

Dear Editor,

We are grateful that you have appreciated the effort to improve this work.

Your technical corrections and comments are in plain font and our responses are in italic and blue font. The revised version of the manuscript with marked changes is also provided.

L19: Please use the math symbol  $\times$ , not the alphabet x.

*This has been corrected (L19)*

L38: Add a period immediately after spp (i.e., spp.).

*This has been corrected (L38)*

L76: Insert a space immediately before Fuhrman.

*This has been corrected (L76)*

L169–170: Please cite a reference for the absorption coefficient of pure Chl-a standard.

*This has been corrected (L170)*

L178–179: Gasol and Del Giorgio (2000)

*This has been corrected (L179)*

L189: The symbol of minus is different from those at L198 and L203. So please amend it.

*This has been corrected (L189)*

L245–L247: Please cite references for the forward and reverse primer pairs for prokaryotes and eukaryotes.

*Both references has been included (L245;L247)*

L302: Not chl-a, but Chl-a.

*This has been corrected (L302)*

L311: Please use  $R^2$  (cf. see L1037).

*This has been corrected (L311)*

L468: Use a semi-colon between 2012 and Barber-Lluch.

*This has been corrected (L468)*

L536: Use an en dash for 0.1–10.

*This has been corrected (L536)*

L982: inorganic

*This has been corrected (L996)*

L1005: The “a” in Chlorophyll-a should be italic.

*This has been corrected (L1020)*

Figures 2 and 5: The “a” in Chl-a should be italic following the text.

*This has been changed*

*Additionally, references has been revised and Table 1 has been included in the text (L985).*

1 **Spatial and temporal variability in the response of**  
2 **phytoplankton and bacterioplankton to B-vitamin**  
3 **amendments in an upwelling system**

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9

10 **Abstract.** We experimentally evaluated the temporal (inter-day and inter-season) and  
11 spatial variability in microbial plankton responses to vitamin B12 and/or B1 supply  
12 (solely or in combination with inorganic nutrients) in coastal and oceanic waters of the  
13 northeast Atlantic Ocean. Phytoplankton and, to a lesser extent, prokaryotes were strongly  
14 limited by inorganic nutrients. Inter-day variability in microbial plankton responses to B-  
15 vitamins was limited compared to inter-season variability, suggesting that B-vitamins  
16 availability might be partially controlled by factors operating at seasonal scale.  
17 Chlorophyll-*a* (Chl-*a*) concentration and prokaryote biomass (PB) significantly increased  
18 after B-vitamin amendments in 13 % and 21 %, respectively, of the 216 cases (36  
19 experiments ~~xx~~ 6 treatments). Most of these positive responses were produced by  
20 treatments containing either B12 solely or B12 combined with B1 in oceanic waters,  
21 which was consistent with the significantly lower average vitamin B12 ambient  
22 concentrations compared to that in the coastal station. Negative responses, implying a  
23 decrease in Chl-*a* or PB, represented 21 % for phytoplankton and 26 % for prokaryotes.  
24 Growth stimulation by B1 addition was more frequent on prokaryotes than in  
25 phytoplankton, suggesting that B1 auxotrophy in the sampling area could be more  
26 widespread in prokaryotes than in phytoplankton. Negative responses to B-vitamins were  
27 generalized in coastal surface waters in summer, and were associated to a high  
28 contribution of Flavobacteriales to the prokaryote community. This observation suggests  
29 that the external supply of B12 and/or B1 may promote negative interactions between  
30 microbial components when B-vitamin auxotrophs are abundant. The microbial response  
31 patterns to B12 and/or B1 amendments were significantly correlated with changes in the  
32 prokaryotic community composition, highlighting the pivotal role of prokaryotes in B-  
33 vitamins cycling in marine ecosystems.

34

## 35 **1 Introduction**

36 Phytoplankton accounts for almost half of the global net primary production (Field et al.,  
37 1998) and may eventually cause toxic episodes, such as those caused by harmful algae  
38 blooms of *Alexandrium* spp. or *Gymnodinium* spp., entailing human health problems and  
39 large economic losses (Hallegraeff, 1993; van Dolah et al., 2001). Recent emerging  
40 evidence suggests the role of biologically active organic compounds, such as B-vitamins,  
41 on the control of marine productivity in both coastal and oceanic waters (Panzeca et al.,  
42 2006; Bertrand et al., 2007; Gobler et al., 2007; Koch et al., 2011; Browning et al., 2017,  
43 2018). B-vitamins act as cofactors for enzymatic reactions and are involved in many  
44 important metabolic pathways (Madigan et al., 2005; Koch et al., 2011; Monteverde et  
45 al., 2017). Vitamin B12 (B12 herein), which is exclusively synthesized by some bacteria  
46 and archaea (Roth et al., 1996; Martens et al., 2002; Warren et al., 2002), acts as a cofactor  
47 of three enzymes in eukaryotes (methionine synthase, methylmalonyl-coA mutase and  
48 ribonucleotide reductase type II) (Helliwell et al., 2011; Bertrand and Allen, 2012). In  
49 comparison, over 20 different B12-dependent enzymes are found in bacteria (Roth et al.,  
50 1996), making B12 critically important also for these organisms. Vitamin B1 (B1 herein)  
51 plays a pivotal role in intermediary carbon metabolism and is a cofactor for a number of  
52 enzymes involved in primary carbohydrate and branched-chain amino acid metabolism  
53 (Croft et al., 2006).

54 Most eukaryote phytoplankton species are auxotrophs for one or more B-vitamins,  
55 consequently requiring an exogenous supply of these molecules (Bertrand and Allen,  
56 2012; Carlucci and Bowes, 1970; Haines and Guillard, 1974; Helliwell et al., 2011).  
57 Moreover, genomic data also indicate widespread B-vitamins auxotrophy among many  
58 bacterial taxonomic groups (Sañudo-Wilhelmy et al., 2014; Paerl et al., 2018), which  
59 implies that phytoplankton and bacterioplankton may eventually compete for the

60 acquisition of these compounds (Koch et al., 2012). Auxotrophic microorganisms may  
61 acquire the required vitamins from the environment or through biotic interactions with  
62 prototrophic (biosynthetically competent) microorganisms (Droop, 2007; Grant et al.,  
63 2014; Kazamia et al., 2012). A well-known example is the mutualistic interaction  
64 between B12 or B12 and B1 dependent phytoplankton and bacterioplankton (Croft et al.,  
65 2005; Amin et al., 2012; Cooper and Smith, 2015; Cruz-López and Maske, 2016).

66 Even though B-vitamins appear to be important and potentially limiting factors for  
67 microbial plankton, our understanding of B-vitamins cycling in the ocean is largely  
68 limited by the complex and still evolving analytical methodology for its quantification in  
69 natural waters (Okbami and Sañudo-Wilhelmy, 2004, 2005; Suffridge et al., 2017).  
70 Sañudo-Wilhelmy et al. (2012) found extensive areas of coastal waters with close to  
71 undetectable B12 concentrations, suggesting that microbes might be well adapted to  
72 thrive under limiting conditions for this growth factor.

73 The factors limiting phytoplankton and bacterial growth in marine ecosystems are known  
74 to vary over different spatial and temporal scales (Cullen et al., 1992; Arrigo, 2005;  
75 Martínez-García et al., 2010b; Moore et al., 2013), in accordance with the dynamic nature  
76 of microbial communities (Pinhassi et al., 2003; Fuhrman et al., 2008; Hernando-Morales  
77 et al., 2018). Compared to mineral nutrient and trace elements, much less is known about  
78 B vitamin limitation and its spatial and temporal variability in marine ecosystems.

79 Some studies have shown enhanced phytoplankton biomass associated to B12  
80 amendments in both temperate coastal and polar waters (Bertrand et al., 2007; Gobler et  
81 al., 2007; Koch et al., 2011, 2012). The simultaneous effect of vitamin B12 supply on  
82 both phytoplankton and bacteria has been barely explored (Koch et al., 2011, Barber-  
83 Lluch et al., 2019). To our knowledge, the effect of B1 amendments on marine natural  
84 microbial plankton community succession has been only assessed by Gobler et al. (2007),

85 who suggested that high concentration of B-vitamins, associated with high bacterial  
86 abundance, caused an increase in auxotrophs, mostly dinoflagellates.

87 The Ría de Vigo (NW Spain) is a coastal embayment affected by intermittent upwelling  
88 of subsurface cold and inorganic nutrient-rich water from March to September and the  
89 downwelling of open ocean surface water from October to March (Fraga, 1981; Barton  
90 et al., 2015). In addition to this seasonality, fluctuations of wind patterns in the area  
91 generate upwelling and downwelling events occurring within each season (Alvarez-  
92 Salgado et al., 1993; Figueiras et al., 2002). A recent study by Barber-Lluch et al. (2019)  
93 at a shelf station off the Ría de Vigo (NW Spain) showed monthly variation in the  
94 response of phytoplankton and bacteria to nutrient and/or B12 additions in surface waters,  
95 likely related to variation in the ambient concentration of B12 and the taxonomic  
96 community composition. Unfortunately, these authors did not specifically assess the role  
97 of these factors on the microbial response to the amendments.

98 Within this context, the aim of our study was to explore spatial (horizontal and vertical)  
99 and temporal (inter-day and inter-season) variability patterns in B12 and B1 vitamin  
100 limitation in relation to the prevailing initial abiotic (e.g., nutrient and B12  
101 concentrations) and biotic (eukaryote and prokaryote community composition)  
102 conditions in this productive ecosystem. We conducted a total of thirty-six microcosm  
103 bioassays in February, April, and August 2016 to evaluate the response of heterotrophic  
104 bacteria and phytoplankton biomasses to the addition of B12 and/or B1.

