spp. under CO₂ enrichment (Feng et al., 2008; Riebesell et al., 2007; Tortell et 96 al., 2008), others report that nano-phytoplankton or Prymnesiophytes replace diatoms under 97 elevated CO₂ and temperature in the Bering Sea (Hare, Leblanc, DiTullio, Kudela, Zhang et 98 al., 2007) and the North Atlantic (Feng et al., 2008), where there was a change from diatoms 99 to Prymnesiophytes under elevated CO₂ and a decrease in inorganic carbon production 100 (calcification). There have been few studies of CO₂ enrichment on changes in phytoplankton 101 community structure and photosynthesis in the oligotrophic gyres (Egleston, Sabine & Morel, 102 2010), even though they occupy the largest areas of the ocean. To date there has only been 103 one study conducted in the sub-tropical North Atlantic at the Bermuda Atlantic Time Series 104 site (BATS), which showed no significant difference in carbon fixation rates at elevated pCO_2 105 (reduced pH) for phytoplankton assemblages dominated by Prochlorococcus sp. and 106 Synechococcus sp. (Lomas, Hopkinson, Losh, Ryan, Shi et al., 2012). By contrast, in the same 107 study nitrogen fixation rates in colonies of Trichodesmium increased by 54 % at pH 7.8 but 108 decreased by 21 % at pH 8.4.

109 The objective of this study was to assess the effect of CO_2 enrichment on

110 phytoplankton photosynthesis and community composition in the North Atlantic sub-tropical

111 gyre through a series of shipboard incubation experiments. Low temperature (LT) emission

112 spectra were used to assess changes in major pigment-protein complexes in the oxygenic

113 photosynthetic membranes. The LT spectra also provided information about the presence of

114 chlorophyll-containing light harvesting antenna complexes and their functional coupling to

115 photosystem reaction centres.

116

117 **2. Methods**

118 2.1. Study area and experimental design

119 Shipboard incubations were conducted aboard the RRS James Cook between 13 October and 120 21 November 2010 during the Atlantic Meridional Transect 20 (Cruise JC053). CO2 121 enrichment experiments were carried out at three stations in the North Atlantic Gyre at 29°N, 122 34°W for experiment (Exp.) I; 18°N, 37°W for Exp. II and 7°N, 30°W for Exp. III (Fig. 1). 123 For each Exp., 80 L of near surface seawater was collected from the ship's underway supply 124 before dawn into a large Nalgene container. The water was sub-sampled into 18 acid cleaned 125 (IOC, 1994) 4 L clear polycarbonate bottles for incubation, leaving a head space of 0.5 L. 126 Nine of the bottles were bubbled with pre-mixed synthetic air with 360 ppm CO₂ (control) 127 and the other nine were bubbled with synthetic air with 760 ppm CO₂ (CO₂ treatment). The 128 gases were supplied to the experimental bottles from gas cylinders via nylon tubing fitted with 129 in-line 0.2 µm sterile Acrovent filters, to prevent contamination by particulates. The nylon 130 tubing entered the bottles through Nalgene caps with Kinesis Omni-Lok fittings and vent 131 tubes to prevent the build-up of pressure in the bottles. To minimise the effect of bubbling on 132 phytoplankton, each experimental bottle was bubbled for an initial 8 h period followed by a 133 further 4 h after 24 h to maintain the CO₂ over the 48 h period. The incubations were 134 conducted in an on-deck incubation system (78 x 60 x 68 cm) supplied with flow through sub-135 surface seawater from the ship's underway supply to maintain the bottles at ambient 136 temperature. The incubation system had no light screen, but the sides of the incubator were opaque which provided shading either side of zenith. pH, alkalinity, pCO₂, HCO₃, and Chl *a*were measured in triplicate at 0 (T0), 6, (T6; except Chl *a*), 12 (T12), 24 (T24) and 48 (T48)
h. Samples for pico- and nanophytoplankton enumeration by flow cytometry, photosynthesisirradiance curves and low emission spectra were measured in triplicate at T0, T24 and T48.
Microscopy counts were made as single measurements at T0 and T48.

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143 2.2. Carbonate chemistry: pH, pCO₂ and alkalinity

pH was determined spectrophotometrically onboard using 2 μ mol L⁻¹ of m-cresol from a 2 144 mmol L^{-1} stock solution which was prepared from pure sodium salt (Sigma-Aldrich, USA). 145 146 This method expresses pH on the total hydrogen concentration scale with 0.01 precision 147 (Dickson, Afghan & Anderson, 2003). Absorbance was recorded before and after addition of 148 m-cresol at 434, 578 and 730 nm on a PerkinElmer Lambda 35 spectrophotometer in a 100 149 mm cuvette. Seawater temperature was measured using an Amarell Ama-Digit 15 150 Temperature probe, and salinity was taken from a Seabird Electronics CTD calibrated against 151 salinity standards. For total alkalinity, samples were stored in 100 mL borosilicate dark bottles 152 with teflon cap liners spiked with 5% mercuric chloride (6.9 nmol L^{-1}). Alkalinity was 153 measured 2 months after the cruise using an automatic Apollo SciTech, Alkalinity Titrator 154 (Model AS-ALK2) in 0.8 M H₂SO₄. pCO₂ was calculated from the pH, temperature, salinity 155 and alkalinity measurements using CO2SYS software (Pierrot & Wallace, 2006), using the 156 constants set according to Mehrbach et al. (1973) and Dickson and Millero (1987) and 157 corrected for differences between *in situ* and measured temperature.