105 Considering that a large fraction of eukaryotic phytoplankton and bacterial taxa require  
106 exogenous B-vitamins and considering the different requirements and capabilities to  
107 synthesize B-vitamins by different microbial taxa, we hypothesize that microbial  
108 community composition play a relevant role in explaining B-vitamins limitation patterns  
109 in microbial plankton.

110

## 111 **2 Methods**

### 112 **2.1 Sampling strategy**

113 Thirty-six enrichment experiments were performed in the upwelling system near Ría de  
114 Vigo on board “B/O Ramón Margalef” in three different oceanographic cruises  
115 (ENVISION I, II & III) conducted in 2016. Two different locations of the East Atlantic  
116 Ocean, one coastal station (C) (42° N, 8.88° W) and one oceanic station (Oc) (42° N, 9.06°  
117 W) (Fig. 1a), were sampled during three different seasons aimed to cover a wide range of  
118 initial hydrographic and ecological conditions. The 10-day cruises were conducted in  
119 February (ENVISION I), coinciding with the spring bloom, and April (ENVISION II)  
120 and August (ENVISION III) during the early and late summer upwelling, respectively.  
121 During each cruise, 12 enrichment experiments were carried out on board, 3 experiments  
122 in each station (C-a, C-b & C-c and Oc-a, Oc-b & Oc-c, respectively) with water from  
123 two different depths. Each experiment began on the first (day 0), third (day 2) and sixth  
124 (day 5) of each cruise for the coast and on the second (day 1), fourth (day 3) and seventh  
125 (day 6) of each cruise for the ocean (Fig. 1b, c). Water was collected using 20 l Niskin  
126 metal-free bottles. Surface (5 m) and sub-surface chlorophyll maximum (SCM) (between  
127 10 m and 50 m according to the CTD data) samples were taken (Fig. 2a-f). We failed to  
128 sample the SCM on two occasions (C-a in February and C-a in April), due to large vertical  
129 displacements between the downward and the upward casts. Vertical profiles of  
130 temperature, salinity and chlorophyll fluorescence were obtained using a regular stainless  
131 CTD-rosette down to 60 m in the coastal station and to 200 m in oceanic station. Samples  
132 for chlorophyll-*a* (Chl-*a*), prokaryotic biomass (PB), dissolved nutrient concentration,  
133 including vitamin B12, and microbial plankton community were collected at the

134 beginning (time zero, hereafter referred to as  $t_0$ ) of each enrichment experiment. Daily  
135 upwelling index (UI) values were computed by the Instituto Español de Oceanografía  
136 ([www.indicedeafloramiento.ieo.es/](http://www.indicedeafloramiento.ieo.es/)) in a  $2^\circ \times 2^\circ$  geostrophic cell centered at  $42^\circ\text{N}$ ,  $10^\circ\text{W}$ , using data from atmospheric pressure at sea level, derived from the WXMAP model  
137  $^\circ\text{W}$ , using data from atmospheric pressure at sea level, derived from the WXMAP model  
138 (Gonzalez-Nuevo et al., 2014). Precipitation data was obtained from the Regional  
139 Weather Forecast Agency-Meteogalicia (<http://www.meteogalicia.gal>) in the  
140 meteorological station Illas Cies (ID 10125).

## 141 **2.2. Experimental design**

142 Seawater samples were gently pre-filtered through a  $200\ \mu\text{m}$  mesh to exclude large  
143 zooplankton in order to ensure good replicability and collected into a 20 l acid-cleaned  
144 polyethylene carboy. It is important to note that incidental trace-metal contamination  
145 could have occurred during water collection. Following sample collection, 300 ml PAR  
146 and UVR transparent, sterile, and non-toxic (whirl-pak) bags were filled and nutrients  
147 were added establishing eight different enrichment treatments as follows: (1) control  
148 treatment (C); (2) inorganic nutrient treatment (I); (3) vitamin B12 (Sigma, V2876)  
149 treatment; (4) vitamin B1 (Sigma, T4625) treatment; (5) Inorganic nutrients and vitamin  
150 B12 (I+B12) treatment; (6) Inorganic nutrients and vitamin B1 (I+B1) treatment; (7)  
151 vitamins B12 and B1 (B12+B1) treatment and (8) Inorganic nutrients with vitamins B12  
152 and B1 (I+B12+B1) treatment (see Table 1 for details). Inorganic nutrients were added to  
153 avoid that inorganic nutrient limitation masked the responses to B vitamins. The nutrient  
154 concentrations of the additions were the same as previously used in similar enrichment  
155 experiments in the sampling area (Martinez-García et al., 2010a). The amount of B12 and  
156 B1 vitamin experimentally added approximated maximum concentrations previously  
157 observed in coastal areas (Okbami and Sañudo-Wilhelmy 2004, 2005, Sañudo-  
158 Wilhelmy et al., 2006). Each treatment had 3 replicates resulting in 24 whirl-pak bags

159 per experiment. To assess short-term effects of nutrient inputs, experimental bags were  
160 incubated on-deck during 72 h. In-situ temperature was reproduced by submerging the  
161 bags in tanks filled with constantly circulating surface seawater. To simulate light  
162 intensity at the SCM the incident light was attenuated by covering the tanks with mesh  
163 screens.

### 164 **2.3 Chlorophyll-*a***

165 Chlorophyll-*a* (Chl-*a*) concentration was measured at t0 and after 72 h incubation as a  
166 phytoplankton biomass proxy. 300 ml of water samples were filtered through 0.2 µm  
167 polycarbonate filters and frozen at -20°C until further analysis. Chl-*a* was extracted with  
168 90 % acetone and kept in darkness at 4°C overnight. Fluorescence was determined with a  
169 TD-700 Turner Designs fluorometer calibrated with pure Chl-*a* standard solution  
170 (absorption coefficient at ~~665-663~~ nm = ~~12.687.7~~; [Lorenzen and Newton Downs, 1986](#)).

### 171 **2.4 Flow cytometry**

172 Samples for prokaryote abundance quantification (2 ml) were preserved with 1 %  
173 paraformaldehyde + 0.05 % glutaraldehyde (final concentrations). Samples were  
174 incubated 20 min for the fixative to act on cells, immersed in liquid nitrogen for 15 min,  
175 and frozen at -80°C. Abundance of prokaryotes was determined using a FACSCalibur  
176 flow cytometer equipped with a laser emitting at 488nm. Samples were stained with  
177 SYBR Green DNA fluorochrome, and bacterial abundance was detected by their  
178 signature of side scatter (SSC) and green fluorescence as described by Gasol and Del  
179 Giorgio, (2000). The empirical calibration between light side scatter (SSC) and cell  
180 diameter described by Calvo-Díaz and Moran (2006) were used to estimate cell  
181 biovolume (BV). BV was converted into biomass by using the allometric factor of

182 Norland (1993:  $\text{fg C cell}^{-1} = 120 \times \text{BV}^{0.72}$ ) for the coastal experiments and using the open  
183 ocean conversion factor for the oceanic experiments ( $\text{fg C cell}^{-1} = 350 \times \text{BV}$ ).

## 184 **2.5 Nutrients**

185 Aliquots for inorganic nutrient determinations (ammonium, nitrite, nitrate, phosphate,  
186 and silicate) were collected before all other variables and directly from the Niskin bottle  
187 in order to avoid contamination. Polyethylene bottles (50 ml) precleaned with 5 % HCl  
188 were filled with the sample using contamination-free plastic gloves and immediately  
189 frozen at  $-20^{\circ}\text{C}$  until analysis using standard colorimetric methods with a Bran-Luebbe  
190 segmented flow analyzer (Hansen and Grasshoff 1983). The detection limit was  $0.1 \mu\text{mol}$   
191  $\text{l}^{-1}$  for nitrate,  $0.02 \mu\text{mol l}^{-1}$  for nitrite and phosphate and  $0.05 \mu\text{mol l}^{-1}$  for ammonium  
192 and silicate. Dissolved inorganic nitrogen (DIN) concentration was calculated as the sum  
193 of the ammonium, nitrite and nitrate concentrations.

## 194 **2.6 Vitamin B12**

195 Seawater samples for dissolved vitamin analysis were taken at surface and SCM depth on  
196 day 1, day 3 and day 5 in the coastal, and on day 1, day 3 and day 6 oceanic station of  
197 each cruise (Table S1 in the Supplement). Samples were filtered through  $0.2 \mu\text{m}$  sterivex  
198 filters and frozen at  $-20^{\circ}\text{C}$  until further analysis. Samples (1 l) were preconcentrated using  
199 a solid-phase extraction with a C18 resin (Bondesil C18, Agilent) at pH 6.5 and rate of  
200  $1 \text{ml/min}$ . Elution was performed with 12 ml of methanol (MeOH) LCMS grade that was  
201 removed via evaporation with nitrogen in a Turbovap. Gas pressure was initially set at 5  
202 PSI and was slowly increased to 15 PSI until 300-500  $\mu\text{l}$  of sample remained. The  
203 concentrated samples were frozen at  $-20^{\circ}\text{C}$  until further analysis using liquid  
204 chromatography coupled to mass spectrometry system.