158

159 2.3. Phytoplankton community structure

160 Nano and picoeukaryote phytoplankton cells from approx. 0.2 to 10 µm were enumerated

161 using a Becton Dickinson FACSortTM flow cytometer (Becton Dickinson, USA) equipped

162 with an air-cooled laser providing blue light at 488 nm, using the methods given in Tarran et 163 al. (Tarran, Heywood & Zubkov, 2006; Tarran, Zubkov, Sleigh, Burkill & Yallop, 2001). 164 Picoeukaryotic phytoplankton and nanophytoplankton were analysed for between 2-4 minutes 165 to measure chlorophyll fluorescence (>650 nm), phycoerythrin fluorescence (585 ± 21 nm) and side scatter (light scattered at 90° to the plane of the laser). Using a combination of 166 167 density plots as described in Tarran et al. (2001), it was possible to identify and quantify the 168 following planktonic groups in the samples: *Prochlorococcus* spp., *Synechococcus* spp., 169 picoeukaryotic phytoplankton (approx. 0.6 to 3 µm, including prasinophyceae, chlorophyceae, 170 pelagophyceae) and nanoeukaryotic phytoplankton (approx. 3 to 10 µm, including 171 cryptophyceaeae and prymnesiophyceae). Instrument flow rate was calibrated daily using 172 Beckman Coulter Flowset fluorospheres of known concentration. Carbon biomass was 173 calculated using median cell diameters measured for each group using the methods described 174 in Tarran et al. (2006) and the carbon conversion factors given in (Bjørnsen, 1986) and 175 (Booth, 1988).

176 Microscopy was used to enumerate diatoms and dinoflagellates. For microscopic 177 counts, 100 mL of seawater samples were collected into brown borosilicate bottles and 178 preserved in Lugol's iodine (final concentration of 1 %) for stations II and III. Dinoflagellates 179 and diatoms were counted 2 months after the cruise using a Leica DM IRB microscope 180 (Leica, Germany). 100 mL of sample was settled in composite sedimentation chambers for 24 181 h. Samples were counted on 26 mm diameter sedimentation slides in replicate vertical and 182 horizontal transects using a x 20 objective. Cell volumes for these groups were calculated 183 using approximate geometric shapes and converted to biomass using the equations given in 184 Menden-Deuer and Lessard (2000).

185 For analysis of Chlorophyll-*a* (Chl *a*), 100 mL samples were filtered onto 25 mm 186 GF/F filters (pore size 0.7 μ m, Whatman). The filter was then placed in a 15 mL FalconTM 187 tube (BD, UK) and 10 mL of 90 % acetone was added. The tubes were then stored in the dark,

188 in a -20°C freezer for 12 h. Chl *a* was then measured onboard fluorometrically following

Holm-Hansen, Lorenzen, Holmes and Strickland (1965) using a Turner Designs, Trilogy
Fluorometer.

- 191
- 192 2.4. Phytoplankton photosynthesis
- 193 2.4.1. Photosynthesis-irradiance curves

194 Photosynthetic-irradiance (P-E) experiments were conducted in linear photosynthetrons 195 illuminated with 50 W, 12 V tungsten halogen lamps following the methods described in 196 Tilstone, Figueiras, Lorenzo and Arbones (2003). Each incubator housed 16 sub-samples in 197 60 mL acid-cleaned polycarbonate bottles, which were inoculated with between 185 kBq (5 μ Ci) and 370 kBg (10 μ Ci) of ¹⁴C labelled bicarbonate. The PE curves were conducted at the 198 199 same time each day to minimise the influence of diel light history. PAR in the incubator was 200 set to ambient levels measured over a 2 h period prior to incubation using a Skye Instruments PAR Sensor (model SKP 200). For Exps I, II and IIII these were 1319, 2166 and 2215 µE m⁻² 201 202 s⁻¹, respectively. The P-E curves were maintained at *in situ* temperature using the ship's non-203 toxic seawater supply. After 1.5 h incubation, the suspended material was filtered through 25 204 mm GF/F filters which were then exposed to 37% hydrochloric acid fumes for 12 h and then immersed in scintillation cocktail. ¹⁴C disintegration time per minute (DPM) was measured 205 206 using the onboard 1414 liquid scintillation counter (PerkinElmer, USA) and the external standard and the channel ratio methods to correct for quenching. Dark ¹⁴C uptake was 207 208 subtracted from light uptake in the other 15 light incubation cells. Photosynthetic rates were 209 calculated from total dissolved inorganic carbon (DIC) and Chl a. DIC was calculated from salinity and alkalinity. The broadband light-saturated Chla-specific rate of photosynthesis, P_m^B 210 [mg C (mg Chla)⁻¹ h⁻¹], and the light-limited slope, α^{B} [mg C (mg Chla)⁻¹ h⁻¹ (µmol m⁻² s⁻¹)⁻¹ 211

¹], were estimated by normalising ¹⁴C uptake to Chl *a* and fitting the data to the model of Platt et al. (1980). Primary production (mg C m⁻³ d⁻¹) was subsequently calculated from photosynthetically active radiation (E_{PAR}), Chl *a* and the photosynthetic parameters using the brand model of Tilstone et al. (2003). E_{PAR} was modelled using the approach of Gregg & Carder (1990) modified to include the effects of clouds (Reed, 1977) and using wind speed and percentage cloud cover from the European Centre for Medium Range Weather Forecasting (ECMWF) following Smyth et al. (2005).