205 The concentrate was filtered again through a cellular acetate membrane 0.2  $\mu\text{m}$   
206 (Phenomenex) prior to the analysis. Ultra Performance Liquid Chromatography tandem  
207 Mass Spectrometry 3Q (UPLC-MS/MS) methodology was adapted from Sañudo-  
208 Wilhelmy et al. (2012), Heal et al. (2014) and Suffridge et al. (2017). Detection and  
209 quantification of dissolved vitamin B12 (cyanocobalamin and hydroxocobalamin) was  
210 conducted using an Agilent 1290 Infinity LC system (Agilent Technologies, Waghaeusel-  
211 Wiesental, Germany), coupled to an Agilent G6460A triple quadrupole mass  
212 spectrometer equipped with an Agilent Jet Stream ESI source. The LC system used a C18  
213 reversed-phase column (Agilent Zorbax SB-C18 Rapid Resolution HT (2.1 inner  
214 diameter  $\times$  50 mm length, 1.8  $\mu\text{m}$  particle size) with a 100  $\mu\text{l}$  sample loop. Agilent  
215 Technologies software was used for data acquisition and analysis. Chromatographic  
216 separation was performed using MeOH and water LCMS grade, both buffered to pH 5  
217 with 0.5 % acetic acid, as mobile phases in a 15 minutes' gradient. Gradient starting at 7  
218 % MeOH for 2 min, changing to 100 % MeOH by minute 11, continuing at 100 % MeOH  
219 until 13.5 min and returning to initial conditions to complete 15 min. Limits of detection  
220 (LODs) and limits of quantification (LOQs) were determined using sequential dilutions  
221 of the lowest point of the calibration curves. LODs were defined as the lowest detectable  
222 concentration of the analyte with a signal-to-noise (S/N) ratio for the qualitative transition  
223 of at least 3. In the same way, LOQs were defined as the lowest quantifiable  
224 concentration with a S/N ratio of 10 for the quantitative transition. S/N ratios were  
225 calculated using the Mass Hunter Workstation software B.04.01. The LODs obtained  
226 were 0.04  $\text{pmol l}^{-1}$  for hydroxocobalamin (OHB12) and 0.01  $\text{pmol l}^{-1}$  for cyanocobalamin  
227 (CNB12), while the LOQs values were 0.05 and 0.025  $\text{pmol l}^{-1}$  for OHB12 and CNB12,  
228 respectively. The average B12 recovery percentage after pre-concentration and extraction  
229 of B-vitamin spiked samples was 93%. B-vitamin free seawater was spiked with CNB12

230 and OHB12 standards for recovery percentage analysis. We failed to detect B1 vitamin  
231 in the pre-concentrated samples, likely due to a low ambient concentration and low pre-  
232 concentration volume.

## 233 **2.7 Microbial plankton community**

234 DNA samples were taken during the experimental period at surface and SCM depth in  
235 the coastal and oceanic station. In particular, sampling of the microbial plankton  
236 community was carried out on day 0, day 1, day 3 and day 5 of each cruise. Community  
237 composition was assessed by sequencing the V4 and V5 regions from 16S rRNA gene  
238 (16S rDNA) for prokaryotes and the V4 region from 18S rRNA gene (18S rDNA) for  
239 eukaryotes. Two liters of water samples were sequentially filtered through 3  $\mu\text{m}$  pore size  
240 polycarbonate filters and 0.2  $\mu\text{m}$  pore size sterivex filter and immediately frozen in liquid  
241 nitrogen and conserved at -80  $^{\circ}\text{C}$ . DNA retained in the 3  $\mu\text{m}$  and 0.2  $\mu\text{m}$  filters was  
242 extracted by using the PowerSoil DNA isolation kit (MoBio Laboratories Inc., CA, USA)  
243 and the PowerWater DNA isolation kit (MoBio Laboratories Inc., CA, USA),  
244 respectively, according to the manufacturer's instructions. Prokaryotic DNA from 0.2  $\mu\text{m}$   
245 filters was amplified using the universal primers "515F and 926R" (Parada et al., 2016)  
246 and eukaryotic DNA from both, 3  $\mu\text{m}$  and 0.2  $\mu\text{m}$  filters, using the primers  
247 "TAREuk454FWD1" and "TAREukREV3" (Logares et al., 2014). Amplified regions  
248 were sequenced in an Illumina MiSeq platform and the sequences obtained were analyzed  
249 with software package DADA2 (Callahan et al., 2016). SILVA reference database (Quast  
250 et al., 2012) was used to taxonomic assignment of 16S amplicon sequence variants  
251 (ASVs) and PR2 (Guillou et al., 2013) and the marine protist database from the BioMarks  
252 project (Massana et al., 2015) were used to taxonomic assignment of 18S ASVs. The data  
253 for this study have been deposited in the European Nucleotide Archive (ENA) at EMBL-  
254 EBI (<https://www.ebi.ac.uk/ena>) under accession numbers PRJEB36188 (16S rDNA

255 sequences) and PRJEB36099 (18S rDNA sequences). ASV table is an analogue of the  
256 traditional OTU table which records the number of times each exact amplicon sequence  
257 variant was observed in each sample (Callahan et al., 2016).

258 The raw ASV tables of prokaryotes and eukaryotes were subsampled to the number of  
259 reads present in the sample with the lowest number of reads, which was 2080 and 1286,  
260 for 16S rDNA and 18S rDNA, respectively. The abundance of ASVs was averaged for  
261 coastal and oceanic samples, differentiating surface and SCM. A total of 1550 unique  
262 ASVs of prokaryotes were identified. As many ASVs of eukaryotes were present in both  
263 size fractions (e.g. those having a cell size range including 3  $\mu\text{m}$ ), we combined datasets  
264 derived from the 0.2 and the 3  $\mu\text{m}$  filters for eukaryotic community analyses. As explained  
265 in Hernández-Ruiz et al. (2018), we normalized the reads from each filter size by the filter  
266 DNA yield, as recommended in Dupont et al. (2015), obtaining 2293 unique ASVs. The  
267 sequence abundances of the subsampled ASV tables were transformed using the centered  
268 log ratio (clr) (Fernandes et al., 2014; Gloor et al., 2017). Before clr transformation, zeros  
269 were replaced by the minimum value that is larger than 0 divided by 2 (Aitchison, 1982;  
270 Martín-Fernández et al., 2003).

## 271 **2.8 Statistical analysis**

272 To compare the effect of different nutrient additions on the response variables,  
273 chlorophyll-*a* concentration and prokaryote biomass, we calculated response ratios (RR)  
274 by dividing each observation (mean of triplicates) of each treatment by the respective  
275 control treatment mean. A value equal to 1 implies no response, a value  $< 1$  implies a  
276 negative response and a value  $> 1$  implies growth stimulation after nutrient addition.  
277 Secondary limitation by B vitamins was calculated by dividing the mean value in the  
278 inorganic nutrients and B vitamin combined treatment by the mean value in the inorganic

279 nutrient addition treatment. In the same way, a value  $< 1$  implies a negative effect of B  
280 vitamins and a value  $> 1$  implies stimulation positive effect of B vitamin treatment  
281 through secondary limitation.

282 Normal distribution was tested by a Kolmogorov-Smirnov test and non-normal variables  
283 such as temperature, salinity, DIN,  $\text{SiO}_4^{2-}$ , and Chl-*a* and PB response ratios, were log  
284 transformed to attain normality. All statistical analysis were considered significant at the  
285 0.05 significance level and p-value was standardized as proposed by Good (1982) in order  
286 to overcome the low number of replicates. Differences between station and depth (spatial  
287 variability) and among sampling months (temporal variability) in the responses to B  
288 vitamins were evaluated with factorial analysis of variance (ANOVA). Bonferroni post  
289 hoc tests analyses were conducted to test which treatments were significantly different  
290 from the control treatment in each experiment. Non-metric multidimensional scaling  
291 (nMDS) was used to analyze the similarities between the samples based on microbial  
292 assemblage structure using the PRIMER6 software (Clarke and Warwick, 2001; Clarke  
293 and Gorley, 2006). The similarities were evidenced in a multidimensional space by  
294 plotting more similar samples closer together. Analysis of similarity (ANOSIM) was used  
295 to verify that microbial community composition from the same season and station were  
296 more similar to each other than to communities from a different season and station. Z-test  
297 was used to test if averaged B vitamins response ratios were significantly different from  
298 1. The RELATE analysis implemented in PRIMER6 was used to relate the B-vitamin  
299 response patterns (Bray-Curtis resemblance matrix built from phytoplankton and bacteria  
300 response ratios) with: (1) environmental factors (Euclidean resemblance matrix built from  
301 normalized values of ammonium, nitrite, nitrate, phosphate, silicate, B12, temperature,  
302 salinity, ~~chl~~Chl-*a* and prokaryote biomass), (2) prokaryote community composition  
303 (Euclidean resemblance matrix built form clr-transformed sequence abundance of major

304 taxonomic groups), or (3) eukaryote community composition (Euclidean resemblance  
305 matrix built from clr-transformed sequence abundance of major taxonomic groups).  
306 RELATE calculates the Spearman rank correlations (Rho) between two resemblance  
307 matrices, and the significance is tested by a permutation test (999 permutations). In order  
308 to highlight which specific taxonomic groups are associated to changes of microbial  
309 plankton (prokaryote plankton and phytoplankton) responses to vitamin B1 and B12, we  
310 conducted a distance based redundancy analysis (dbRDA) combined with a distance  
311 linear-based model (DistLM) using a step-wise procedure and adjusted  $F^2-R^2$  as selection  
312 criteria using the PRIMER6 software.

313

### 314 **3 Results**

#### 315 **3.1 Initial conditions**

316 Different hydrographic conditions were found during each cruise (Fig. 1 and Fig. 2). In  
317 February, heavy rainfall (Fig. 1c) combined with relaxed winds caused a halocline at 10  
318 m depth (Fig. 2m). High levels of Chl-*a* (as derived from the calibrated CTD fluorescence  
319 sensor) were observed at the coastal station, being maximum ( $4.97 \mu\text{g l}^{-1}$ ) by the end of  
320 the cruise (Fig. 2a). At the oceanic station, Chl-*a* levels remained low (less than  $3 \mu\text{g l}^{-1}$ )  
321 throughout the cruise, being slightly higher in the subsurface layer (Fig. 2d).

322 Strong precipitation during the April cruise (Fig. 1c) caused a persistent surface halocline  
323 at the coastal station (Fig. 2n). Maximum Chl-*a* concentrations ranged from 0.99 to  $2.73$   
324  $\mu\text{g l}^{-1}$ , declining from day 5 onwards (Fig. 2b), coinciding with an increase in water  
325 temperature associated to a downwelling situation. At the oceanic station, a persistent  
326 subsurface Chl-*a* maximum (up to  $1.61 \mu\text{g l}^{-1}$ ) was observed throughout the cruise (Fig.  
327 2e).

328 In August, strong thermal stratification was observed at both stations (Fig. 2i and Fig. 2l).  
329 At the beginning of the cruise, high Chl-*a* concentration (close to 20  $\mu\text{g l}^{-1}$ ) was observed  
330 in subsurface water (Fig. 2c). Chl-*a* was relatively low at the oceanic station, and  
331 increased by the end of the sampling period (Fig. 2f) as a consequence of an upwelling  
332 event (Fig. 1b), that brought cold and nutrient rich water to the surface, at day 5.

333 Abiotic and biotic conditions at the beginning of each experiment are shown in Fig. 3 and  
334 in the supplementary Table S2. Overall, the concentration of dissolved inorganic nitrogen  
335 (DIN) was higher at the coastal than at the oceanic station, where very low levels were  
336 measured in August (Fig. 3i). At the coastal station, higher DIN concentrations were  
337 observed in surface compared to subsurface waters. The DIN:DIP (dissolved inorganic  
338 phosphorous) ratio was always lower in open ocean than in the coastal station and mostly  
339 below the Redfield ratio (16:1). Phosphorous limitation (DIN:DIP > 16) was frequent in  
340 coastal surface waters in February and April (Fig. 3j and Fig. 3k).

341 On average, chl-*a* concentration varied greatly between stations and months but was  
342 always higher at the coastal than at the oceanic station (Fig. 3a-c). Prokaryote biomass  
343 (PB) increased from winter (February) to summer (August) at the two stations (Fig. 3d-  
344 f). In February, Chl-*a* concentrations increased by the end of the cruise at both coastal  
345 and oceanic stations (Fig. 3a), while PB remained very low throughout this sampling  
346 period (Fig. 3d). In April, both PB and Chl-*a* were similar in the ocean and the coast, and  
347 showed reduced temporal variability (Fig. 3b and Fig. 3e), irrespective of the observed  
348 nutrient variability (Fig. 3h). In August, Chl-*a* concentration was much higher at the  
349 coastal than at the oceanic station, and showed reduced temporal variability (except at the  
350 SCM in the coast) (Fig. 3c). At the beginning of the sampling period, PB was higher in  
351 the ocean than in the coast, and tended to decline by the end of the cruise (Fig. 3f).

352 A MDS analysis revealed that microbial community composition showed a relatively  
353 reduced variability within period, with samples clustering according to the sampling  
354 period (ANOSIM,  $p = 0.001$ ) and station (ANOSIM,  $p = 0.001$ ) (Fig. S1 in the  
355 Supplement). Consequently, we averaged the microbial community composition for each  
356 period and sampling site. The sampling period-averaged composition of the eukaryote  
357 community showed a clear variability among cruises, while differences between sampling  
358 locations and depths were less pronounced (Fig. 4a). At the coastal location,  
359 Mamiellophyceae (*Ostreococcus* and *Micromonas*) were relatively abundant in February  
360 and April, but their relative abundance sharply decreased in August. By contrast, the  
361 relative abundance of Dinophyceae was highest in August at both sampling locations.  
362 The contribution of diatoms (Bacillariophyta) was very low in summer at the oceanic  
363 station and marine alveolates (MALV) groups (MALV-I and MALV-II) were most  
364 representative in February at both locations. Flavobacteriales and Rhodobacterales were  
365 the dominant prokaryotes (Fig. 4b) in coastal waters, particularly in August, when both  
366 represented more than 80 % of sequences, while the Cyanobacteria *Synechococcus* were  
367 mostly present in February and April. In oceanic waters, Flavobacteriales and  
368 *Synechococcus* were the dominant prokaryotes. SAR11 clade and Archaea  
369 (Euryarchaeota and Thaumarchaeota) were most abundant in February at both sampling  
370 locations.

371 B12 concentration was low, ranging from 0.06 to 0.66 pmol l<sup>-1</sup> (Table S1 in the  
372 Supplement). Average B12 concentration was significantly higher in the coast (0.30±0.13  
373 pmol l<sup>-1</sup>) than in the ocean (0.15±0.12 pmol l<sup>-1</sup>) (t-test,  $t = 3.17$ ,  $df = 10$ ,  $p = 0.01$ ), and  
374 showed less variability at the coastal than at the oceanic station (Fig. 4c).

### 375 **3.2 Short-term phytoplankton and prokaryote responses to inorganic nutrients and** 376 **vitamin additions**

377 The temporal development of the phytoplankton (as estimated from changes in Chl-*a*  
378 concentration) and prokaryote biomass in the control treatments showed different  
379 patterns. Chl-*a* remained either stable or increased after 72 h of incubation in 87.5% of  
380 the experiments conducted in February and April. However, Chl-*a* mostly decreased in  
381 the coastal experiments conducted in August (Fig. 5a and Fig. 5c). A very similar pattern  
382 was observed for prokaryote biomass, although the decrease in biomass occurred both in  
383 the coastal and in the oceanic stations during summer (Fig. 6).

384 The response ratios (RRs) of Chl-*a* and prokaryote biomass were calculated as a measure  
385 of the magnitude of phytoplankton and prokaryote responses to nutrient and vitamin  
386 treatments (Fig S2, S3 and S4 in the supplement). The RRs differed between sampling  
387 stations (ANOVA,  $F(1,502) = 18.059$ ,  $p < 0.001$ ) and among sampling periods (ANOVA,  
388  $F(2,501) = 6.54$ ,  $p = 0.002$ ). The most prominent responses of phytoplankton, compared  
389 to the control treatment, occurred after inorganic nutrient amendments, especially in  
390 surface oceanic waters (Fig. 5c and Fig. S2b, f and j in the Supplement). The magnitude  
391 of the phytoplankton response to inorganic nutrients was significantly higher in oceanic  
392 than in coastal waters (ANOVA,  $F(1,34) = 5.22$ ,  $p = 0.028$ ). Prokaryotes responded less  
393 than phytoplankton to inorganic nutrients and, in addition, heterotrophic prokaryote  
394 responses to inorganic nutrients were similar between coastal and oceanic waters  
395 (ANOVA,  $F(1,34) = 1.68$ ,  $p = 0.203$ ). The addition of inorganic nutrients caused  
396 significant increases in Chl-*a* in 31 out of the 36 experiments (Fig. 5 and Fig S2 in the  
397 supplement), while prokaryotes increased their biomass in 19 out of 36 experiments (Fig.  
398 6 and Fig. S2 in the Supplement).

399 The addition of B12 stimulated phytoplankton in 5 out of 36 experiments (Fig. 5 and Fig.  
400 S3 in the Supplement) and prokaryotes in 6 experiments (Fig. 6 and Fig. S4 in the  
401 Supplement). Chl-*a* increased in 3, and prokaryote biomass in 7 out of 36 experiments

402 after adding B1 (Fig. 5 and Fig. 6). B vitamins also caused negative responses of  
403 phytoplankton (Fig. 5 and Fig. S3 in the Supplement) and prokaryote biomass (Fig. 6 and  
404 Fig. S4 in the Supplement). The addition of vitamins induced decreases of Chl-*a* in 6  
405 experiments (4 after adding B12 and 2 after adding B1) and prokaryote biomass in 14  
406 experiments (6 after adding B12 and 8 after adding B1). Secondary limitation by B1  
407 and/or B12 was occasionally observed when inorganic nutrients were limiting, leading to  
408 a higher biomass increase in the treatments including both inorganic nutrients and  
409 vitamins as compared to the inorganic nutrient addition alone (Fig. 5, Fig. 6 and Fig. S3  
410 and Fig. S4 in the Supplement). In the case of Chl-*a*, secondary limitation by B-vitamins  
411 was found in the C-b-surface, Oc-a-SCM and Oc-b-SCM experiments in February, in the  
412 C-b-surface and C-b-SCM experiments in April, and in the C-b-SCM, Oc-b-SCM and  
413 Oc-c-surface experiments in August (Fig. 5).