219

220 2.4.2. Low temperature fluorescence emission spectra

221 Samples were filtered onto 25 mm GF/F filters from ~1 L of sea water and processed

following standard protocols (Suggett, Stambler, Prasil, Kolber, Quigg et al., 2009). A 4 x 10

223 mm piece of the filter was cut and placed onto a holder, flash frozen in liquid nitrogen and

224 measured immediately using a SM-9000 spectrometer (Photon Systems Instruments, Brno).

225 The instrument is based on a Zeiss MCS CCD spectrometer; the detector has 1044 pixels, the

grating images from 200-980 nm, the wavelength accuracy is 0.5 nm and the spectral

resolution is 3.5 nm (FWHM). Two gaussian shaped lines are separated at >3.5nm, with a

resolution of 3.2 nm from 0.8nm dispersion by 4 pixels. Whole fluorescence emission spectra

229 were determined using an integration time of 1000 ms. The fluorescence emission of blanks

230 (seawater pre-filtered through a 0.7 μm GF/F) was subtracted and the spectra were normalized

- 231 to 686 nm for de-convolution (Kaňa, Kotabová, Komárek, Papageorgiou, Govindjee et al.,
- **232 2012**).

233

234 2.5. CTD and Nutrients.

A Seabird 911 plus CTD was deployed at each station at which experiments were conducted.

The data was processed using SeaBird software v7.21 and up and down casts were then

237	merged to 1 m binned resolution. Micro-molar nutrients were analysed using a 5 channel
238	nitrate (Brewer & Riley, 1965), nitrite (Grasshoff, 1976), phosphate, silicate (Kirkwood,
239	1989) and ammonia (Mantoura & Woodward, 1983) Bran & Luebbe AAIII segmented flow,
240	colourimetric, autoanalyser. Water samples were taken from a 24 x 20 litre bottle stainless
241	steel framed CTD / Rosette system (Seabird). These were sub-sampled into clean (acid-
242	washed) 60 mL HDPE (Nalgene) sample bottles. Subsequent nutrient analysis was complete
243	within 1-2 hours of sampling. Clean handling techniques were employed to avoid
244	contamination of the samples.
245	
246	2.6. Statistical analysis
247	Paired T-tests samples were employed to test for significant differences between
248	treatments at T48 on individual Exp.'s and for all Exp.'s. Results of T-test analyses from all
249	Exp.'s (I-III) at 48 hrs are given in Table 2. Results of T-test analyses from individual Exp.'s
250	at 48 hrs are given on each sub-figure. The T-test results are given as $T_{1,19} = x$ and $P = y$
251	where T is the deviation of the sample mean from the normally distributed parametric mean to

parametric standard deviation ratio, the sub-script numbers (1, 23) denote the degrees offreedom and P is the T-test critical significance value.

254

255 **3. Results**

256 3.1. Initial hydrographic conditions

Temperature varied from 26 °C to 29 °C in the surface waters of the sub-tropical Atlantic (Figs. 1, 2). These stations were typically oligotrophic with a deep thermocline at between 50 and 75 m, low nitrate and nitrite (<0.02 μ M), low surface Chl *a* (0.03-0.05 mg m⁻³) and deep Chl *a* maxima between 70 and 130 m of between 0.1 and 0.3 mg m⁻³ (Fig. 2B, C, D). The 261 concentrations of nitrate+nitrite and silicate in the surface waters were beyond the detection
262 limit (Harris, 2011).

- 263
- 264 *3.2. Changes in carbonate chemistry*

265 For all experiments, by 48 h the CO₂ treatment resulted in an average concentration of 748 266 ppm and 7.97 pH units compared to 468 ppm and 8.14 pH units in the control (Fig. 3). For the 267 control, bubbling pre-mixed synthetic air with a CO₂ concentration of 360 ppm to natural 268 seawater, resulted in > 460 ppm (Fig. 3). At 48 h the difference in CO_2 content and pH 269 between the CO₂ treatment and control was ~280 ppm and 0.17 pH units, respectively (Fig. 270 3). Similarly, the mean HCO_3^- in the CO_2 treatment during these Exp.'s was 2013 ppm and 271 1860 ppm in the control (Fig. 4A, B, C) and the mean HCO₃⁻: CO₂ ratio was 2.7 in the CO₂ 272 treatment and 3.99 in the control (Fig. 4D, E, F).

273 The corresponding changes in total carbon (TC) and alkalinity (TA) during each experiment 274 are given in Table 1. For all Exp.'s, the TA in the control and CO₂ enrichment was similar and 275 consequently there was no significant difference in TA for Exp. II & III and all Exp.'s 276 together (Table 2). There was a significant difference in TA between CO₂ and control at T48 277 for Exp. I (T = 55.25, P = 0.012). By contrast, TC was significantly higher in the CO_2 278 treatment during Exp. II (T = 4.42, P = 0.048) & III (T = 7.08, P = 0.019) and all Exp.'s 279 together (Table 2), but this was not significant for Exp. I. Carbonate chemistry in the bottles 280 equilibrated at between 12 & 24 h. For all Exp.'s, there was a significant difference in CO₂, pH, HCO₃⁻ and HCO₃⁻ : CO₂ TC, TA, (Table 2) between control and CO₂ treatments. At T48 281 282 in the CO_2 treatment, the CO_2 concentration was significantly higher than at T0 (Figure 3; 283 Table 2).

During Exp.'s I & II there was no significant difference in Chl a between CO₂ and 286 287 control treatments (Fig. 5) and as a consequence, for all Exp.'s. there was no significant 288 difference between treatments and with T0 (Table 2). In Exp. III, Chl a was significantly 289 higher in the control (Fig. 5C). Of the phytoplankton groups enumerated by flow cytometry, 290 nanoeukaryotes had the highest biomass in Exp.'s I and II, whereas in Exp. III the biomass of 291 nano- and picoeukaryotes and Synechococcus spp. were similar (Fig. 6, 7). During all Exp.'s, 292 there was no difference in nanoeukaryotes and picoeukaryotes between control and CO₂ 293 treatments (Table 2). During Exp. I, there was little change in the biomass of nanoeukaryotes 294 and picoeukaryotes over the duration of the Exp.'s and there was no significant difference 295 between control and CO₂ treatment (Fig. 6A, D). During Exp. II, the initial biomass of 296 nanoeukaryotes and picoeukaryotes decreased at 24 h followed by a slight increase at 48 h, 297 but there was no significant difference in biomass between control and CO₂ treatments at 48 h 298 (Fig. 6B, E). In Exp. III, the biomass of these groups initially decreased, but at 48 h both 299 nanoeukaryotes and picoeukaryotes were significantly higher in the control compared to the 300 CO₂ treatments and TO (Fig. 6C, F). During all experiments there was no significant 301 difference between control and CO₂ treatments for Synechococcus spp. and Prochlorococcus 302 spp. (Fig. 7, Table 2). The biomass of Prochlorococcus spp. and Synechococcus spp. 303 decreased from T0 to T48 and T0 was significantly higher compared to the control and the 304 CO₂ treatment (Fig. 7, Table 2). For individual Exp.'s, there was no significant difference 305 between treatments for both Synechococcus spp. and Prochlorococcus spp. in Exp. I, II & III 306 (Fig. 7).

307 From the microscopy samples, there was no significant difference in diatoms biomass 308 between treatments during Exp.'s II and III (Table 2). The total biomass in Exp.'s II was low 309 3.5 and 4.5 μ g C L⁻¹ at 48 h in CO₂ and control treatments, respectively when *Navicula* spp. 310 accounted for the highest biomass. In Exp. II, the diatom biomass was higher and 17 and 18 μ g C L⁻¹ in CO₂ and control treatments, respectively when *Rhizosolenia* spp. and *Navicula* spp. dominated (Fig. 8A, B). In this Exp. *Navicula* spp. dominated the biomass in the CO₂ treatment, whereas in the control *Rhizosolenia* spp. biomass was higher than any other group (Fig. 8A, B). The biomass of diatoms was significantly higher in T0 compared to both the control and CO₂ treatments (Fig. 8A, B, Table 2). The diatom biomass may have been reduced through silicate limitation.

Dinoflagellates exhibited the highest biomass of all phytoplankton groups during Exp.'s II and III, reaching on average $60 \ \mu g \ C \ L^{-1}$ under CO₂ treatment and $30 \ \mu g \ C \ L^{-1}$ in the control, though the differences between treatments were not significant (Table 2). *Gymnodinium* spp. accounted for the majority of the biomass, especially in Exp. III (Fig. 8D). The dinoflagellate biomass was also significantly higher in the control at T0 compared to T48, but this was not the case for the CO₂ treatment (Fig. 8C, D, Table 2). T-test was not performed on individual Exp.'s, since only single samples were enumerated at T48.

324

325 3.4. Changes in photosynthesis and emission spectra

For all Exp.'s there was no significant difference in P_m^B between control and CO₂ treatment (Fig. 9A, B, C, Table 2). For the individual Exp.'s, there was no difference in P_m^B between control and CO₂ treatment in Exp. I (Fig. 9A), but P_m^B was significantly higher in the CO₂ treatment compared to the control in both Exp.'s II and III (Fig. 9B, C). In all Exp.'s, α^B was significantly higher in the CO₂ treatment compared to the control (Fig. 9D, E, F, Table 2). There was no significant difference in α^B in the control between T0 and T48, but α^B was significantly higher at T48 in the CO₂ treatment compared to T0 (Table 2).

Using spectral de-convolution of LT emission spectroscopy measurements, the fluorescence peak at 678 nm is associated with uncoupled chlorophyll-containing PS antenna, while 686 nm represents a mixture of signals from PS II chlorophylls and terminal emitters of 336 phycobilisomes (PBS) (Rakhimberdieva, Vavilin, Vermaas, Elanskaya & Karapetyan, 2007). 337 Generally, the peak area at 678 nm decreased by more than three times from T0 to T48 in all 338 Exp.'s and in both treatments (Fig. 10, Table 3). In all of the CO_2 treatments however, there 339 was a significantly lower signal at 678 nm (Table 3), indicating reversible antenna connection 340 after CO_2 enrichment. In addition, the spectral emission of the PSII core antenna increased 341 from T0 to T48 (Fig. 10).