414 In order to quantify the relevance of inter-day variability, we calculated the mean  
415 coefficient of variation (CV) of the responses to B vitamins (i.e., excluding the responses  
416 to inorganic nutrients, and normalizing the responses of the nutrient and vitamin  
417 combined treatments to the corresponding response to inorganic nutrients alone) within  
418 sampling periods for each sampling point (2 stations and 2 depths). The CV ranged from  
419 9%, in subsurface oceanic waters in April, to 34% in surface coastal waters in April,  
420 averaging  $16 \pm 6$  (SD) % (data not shown). Considering that short-term (within sampling  
421 period) variability was overall very low, and for simplicity, we averaged the responses to  
422 B vitamins in the 3 experiments conducted at each of the 12 sampling points to further  
423 describe spatial and temporal patterns in the response to B vitamin amendments (Fig. 7).

424 When averaging the responses within each sampling point (Fig. 7), some general patterns  
425 emerge. Both phytoplankton and prokaryotes showed more negative than positive  
426 responses to B1 and/or B12 amendments. Most positive responses occurred at the oceanic

427 station (83.3%), while negative responses dominated in the coast (61.5%). Phytoplankton  
428 significant positive responses mostly occurred in February, showing an average increase  
429 of up to 1.2-fold in coastal subsurface waters after B12+B1 amendment (Fig. 7a). The  
430 largest significant increase in Chl-a (ca. 1.4-fold) occurred in April after the combined  
431 addition of B12 and B1 in coastal surface waters. Significant positive prokaryote  
432 responses mainly occurred in August, when the largest increase (ca. 1.3-fold) occurred in  
433 coastal subsurface waters after B1 amendment (Fig. 7b). Most positive responses were  
434 associated with treatments containing B12 either alone or combined with B1 (Fig. 7b).  
435 Phytoplankton primary B1 limitation was only found at the oceanic SCM in February  
436 (Fig. 7a), while prokaryote primary B1 limitation only occurred at the coastal SCM in  
437 August. In addition, prokaryote secondary B1 limitation occurred in oceanic surface  
438 waters in February and August.

### 439 **3.3 B-vitamin response patterns in relation to environmental factors and prokaryote** 440 **and eukaryote community composition**

441 In order to explore the controlling factors of the observed B-vitamin response patterns,  
442 the correlation between the B-vitamin response resemblance matrix and the  
443 corresponding resemblance matrices obtained from the initial environmental factors, the  
444 initial prokaryotic community composition, or the initial eukaryotic community  
445 composition were calculated. While eukaryotic community composition did not show a  
446 significant correlation with the B-vitamin responses (Spearman Rho = 0.05,  $p = 0.39$ ), the  
447 prokaryotic community composition was significantly correlated with the B-vitamin  
448 responses (Spearman Rho = 0.31,  $p = 0.041$ ). We then used distance-based linear  
449 modelling (DistLM) to identify the prokaryotic taxa which best explained the microbial  
450 plankton responses to B-vitamins (Fig. 8). The resulting model explained 78% of the  
451 variation and included seven prokaryotic groups: *Planktomarina* (24%), Actinobacteria

452 (14%), SAR11 (8.2%), Cellvibrionales (8.5%), Euryarchaeota (8.7%), Flavobacteriales  
453 (9%) and *Synechococcus* (6.1%). The sequential test identified *Planktomarina* and  
454 Actinobacteria as the taxa explaining the largest fraction of variation (ca. 24 % and 14%,  
455 respectively, data not shown). The total variation explained by the db-RDA1 (34.9%) and  
456 db-RDA2 (24.5%) was 59.4 %, both represented as x and y axis, respectively (Fig. 8).  
457 The db-RDA1 axis separated, to some extent, coastal samples, where negative responses  
458 to B vitamins dominated, from oceanic samples, where most positive responses were  
459 found (Fig. 7). The db-RDA plot showed that Cellvibrionales and *Planktomarina*  
460 positively correlated with axis 1, while SAR11 and *Synechococcus* showed negative  
461 correlation with axis 1. Flavobacteriales and Actinobacteria mostly correlated with the  
462 db-RDA2 axis.

463

#### 464 **4 Discussion**

465 Although the dependence of phytoplankton on B vitamin has been previously observed  
466 in cultures (e.g. Croft et al., 2006; Droop, 2007; Tang et al., 2010) and in natural microbial  
467 assemblages in coastal areas (e.g. Sañudo-Wilhelmy et al., 2006; Gobler et al., 2007;  
468 Koch et al., 2011, 2012; Barber-Lluch et al., 2019), this is, to the best of our knowledge,  
469 the most complete study about responses of phytoplankton and prokaryotes to vitamin  
470 B12 and/or B1 addition. The 36 experiments developed in this study contributed to  
471 increase our understanding of the role of vitamins B12 and B1 at different spatial and  
472 temporal scales.

473 Considering the high short-time variability of the hydrographic conditions in the area  
474 (Alvarez-Salgado et al., 1996), we expected a large inter-day variation in the responses  
475 to B vitamin amendments. By contrast, inter-day variability of microbial responses to B

476 vitamins and microbial plankton community composition was relatively small (Fig. 5,  
477 Fig. 6, Fig. S1 and Fig. S2 in the supplement). The reduced short-term variability in the  
478 responses to B vitamins additions suggested that B vitamin availability might be  
479 controlled by factors operating at larger temporal scales, such as the succession of  
480 microbial communities associated to seasonal environmental variation (Hernández-Ruiz  
481 et al., 2018; Hernando-Morales et al., 2018). Considering this, and for further discussion,  
482 we averaged the responses from the three experiments conducted during each sampling  
483 period, resulting in 12 experimental situations (2 stations  $\times$  2 depths  $\times$  3 periods). Overall,  
484 phytoplankton and/or prokaryote growth enhancement in at least one B vitamin treatment  
485 was frequent but relatively small in this productive ecosystem, showing 1.1 to 1.3-fold  
486 increases in 75% of the experimental situations for phytoplankton and in 50% for  
487 prokaryotes. On the other hand, negative responses to at least one B vitamin treatment  
488 occurred in 83% of the experimental situations for phytoplankton and in 67% for  
489 prokaryotes (Fig. 7). The low and constant B12 ambient concentration (Fig. 4c) and the  
490 reduced magnitude of microbial responses suggest a close balance between production  
491 and consumption of this growth factor. Different patterns of response to B-vitamin  
492 amendments were observed in phytoplankton and prokaryotes (Fig. 7), which appear to  
493 be mostly explained by the prokaryotic community composition (Fig. 8).

#### 494 **4.1 Positive responses to vitamin B1 and B12 amendments**

495 The experimental design allowed the detection of two categories of B vitamin dependency  
496 of the microbial plankton community. A primary limitation by B vitamins occurs when  
497 microorganisms respond to additions of B vitamins alone. A secondary limitation by B  
498 vitamins arises when the response to the combined addition of B vitamins and inorganic  
499 nutrients is significantly higher than that to inorganic nutrients alone. Such response  
500 occurs because of the ambient B-vitamin depletion associated to the plankton growth after

501 inorganic nutrient enrichment. Most positive (72% for phytoplankton and 60 % for  
502 prokaryotes) responses occurred after single B-vitamins additions, suggesting that  
503 inorganic nutrient availability enhance B-vitamin production by the prototrophic  
504 microbes. Under nutrient-limiting conditions, the external supply of vitamins could  
505 reduce the energy costs associated to its synthesis (Jaehme and Slotboom, 2015),  
506 stimulating the growth not only of auxotrophs but also of prototrophs.

507 The significant positive effects of B12 and/or B1 addition, suggest that these compounds  
508 may be eventually limiting microbial growth in marine productive ecosystems, as  
509 previously observed by other authors (e.g., Panzeca et al., 2006; Sañudo-Wilhelmy et al.,  
510 2006; Bertrand et al., 2007; Gobler et al., 2007; Koch et al., 2011; 2012; Barber.-Lluch et  
511 al., 2019). Most positive responses to B vitamin amendments were observed in oceanic  
512 waters, where B12 concentration was significantly lower than in coastal waters (Fig. 4c).  
513 Unfortunately we lack B1 measurements in this study, but, according to previous field  
514 studies in other oceanographic regions, a similar pattern to that observed for B12 can be  
515 expected (Cohen et al., 2017; Sañudo-Wilhelmy et al., 2012; Suffridge et al., 2018). The  
516 overall low and stable concentration of B12 at both sampling locations suggests a high  
517 turnover time of this compound in these productive, well-lit waters. Rapid cycling of B12  
518 in surface waters may occur due to high biological uptake rates (Taylor and Sullivan,  
519 2008; Koch et al., 2012) and/or photochemical degradation (Carlucci et al., 1969;  
520 Juzeniene and Nizauskaite, 2013; Juzeniene et al., 2015). The measured B12  
521 concentrations were in the lower range reported for coastal sites, and similar to that found  
522 in the upwelling system off the California coast in the San Pedro Basin during winter,  
523 spring and summer (Panzeca et al., 2009).

524 The increase of Chl-*a* was mostly associated to B12 amendments, which is consistent  
525 with the known incapability of eukaryotes to synthesize this vitamin (Croft et al., 2005;

526 Tang et al., 2010; Sañudo-Wilhelmy et al., 2014). Considering the very low concentration  
527 of B12 in the sampling area, the relatively limited phytoplankton response to B vitamins  
528 suggests that the existing species might have adapted to overcome B12 shortage. For  
529 example, changes in external B12 availability may cause shifts from vitamin B12-  
530 dependence to vitamin B12-independence in taxa possessing the vitamin B12-  
531 independent methionine synthase (MetE) gene (Bertrand et al., 2013; Helliwell et al.,  
532 2014). Other strategies used by phytoplankton to cope with low B12 concentration  
533 include, increased cobalamin acquisition machinery, decreased cobalamin demand, and  
534 management of reduced methionine synthase activity through changes in folate and S-  
535 adenosyl methionine metabolism (Bertrand et al., 2012). The available data on B12 half-  
536 saturation constants for phytoplankton ( $0.1\text{--}10\text{ pmol l}^{-1}$ ) (Droop, 1968, 2007; Taylor and  
537 Sullivan, 2008; Tang et al., 2010; Koch et al., 2011) are similar or higher than the B12  
538 concentrations measured here ( $0.3\text{ pmol l}^{-1}$  in the coastal and  $0.15\text{ pmol l}^{-1}$  in the oceanic  
539 waters, on average), reinforcing the hypothesis of a phytoplankton community adapted to  
540 B12 limiting concentrations in this upwelling system.

541 The positive responses of phytoplankton in surface oceanic waters in February seemed to  
542 be associated with high abundance of *Synechococcus* and SAR11 (Fig. 4b and Fig. 8).  
543 *Synechococcus* produce a B12 analog known as pseudocobalamin, where the lower ligand  
544 base adenine replaces 5,6-dimethylbenzimidazole (DMB) (Helliwell et al., 2016). In  
545 natural conditions, pseudocobalamin is considerably less bioavailable to eukaryotic algae  
546 than other cobalamin forms (Helliwell et al., 2016; Heal et al., 2017). SAR11 do not  
547 require B12 and do not have pathways for its synthesis (Sañudo-Wilhelmy et al., 2014;  
548 Gómez-Consarnau et al., 2018), suggesting that B12 synthesis could be limited in oceanic  
549 waters in winter, due to the low abundance of potential B12 producers.

550 Microbial responses to B vitamins in subsurface oceanic waters in February were  
551 associated to high abundance of *Synechococcus* and, to some extent, of Actinobacteria  
552 (Fig. 8). In these experiments, positive effects of B1 addition on phytoplankton and  
553 prokaryotes were observed (Fig. 7). While *Synechococcus* is capable of B1 synthesis  
554 (Carini et al., 2014; Sañudo-Wilhelmy et al., 2014; Gómez-Consarnau et al., 2018),  
555 Actinobacteria seems to have a strong dependence on this vitamin (Gómez-Consarnau et  
556 al., 2018). Among the sequenced eukaryote genomes, only Stramenopiles contain genes  
557 codifying for the synthesis of thiamine monophosphate (Sañudo-Wilhelmy et al., 2014;  
558 Cohen et al., 2017). While Stramenopiles, dominated by Bacillariophyta, were ubiquitous  
559 in the sampling area, their relative contribution was lower in oceanic waters (Fig. 4a).  
560 The simultaneous stimulation of phytoplankton and prokaryotes by B1 addition in  
561 subsurface oceanic waters in winter suggest a strong demand for this compound under  
562 these particular conditions, however what triggers the observed responses remain unclear.  
563 Even though B1 caused a significant effect on phytoplankton only in subsurface waters  
564 in winter, half of the positive responses of prokaryotes were associated to B1 supply (Fig.  
565 7b). This pattern is consistent with the recently described widespread dependence of  
566 bacterioplankton on external B1 supply (Paerl et al., 2018). B1 stimulated prokaryote  
567 growth in subsurface coastal waters and surface oceanic waters in summer (Fig. 7b), when  
568 the B vitamin response patterns were associated to high abundance of *Planktomarina* and  
569 Actinobacteria (Fig. 8), which are expected to strongly depend on external B1 sources  
570 (Giebel et al., 2013; Gómez-Consarnau et al., 2018). The generalized significant and  
571 positive responses of prokaryotes to vitamin treatments in surface oceanic waters in  
572 summer, when the prokaryote biomass was high and dissolved inorganic nitrogen  
573 concentration was very low (Fig. 3i), suggest that prokaryotes may have an advantage in  
574 the uptake and assimilation of B vitamins under nitrogen limiting conditions. This is

575 consistent with the observation of small (0.7–3  $\mu\text{m}$ )-plankton cells containing more B1  
576 than larger cells (Fridolfsson et al., 2019). Following this, it has been speculated that  
577 bacteria and small phytoplankton can transfer B1 to large cells through predation by  
578 acting as an important source of this compound in the marine environment (Fridolfsson  
579 et al., 2019).

#### 580 **4.2 Negative responses to vitamin B1 and B12 amendments**

581 Similar experiments conducted in this area also reported negative responses of microbial  
582 plankton to vitamin B12 additions (Barber-Lluch et al., 2019). The predominantly  
583 negative prokaryote responses after vitamin amendments in the coast during summer (Fig.  
584 7b), when nutrient concentrations were low (Fig. 3), suggest either a strong competition  
585 between phytoplankton and prokaryotes or a stimulation of predation. Dinoflagellates  
586 were particularly abundant in summer at both sampling sites and depths. Many  
587 dinoflagellate species are auxotrophs for B1 and/or B12 (Croft et al, 2006; Tang et al.,  
588 2010), and also many of them are phagotrophs (Stoecker and Capuzzo, 1990; Smayda,  
589 1997; Sarjeant and Taylor, 2006; Stoecker et al., 2017), thus the external supply of B  
590 vitamins may have promoted their growth, ultimately leading to net decreases in  
591 microbial biomass at the end of the experiments. Several studies demonstrated that  
592 vitamin B12 is implicated in the occurrence of dinoflagellate blooms around the world  
593 (Aldrich, 1962; Carlucci and Bowes, 1970; Takahashi and Fukazawa, 1982; Yu and  
594 Rong-cheng, 2000). It has been suggested that the B12-dependent enzyme  
595 methylmalonyl-CoA mutase in dinoflagellate, euglenoid, and heterokont algae allows  
596 them to grow heterotrophically when B12 is available (Croft et al., 2006). Therefore, the  
597 B12 enrichment could trigger such nutritional strategy, particularly in summer, when  
598 mineral nutrients are less available, resulting in an increased predation pressure on  
599 prokaryotes.

600 The B vitamin response patterns in surface coastal waters in summer (Fig. 7), seemed to  
601 be associated with high abundance of Flavobacteriales (Fig. 8). All isolates of  
602 Bacteroidetes sequenced so far are predicted to be B12 auxotrophs (Sañudo-Wilhelmy et  
603 al., 2014; Gómez-Consarnau et al., 2018) and recent metatranscriptomic analyses reveal  
604 that B1 synthesis gene transcripts are relatively low in Flavobacteria as a group (Gómez-  
605 Consarnau et al., 2018). As both phytoplankton and prokaryotes are dominated by  
606 potentially B12 and B1 auxotrophs (dinoflagellates and Flavobacteriales) in the coast  
607 during summer (Fig. 4b), the negative responses could be the result of strong competition  
608 for B vitamins. However, the negative responses to B vitamins of both phytoplankton and  
609 prokaryotes in surface coastal water in summer suggests an increase in phytoplankton and  
610 prokaryote predation by mixotrophs rather than competition between them. By contrast,  
611 prokaryotes and phytoplankton showed opposite patterns of response to B vitamins in  
612 subsurface coastal waters in summer, which suggests competition between both microbial  
613 compartments (Fig. 7). While phytoplankton negatively responded only to single B  
614 vitamin additions, prokaryotes responded negatively only when both inorganic nutrients  
615 and B vitamins were added (Fig. 7). It is conceivable that phytoplankton had an advantage  
616 over prokaryotes when mineral nutrients were added. This hypothesis contrasts with  
617 previous studies reporting that B12 and B1 vitamin uptake is dominated by picoplankton  
618 (Koch et al., 2011, 2012), strongly suggesting that bacteria could outcompete larger  
619 phytoplankton for vitamin uptake. By contrast, Koch et al. (2014), found that carbon-  
620 specific B12 uptake by large phytoplankton was significantly lower during non-bloom  
621 (low nutrient concentration) compared to bloom conditions (high nutrient concentration),  
622 which suggests better competitive ability under nutrient-rich conditions.

## 623 **5 Conclusions**

624 In conclusion, our findings suggest that the heterogeneous responses of microbial  
625 plankton to B1 and B12 vitamins supply in this coastal upwelling system could be  
626 partially controlled by the composition of the prokaryote community, which is consistent  
627 with their previously reported major role as B12 producers and B1 consumers. Even  
628 though we lack data on B1 concentration, the overall moderate responses together with  
629 the low ambient B12 concentration, suggest that the microbial plankton community in  
630 this area could be well adapted to cope with B vitamin shortage and that a close balance  
631 exists between production and consumption of these important growth factors.

632

633 *Author contribution.*

634 Eva Teira designed the experiments and Vanessa Joglar carried them out with  
635 contributions from all co-authors. Vanessa Joglar analyzed the data, Vanessa and Eva  
636 Teira interpreted the results and Vanessa Joglar prepared the manuscript under Eva Teira  
637 supervision.

638 *Competing interests.* The authors declare that they have no conflict of interest.

639 *Acknowledgements*

640 We thank all the people involved in the project ENVISION for helping with sampling  
641 and analytical work. We also thank the crew of the Ramón Margalef for their help during  
642 the work at sea. From IIM-CSIC, V. Vieitez and M.J. Pazó performed the nutrient  
643 analyses. This research was supported by the Spanish Ministry of Economy and  
644 Competitiveness through ENVISION (CTM2014-59031-P) and INTERES (CTM2017-  
645 83362-R) projects. Vanessa Joglar was supported by a FPI fellowship from the Spanish  
646 Ministry of Economy and Competitiveness.