- 342
- 343 **4. Discussion**

344 *4.1. Changes in the phytoplankton community in relation to carbonate chemistry*

345 Experiments on CO₂ enrichment of natural samples in the Atlantic Ocean have mostly 346 focused on the interactions between diatoms, Phaeocystis spp. and prymnesiophytes. The sub-347 tropical and tropical oligotrophic gyres are dominated by Cyanobacteria which make a 348 significant contribution to the carbon fixation in the global ocean (Bell & Kalff, 2001). In the 349 North Atlantic gyre, the phytoplankton assemblage at the deep Chl *a* maximum is typically 350 dominated by picoeukaryotes, Prochloroccocus spp. At the DCM and Synechococcus spp. in 351 the suface mixed layer (Tarran et al., 2006; Zubkov, Sleigh, Tarran, Burkill & Leakey, 1998), 352 which exist at very low (beyond detection limit) nitrate and phosphate concentrations. Under 353 climate change scenarios of a warming ocean, increased stratification is likely to reduce 354 nutrient concentrations in the sub-tropical gyres (Sarmiento et al., 2004). We conducted 355 experiments with elevated CO₂ in the North Atlantic sub-tropical gyre during autumn to 356 evaluate the effects on phytoplankton community structure and photosynthetic rates. The 357 Exp.'s were conducted with no nutrient addition to mimic the oligotrophic conditions of the 358 gyre. The temperature at the experimental stations was 26 to 28°C and typical of stratified 359 sub-tropical waters where irradiance is high. During the Exp.'s, the phytoplankton community 360 was dominated by dinoflagellates, which was 55 to 80 times higher than the other groups, but there was no significant difference between treatments and in some cases the biomassdecreased from T0 to T48 (Fig. 8).

363 Understanding the response of dinoflagellates to both increases in CO₂ and 364 temperature is key to detecting climate-driven perturbations in the ecosystem. In culture 365 experiments on autotrophic dinoflagellates, some strains of Alexandrium fundyense 366 (Hattenrath-Lehmann, Smith, Wallace, Merlo, Koch et al., 2015), Karenia brevis (Errera, Yvon-Lewis, Kessler & Campbell, 2014), Karlodinium veneficum (Fu, Place, Garcia & 367 368 Hutchins, 2010), Prorocentrum minimum and Heterosigma akashiwo (Fu, Zhang, Warner, 369 Feng, Sun et al., 2008) grow significantly faster at high pCO₂. Additionally, Alexandrium 370 fundyense experiences a significant increase in cell toxicity under elevated CO₂ (Hattenrath-371 Lehmann et al., 2015). There have been few studies on the response of Gymnodinium spp. and 372 Gyrodinium spp., which exhibited high biomass in this study. Most Gymnodinium spp. are 373 autotrophic, though some (e.g. G. abbreviatum, G. heterostriatum) are known heterotrophs 374 (Tomas, 1996). Similarly many Gyrodinium spp. are autotrophic, though some (e.g. G. 375 lachryma, G. pingue, G. spirale) are heterotrophic (Tomas, 1996), so the patterns shown in 376 Figures 8C, D may also include some heterotrophs as we were not able to identify all 377 Gymnodinium spp. and Gyrodinium spp. to species level. Calbet, Sazhin, Nejstgaard, Berger, 378 Tait et al. (2014) studied the response of natural phytoplankton communities in a Norwegian 379 Fjord using mesocosm experiments and found at elevated temperature and lower pH there was 380 no difference in Gyrodinium spp. abundance. Wynn-Edwards, King, Davidson, Wright, 381 Nichols et al. (2014) found that in continuous batch culture of Antarctic strains of 382 Gymnodinium spp. that there was no difference in growth rates and cell size between CO₂ 383 enrichment to 993 ppm and ambient, but there was a significantly higher cell polyunsaturated 384 fatty acid content at the higher CO₂ concentrations.

385 In our Exp.'s Diatoms had the second highest biomass, but there was no difference 386 between the elevated CO₂ treatment and ambient (Fig. 8, Table 2). In other studies, the growth 387 rates of the diatoms Skeletonema costatum (Grev.) Cleve and Chaetoceros spp. increased at 388 elevated CO₂ by 100% and 24%, respectively (Beardall & Raven, 2004; Kim, Lee, Shin, 389 Kang, Lee et al., 2006). Prochlorococcus spp. exhibited the third highest biomass, however 390 there was also no significant difference in the biomass of *Prochlorococcus* spp., and that of 391 Synechococcus spp., picoeukaryotes, and nanoeukaryotes except in one Exp. III, in which 392 there was a higher biomass of both picoeukaryotes and nanoeukaryote in the control (Fig. 6). 393 The response of Synechococcus spp. in our experiments on natural samples contrasts the 394 trends observed in culture experiments. For example, Fu et al. (2007) found that 750 ppm CO₂ 395 stimulated the growth rate of Synechococcus spp. The different responses between 396 Synechococcus spp. and Prochlorococcus spp. were thought to be due to differences in inorganic carbon acquisition systems associated with carbon limitation at low pCO2 in 397 398 Synechococcus spp. (Fu et al. 2007). In contrast, we observed no difference in biomass 399 between treatments under limiting nutrients. Similarly in natural phytoplankton communities 400 Lomas et al. (2012) found that under replete P and Fe there was no significant response in the 401 carbon-fixation of Prochlorococcus and Synechococcus spp. dominated communities to elevated CO₂ and a decrease in pH. In their study, pH was adjusted by acid-base manipulation 402 403 (rather than direct enrichment with CO₂) and nutrient media were added to the incubations 404 even though nitrate and phosphate concentrations were below detection on first collection of 405 the samples. By contrast, in mesocosm exp.'s using natural phytoplankton communities with 406 replete nutrient concentrations, Paulino, Egge and Larsen (2008) reported a lower abundance 407 of Synechococcus spp. and a higher abundance of picoeukaryotes at high CO₂. By contrast, 408 Low-Decarie, Fussmann and Bell (2011) found in experimental mesocosms that increases in 409 CO₂ enhanced the competitive ability of chlorophytes relative to cyanobacteria. Similarly in 410 the North Atlantic, Feng et al. (2009) suggested that nanophytoplankton (coccolithophores) 411 may further increase in abundance relative to other phytoplankton groups in the later stages of 412 the spring bloom if future CO_2 concentrations and sea temperature continue to rise.