647

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983

984

985 **Table 1**

986

987

	Treatment	Nutrient included	Concentration
1.	Control (C)	No nutrient added	
2.	Inorganic nutrients (I)	$\text{NO}_3^-$	5 $\mu\text{mol l}^{-1}$
		$\text{NH}_4^+$	5 $\mu\text{mol l}^{-1}$
		$\text{HPO}_4^{2-}$	1 $\mu\text{mol l}^{-1}$
		$\text{SiO}_4^{2-}$	5 $\mu\text{mol l}^{-1}$
3.	Vitamin B12 (B12)	B12	100 pmol l <sup>-1</sup>
4.	Vitamin B1 (B1)	B1	600 pmol l <sup>-1</sup>
5.	B12 + B1	B12	100 pmol l <sup>-1</sup>
		B1	600 pmol l <sup>-1</sup>
6.	I + B12	$\text{NO}_3^-$	5 $\mu\text{mol l}^{-1}$
		$\text{NH}_4^+$	5 $\mu\text{mol l}^{-1}$
		$\text{HPO}_4^{2-}$	1 $\mu\text{mol l}^{-1}$
		$\text{SiO}_4^{2-}$	5 $\mu\text{mol l}^{-1}$
		B12	100 pmol l <sup>-1</sup>
7.	I + B1	$\text{NO}_3^-$	5 $\mu\text{mol l}^{-1}$
		$\text{NH}_4^+$	5 $\mu\text{mol l}^{-1}$
		$\text{HPO}_4^{2-}$	1 $\mu\text{mol l}^{-1}$
		$\text{SiO}_4^{2-}$	5 $\mu\text{mol l}^{-1}$
		B1	600 pmol l <sup>-1</sup>
8.	I + B12 + B1	$\text{NO}_3^-$	5 $\mu\text{mol l}^{-1}$
		$\text{NH}_4^+$	5 $\mu\text{mol l}^{-1}$
		$\text{HPO}_4^{2-}$	1 $\mu\text{mol l}^{-1}$
		$\text{SiO}_4^{2-}$	5 $\mu\text{mol l}^{-1}$
		B12	100 pmol l <sup>-1</sup>
		B1	600 pmol l <sup>-1</sup>

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992 **6 Tables and Figures**

993 **Table 1:** Eight different treatments were applied consisting of: (1) control treatment (C):  
994 no nutrients added; (2) inorganic (I) nutrient treatment: 5  $\mu\text{M}$  nitrate ( $\text{NO}_3^-$ ), 5  $\mu\text{M}$   
995 ammonium ( $\text{NH}_4^+$ ), 5  $\mu\text{M}$  silicate ( $\text{SiO}_4^{2-}$ ) and 1  $\mu\text{M}$  phosphate ( $\text{HPO}_4^{2-}$ ); (3) vitamin B12  
996 treatment: 100  $\text{pmol l}^{-1}$ ; (4) vitamin B1 treatment: 600  $\text{pmol l}^{-1}$ ); (5) ~~Inorganic-inorganic~~  
997 nutrients and vitamin B12 (I+B12) treatment; (6) Inorganic nutrients and vitamin B1  
998 (I+B1) treatment; (7) vitamins B12 and B1 (B12+B1) treatment and (8) Inorganic  
999 nutrients with vitamins B12 and B1 (I+B12+B1) treatment.

1000

1001 **Figure 1:** (a) The NW Iberian margin (rectangle) and locations of the stations that were  
1002 sampled in the Ría de Vigo (C) and on the shelf (Oc) (diamonds), (b) distribution of daily  
1003 coastal upwelling index (UI) and (c) registered precipitations during each sampling period  
1004 showing the initial time of each experiment (C-a, C-b, C-c and Oc-a, Oc-b, Oc-c).

1005

1006 **Figure 2:** Vertical distribution over time in the coastal station of Chl-*a* ( $\mu\text{g l}^{-1}$ ) in (a)  
1007 February, (b) April and (c) August; temperature ( $^{\circ}\text{C}$ ) in (g) February, (h) April and (i)  
1008 August; and salinity (PSU) in (m) February, (n) April and (o) August. Vertical  
1009 distribution over time in the oceanic station of Chl-*a* ( $\mu\text{g l}^{-1}$ ) in (d) February, (e) April  
1010 and (f) August; temperature ( $^{\circ}\text{C}$ ) in (j) February, (k) April and (l) August; and salinity  
1011 (PSU) in (p) February, (q) April and (r) August Dots show the  $t_0$  of the experiments. Chl-  
1012 *a*: Chlorophyll-*a* concentration.

1013

1014 **Figure 3:** Initial biological conditions and abiotic factors at the coastal and oceanic  
1015 sampling stations. Each bar corresponds to one of the 3 experiments performed in each  
1016 depth and station during February, April and August. (a, b, c), Chl-*a*, total Chl-*a* ( $\mu\text{g l}^{-1}$ ).

1017 Note that the y-axis is broken; (d, e, f) PB, prokaryote biomass ( $\mu\text{g C l}^{-1}$ ); (g, h, i) DIN,  
1018 dissolved inorganic nitrogen ( $\mu\text{mol l}^{-1}$ ) and (j, k, l) DIN:DIP, ratio inorganic  
1019 nitrogen:phosphate. The blue line shows the Redfield ratio (16:1) and SCM refers to the  
1020 sub-surface chlorophyll maximum. Chl-*a*: Chlorophyll-*a* concentration.

1021

1022 **Figure 4:** Averaged relative contribution of reads to the major taxonomic groups of (a)  
1023 eukaryotes and (b) prokaryotes at surface (surf) and SCM in the coastal and oceanic  
1024 station in February, April and August. (c) Averaged B12 concentration ( $\text{pmol l}^{-1}$ ) at  
1025 surface (surf) and SCM in the coastal and oceanic station in February, April and August.  
1026 Error bars represent standard error. SCM refers to the sub-surface chlorophyll maximum.

1027

1028 **Figure 5:** Chlorophyll-*a* concentration ( $\mu\text{g l}^{-1}$ ) in the t0 of each experiment (striped bars)  
1029 and in the endpoint of each treatment (colored bars) in the experiments conducted at (a)  
1030 surface and (b) SCM in the coastal and at (c) surface and (d) SCM in the oceanic station  
1031 in February, April and August. Error bars represent standard error. Note that the y-axis is  
1032 broken. SCM: sub-surface chlorophyll maximum.

1033

1034 **Figure 6:** Prokaryote biomass ( $\mu\text{g C l}^{-1}$ ) in the t0 of each experiment (striped bars) and in  
1035 the endpoint of each treatment (colored bars) in the experiments conducted at (a) surface  
1036 and (b) SCM in the coastal and at (c) surface and (d) SCM in the oceanic station in  
1037 February, April and August. Error bars represent standard error. Note that the y-axis is  
1038 broken. SCM: sub-surface chlorophyll maximum.