413 During our Exp.'s, dinoflagellates, diatoms and Prochlorococcus spp. decreased from 414 the initial T0 biomass, which may in part be due to the nitrate and silicate limitation or 415 grazing by heterotrophic and / or mixotrophic dinoflagellates, through confinement in bottles. 416 Nitrate and nitrite were low prior to confinement in all Exp.'s (Fig. 2) and silicate was beyond 417 the detection limit. The limited silicate availability meant that biomass of diatoms was low, 418 though it did increase in Exp. III. The biomass at TO suggests that the cyanobacteria were 419 adversely affected by containment, which may have allowed the other groups to thrive due to 420 lack of competition and / or the recycled nutrients released during the decline of the 421 cyanobacteria biomass. Despite the initial decline, there was no clear response to elevated 422 CO₂ in the phytoplankton assemblage. Most studies on changes in the phytoplankton 423 community to elevated CO₂ report significant shifts or responses, however this may provide a 424 biased view since non-significant responses are rarely published (Lomas et al., 2012).

425

426 4.2. Effects of CO₂ enrichment on phytoplankton photosynthesis and primary production.

427 Aquatic photosynthesis is inherently under-saturated with respect to CO₂ (Tortell, Rau 428 & Morel, 2000), therefore increases in seawater CO₂ can enhance photosynthetic carbon 429 fixation (Riebesell, 2004). Some studies have observed a 40% increase in the uptake of DIC at 430 elevated CO₂ compared to present levels (Riebesell et al., 2007). It has been suggested that 431 the photosynthesis of dinoflagellates may be limited by current water column CO₂ 432 concentrations and that they may benefit from rising atmospheric pCO₂ (Eberlein, de Waal & 433 Rost, 2014). There are a number of studies that contradict this hypothesis, for example, in 434 dilute batch cultures of the calcareous dinoflagellate Scrippsiella trochoidea and the toxic 435 dinoflagellate Alexandrium tamarense grown at a range of CO₂ concentrations (180-1200 436 ppm), there was no significant difference in photosynthetic and growth rates between 437 treatments, though A. tamarense did exhibit greater respiration rates at higher pCO₂ (Eberlein 438 et al., 2014). In our experiments, photosynthetic rates were significantly greater in the CO_2 439 enriched treatment (Fig. 9) and the dinoflagellates were probably able to maximise carbon 440 fixation due to lack of competition, since the biomass of the other groups was low (Fig. 6, 7, 441 8A, B). In Exp.'s II and III, when the dinoflagellates Gymnodinium and Gyrodinium spp. dominated the biomass, significantly higher P_m^B and α^B at elevated CO₂ were associated with 442 a lower signal of unconnected PS antennae, suggesting that these previously unconnected PS 443 444 antennae become connected to facilitate the higher photosynthetic rates. In addition, there was 445 also a slight increase in fluorescence signal at 686 nm, due to a lower abundance of 446 Prochlorococcus spp. in the CO₂ treatment compared to the control, since this group do not 447 have PBS antennae (Partensky, Hess & Vaulot, 1999).

448 Photosynthetic carbon fixation is mediated by both pH and temperature dependent 449 enzymatic reactions via the Calvin-Benson Cycle (Giordano, Beardall & Raven, 2005; Portis 450 & Parry, 2007). Varying climate change scenarios of increased CO₂, temperature and a 451 decrease in pH may affect intra-cellular DIC uptake differently in different phytoplankton groups or species (Badger et al., 1998). P_m^B and α^B and temperature are tightly coupled (Uitz, 452 453 Huot, Bruyant, Babin & Claustre, 2008), and different phytoplankton species are adapted to a 454 narrow temperature band with sharp decrease in photosynthetic rates beyond 20°C (Eppley, 455 1972; Raven & Geider, 1988) which can be modified through the synthesis of photoprotecting 456 rather than photosynethetic pigments (Kiefer & Mitchell, 1983). The enzyme that is primarily 457 responsible for photosynthetic carbon fixation is Ribulose Bisphosphate Carboxylase 458 (RuBisCO), though this does have some catalytic limitations (Giordano et al., 2005). 459 Phytoplankton have therefore adapted two strategies to maximise the performance of

RuBisCO; by either altering the affinity of RuBisCO or through CO₂-concentrating 460 461 mechanisms (CCMs). CCMs have a higher affinity for inorganic carbon (Giordano et al., 462 2005; Raven, 2011a; Raven, 2011b; Reinfelder, 2011; Tortell et al., 2000), and through a 463 combination of RuBisCO activity or CCM, most species achieve photosynthetic saturation at 464 environmental CO₂ levels under light and nutrient replete conditions (Raven, 1997; Raven & 465 Johnston, 1991). The differences in photosynthetic efficiency and rates between different 466 phytoplankton groups can be partially explained by CCM and / or RuBisCO kinetics (Raven, 467 1997). The effect of an increase in CO_2 and decrease in pH may select phytoplankton species 468 or genotypes based on the efficiency of RuBisCO or CCM. Dinoflagellates possess Form II 469 RubisCO (Morse, Salois, Markovic & Hastings, 1995), which has low kinetic properties and 470 therefore a low photosynthetic efficiency (Tortell et al., 2000). They are abundant in low 471 nutrient oceanic waters (Margalef, 1979), have low growth rates attributed to their high basal 472 respiration rates (Tang, 1996), do not need to compete for their position in the ecosystem and 473 therefore rely on diffusive CO_2 uptake (Giordano et al., 2005). Dinoflagellates possess less 474 efficient CCMs and/or low affinities for CO₂, and may therefore benefit from living in a high 475 CO₂ world (Reinfelder 2011; Fu et al. 2012) and be highly sensitive to CO₂ (Rost et al., 476 2008). Form II RUBISCO has a much lower selectivity for CO₂ over O₂ (Giordano et al., 477 2005). To compensate for this, some marine dinoflagellates, such as *Prorocentrum minimum*, 478 Heterocapsa triquetra, Ceratium lineatum, Protoceratium reticulatum and Peridinium 479 gatunense have a CCM linked to carbonic anhydrases (CAs), the expression of which changes 480 in response to the external CO₂ concentration (Berman-Frank, Erez & Kaplan, 1998; Brading 481 et al., 2011; Ratti, Giordano & Morse, 2007; Rost, Riebesell & Sultemeyer, 2006). This 482 suggests that an increase in CO₂ stimulated efficiency in light absorption and utilization, 483 which resulted in the higher rates of carbon fixation shown in Figure 9. Using spectral de-484 convolution of low temperature emission spectroscopy measurements, the peak at 691 nm is

485 emitted from active PS II chlorophylls and 711 nm from PS I chlorophylls (Kaňa et al., 2012). 486 The fluorescence peak at 678 nm was assigned to uncoupled chlorophyll-containing PS 487 antenna, while 686 nm represents a mixture of signals from PS II chlorophylls and terminal 488 emitters of phycobilisomes (PBS) (Rakhimberdieva et al., 2007). The peak at 686 nm 489 increased by a factor of two in all Exp.'s, which indicates either an increase in PBS content or 490 the presence of the iron-stress protein, IsiA (Chauhan, Folea, Jolley, Kouril, Lubner et al., 491 2011). The lower signal of unconnected PS antennae in our Exp.'s maybe indicative of CCM 492 functionality as the dominant dinoflagellates connect more antennae to PS II to achieve the higher P_m^B and α^B rates. The PS I chlorophyll peak at 711 nm remained constant and there 493 494 was no obvious difference between treatments (Fig. 10, Table 2). Although semi-quantitative, 495 the advantage of LT spectroscopy is that when it is applied to time series and bioassays, 496 gradual changes in phytoplankton photochemical processes can be monitored. This was the 497 case in the Exp.'s on this cruise, where more detailed biochemical or biophysical analyses 498 cannot usually be performed. Further work is required to ascertain whether the higher 499 photosynethetic rates we observed in Gymnodinium, and Gyrodinium spp. under CO₂ 500 enrichment is a species or genotypic response. This response also needs to be further 501 invetigated under different light conditions. For example, fluctuating light can affect 502 dinoflagellate growth and photosynthesis with constant light and high CO₂ having a negative 503 effect on the growth and quantum yield of Prorocentrum micans (Zheng, Giordano & Gao, 504 2015), which has implications for ocean warming and high stratification in a high CO_2 world. 505 Cyanobacteria and chlorophytes possess Form IB RuBisCO (Badger & Price, 2003). 506 Cyanobacteria utilize active HCO_3^- and CO_2 pumps as their CCM to facilitate CO_2 fixation 507 and maintain rapid growth at low external DIC concentrations (Badger & Price, 2003; Badger, 508 Price, Long & Woodger, 2006). In Exp. III, Prochlorococcus sp. biomass was significantly

509 higher in the control, possibly as a result of their high affinity for CO_2 uptake at low

510 concentrations or grazing of *Prochlorococcus* sp. biomass by heterotrophs or mixotrophs (Fig. 511 3D), but the photosynthetic rates in the control were significantly lower than the CO_2 512 treatment, principally because the dinoflagellate biomass was far higher in this Exp. By 513 comparison, Fu et al. (2007) also observed that the photosynthetic rate and light harvesting 514 efficiency of Prochlorococcus sp. in vivo was not affected by increasing CO₂. Similar to the 515 patterns we saw in Exp.'s II & III, Fu et al. (2007) also observed only slight differences in 516 growth and photosynthetic rates of Synechococcus spp. at 750 ppm CO₂ and control, though 517 this changed when the temperature was elevated by 4 °C. By comparison, Lomas et al. (2012) 518 observed that natural phytoplankton assemblages dominated by cyanobacteria were able to 519 adapt to changes in pH through modification of the assimilation number, which resulted in no 520 significant differences in photosynthetic rates. We found in two of our Exp.'s that the 521 assimilation number was significantly higher at elevated CO₂. The duration of our 522 experiments represent short term responses to CO₂ enrichment and do not account for longer 523 term adaptation and evolution, which can be complex (Collins & Bell, 2006). Our 524 experiments also do not account for interactive synergies between CO₂, temperature, nutrients 525 and light which may affect phytoplankton, as shown by other studies in the North Atlantic 526 (Feng et al., 2009; Hare et al., 2007). Further work is required to elucidate these dual effects 527 on phytoplankton community structure in the North Atlantic sub-tropical gyre.

528 What is the consequence of increases in CO_2 on water column integrated primary 529 production? Some studies have reported no significant changes in primary production under 530 elevated CO_2 (Goldman, 1999; Raven & Falkowski, 1999; Tortell & Morel, 2002; Tortell et 531 al., 2000). We found in the North Atlantic sub-tropical gyre when dinoflagellates dominated, 532 that there was a 25% increase in primary production over all Exp.'s at 760ppm CO_2 and that 533 this was significant in Exp.'s II & III (Fig. 11). This is similar to the findings of Hein and SandJensen (1997) who suggested that increasing CO₂ concentrations in seawater would
increase primary production by 15 to 19%.

536

537 **5. Conclusions**

538 Three CO₂ enrichment experiments were conducted on natural phytoplankton 539 assemblages in the North Atlantic sub-tropical gyre. The dinoflagellates, Gymnodinium, and 540 Gyrodinium spp., dominated the biomass but there was no significant difference in the 541 biomass of these and other phytoplankton groups at elevated CO₂ concentrations of 760 ppm 542 and 7.94 pH compared to ambient (468 ppm CO₂ and 8.33 pH). There were however, 543 significantly higher photosynthetic rates in the elevated CO₂ treatment which was due to the 544 connection of reversible photosystem antennae at higher CO₂ concentrations, which resulted 545 in a 25% increase primary production.

546

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- 560
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- 817
- 818 Figure Legends.
- 819 Fig. 1. Stations in the North Atlantic at which CO₂ enrichment experiments were conducted
- 820 overlaid on MODIS-aqua Sea Surface Temperature monthly composite image for October
- 821 2010.
- 822
- 823 Fig. 2. Initial profiles of temperature (blue line), nitrate (black line, open squares) and
- 824 chlorophyll a (green line, filled squares) at stations sampled for the CO₂ enrichment
- 825 experiments (A.) I, (B.) II, (C.) III. Dotted line is the euphotic depth.
- 826
- **Fig. 3.** Changes in pCO₂ during experiments (A) I, (B) II, (C) III, and in pH during
- 828 experiments (D) I, (E) II, (F) III in control (open circles) and 760 ppm CO₂ (closed circles).
- 829 Broken line in (C) and (F) represents phases of (space) and dark (line) during the experiments.

- **Fig. 4.** Changes in HCO₃⁻ during experiments (A) I, (B) II, (C) III, and in HCO₃⁻: CO₂ during
- 832 experiments (D) I, (E) II, (F) III in control (open circles) and 760 ppm CO₂ (closed circles).
- 833 Broken line in (C) and (F) represents phases of light (space) and dark (line) during the
- 834 experiments.

836	Fig. 5. Changes in Chlorophyll <i>a</i> during experiments (A) I, (B) II, (C) III. Broken line in (C)
837	represents phases of light (space) and dark (line) during the experiments.
838	
839	Fig. 6. Changes in the biomass of nanoeukaryotes in experiments (A.) I, (B.) II, (C.) III, and
840	changes in biomass of picoeukaryotes in experiments (D.) I, (E.) II, (F.) III in control (open
841	circles) and 760 ppm CO ₂ (closed circles). Broken line in (C) and (F) represents phases of
842	light (space) and dark (line) during the experiments.
843	
844	Fig. 7. Changes in the biomass of Synechococcus spp. in experiments (A.) I, (B.) II, (C.) III,
845	and changes in biomass of Prochlorococcus spp. in experiments (D.) I, (E.) II, (F.) III in
846	control (open circles) and 760 ppm CO ₂ (closed circles). Broken line in (C) and (F) represents
847	phases of light (space) and dark (line) during the experiments.
848	
849	Fig. 8. Changes in diatom biomass during experiments (A) II and (B) III, and dinoflagellate
850	biomass during experiments (C) II and (D) III at time(T) 0 and 48 h in control (AIR) and 760
851	ppm CO ₂ enrichment. In (A) and (B) brown is <i>Navicula</i> spp., orange is pennate datioms, light
852	green is Nitszchia spp., dark green is Thalassiosira spp., light blue is Rhizosolenia spp. and
853	dark blue is total diatom biomass. In (C) and (D) black is Amphidinium spp., dark blue is
854	Gonyaulax spp., light blue is Gymnodinium spp., green is Gyrodinium spp., yellow is total
855	dinoflagellate biomass (N=1; microscope transects = 4).
856	
857	Fig. 9. Changes in P_m^B [mg C (mg Chla) ⁻¹ h ⁻¹] during experiments (A.) I, (B.) II and (C.) III.
858	Changes in α^{B} [mg C (mg Chla) ⁻¹ h ⁻¹ (µmol m ⁻² s ⁻¹) ⁻¹] during experiments (D.) I, (E.) II and

(F.) III. Broken line in (C) and (F) represents phases of light (space) and dark (line) during
the experiments.

- 862 Fig. 10. Changes in low temperature emission during experiment (A) I CO₂ treatment, (B) I
- 863 control, (C) II CO₂ treatment, (D) II control, (E) III CO₂ treatment, and (F) III control (N=1).
- 864 Spectra are: black phycoerythrin; red phycocyanin; light green unconnected antenna;
- 865 dark-blue photosystem (PS) II or phycobilisomes anchoring pigments; yellow PSII core
- 866 antenna; cyanin PSI; light-blue vibrational peak; grey cumulative sum of all spectra.
- 867
- **Fig. 11.** Changes in Primary production $[mg C m^{-3} d^{-1}]$ at 48h in control (grey bars) and 760
- 869 ppm CO_2 (black bars) for experiments I-II.