1039

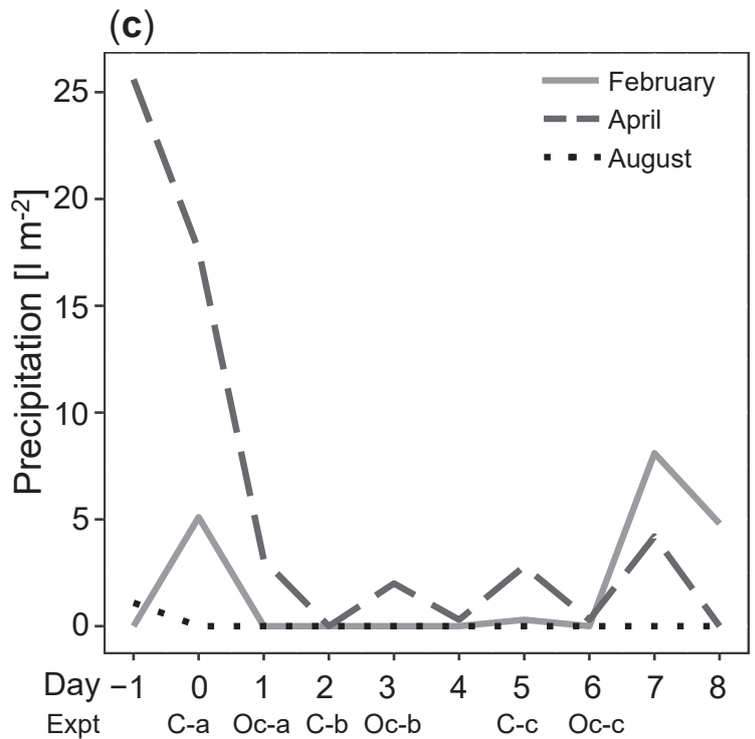
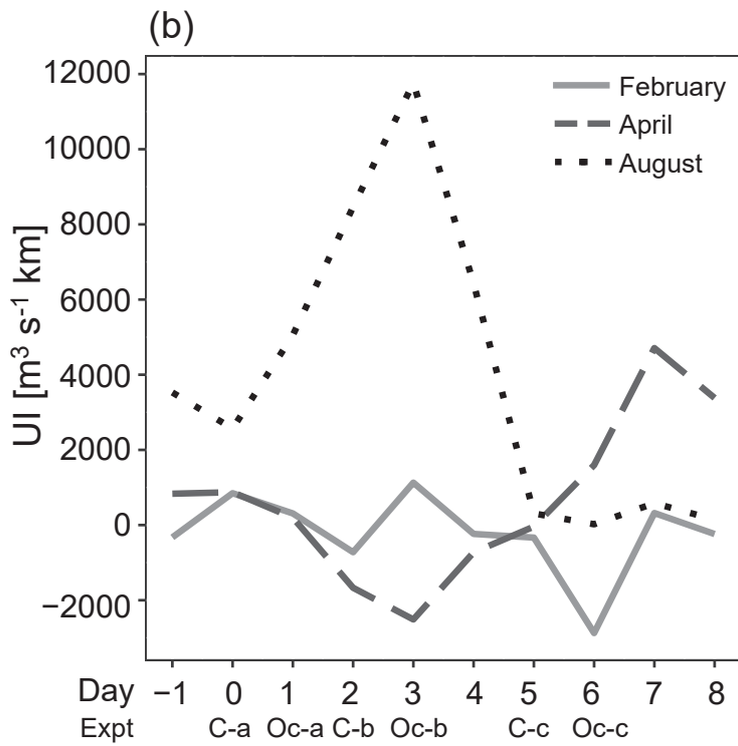
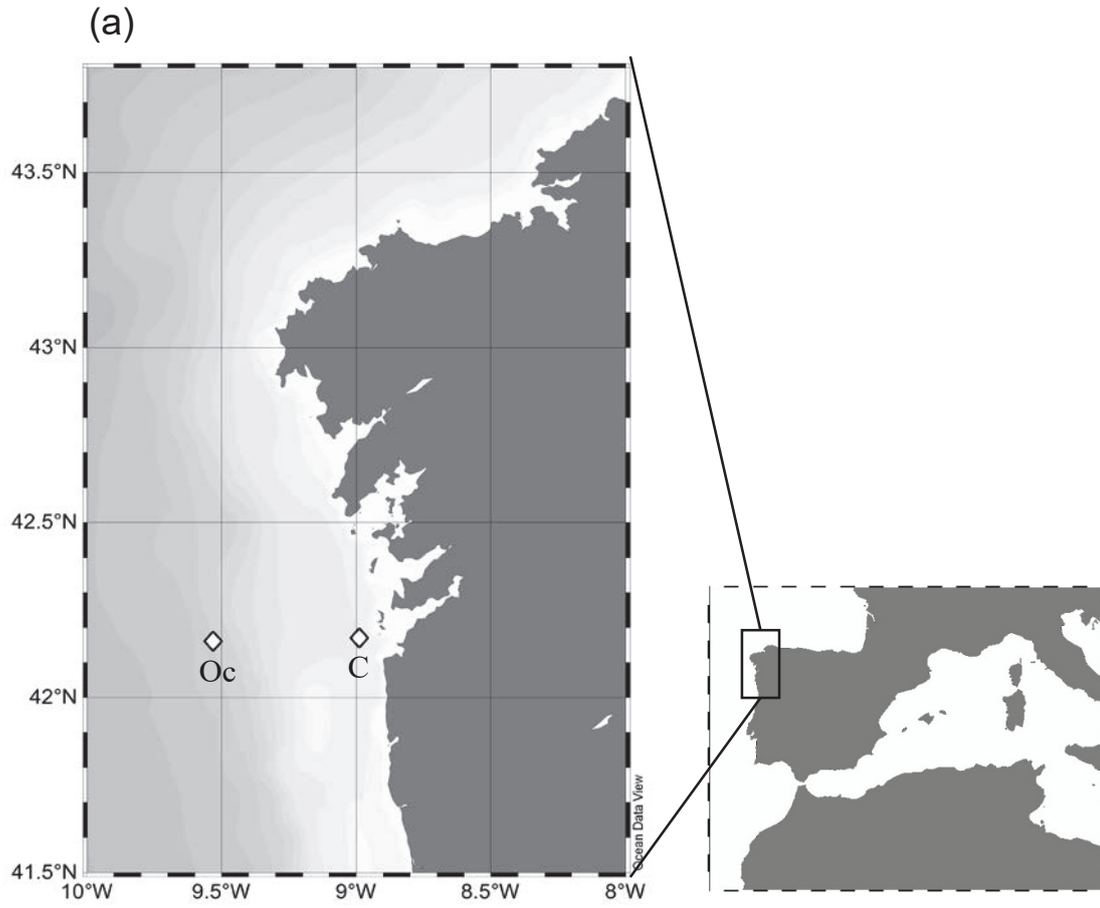
1040 **Figure 7:** Monthly averaged response ratio (RR) of (a) Chl-*a* or (b) prokaryote biomass  
1041 at surface and SCM in the coastal and oceanic station. Horizontal line represents a

1042 response equal to 1, that means no change relative to control in the pink dots (treatments  
1043 with vitamins alone) and no change relative to inorganic (I) treatment in the green dots  
1044 (vitamins combined with I treatments). Asterisks indicate averaged RRs that were  
1045 significantly different from 1 (Z-test; \*  $p < 0.05$ ) and “a” symbols indicate averaged RRs  
1046 that were marginally significant (Z-test; <sup>a</sup>  $p = 0.05-0.06$ ). Error bars represent standard error.  
1047 SCM: sub-surface chlorophyll maximum.

1048

1049 **Figure 8:** Distance based redundancy analysis (dbRDA) of B vitamin responses by  
1050 phytoplankton and prokaryotes based on Bray-Curtis similarity. Only prokaryotic taxa  
1051 that explained variability in the B vitamin responses structure selected in the DistLM  
1052 model (step-wise procedure with adjusted  $R^2$  criterion) were fitted to the ordination.  
1053 Filled and open symbols represent samples from coastal and oceanic station, respectively,  
1054 triangles and circles represent samples from surface and SCM, respectively, and colours  
1055 correspond to the months: (green) February, (blue) April and (pink) August. SCM: sub-  
1056 surface chlorophyll maximum.

Fig. 01



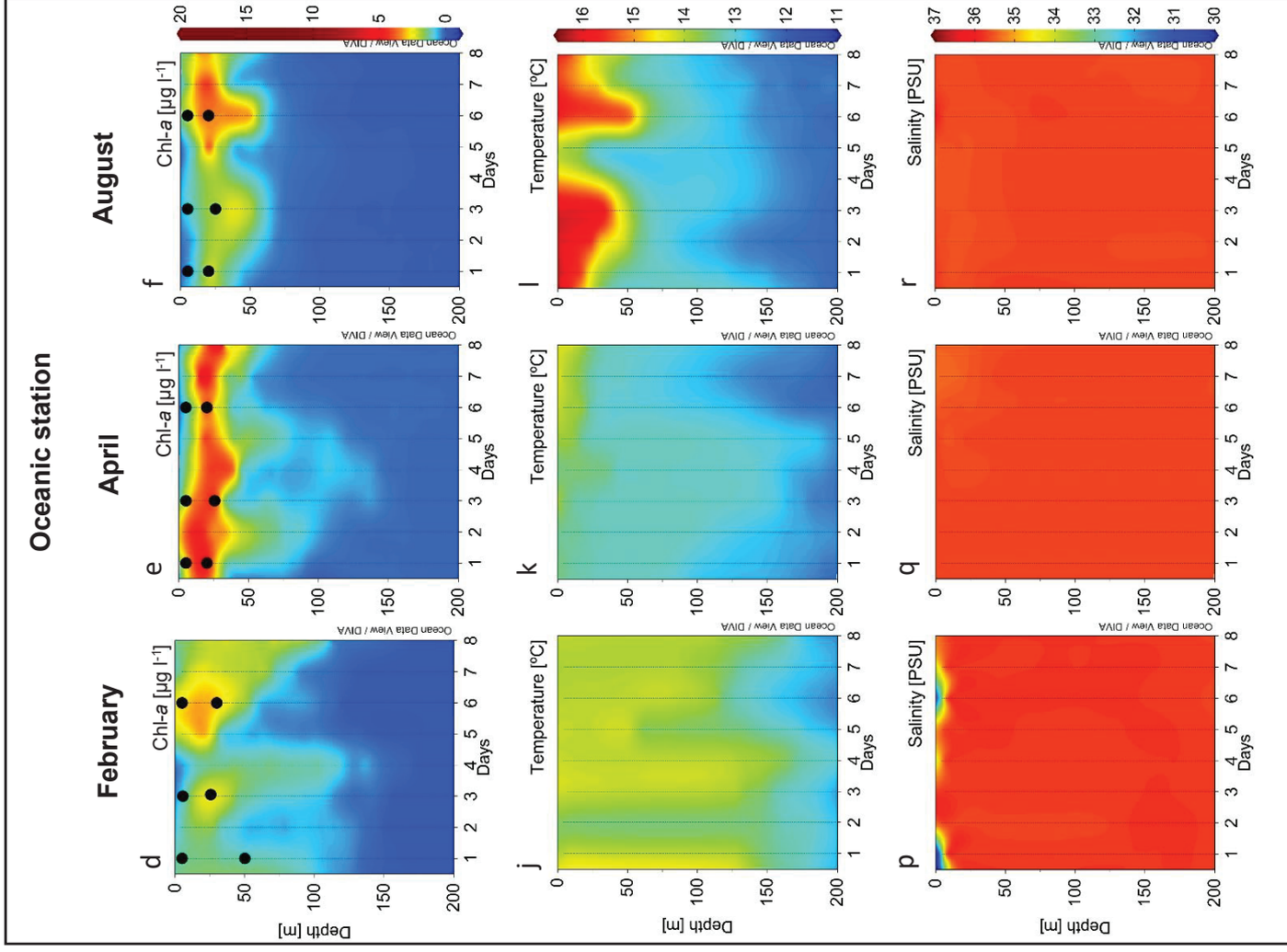
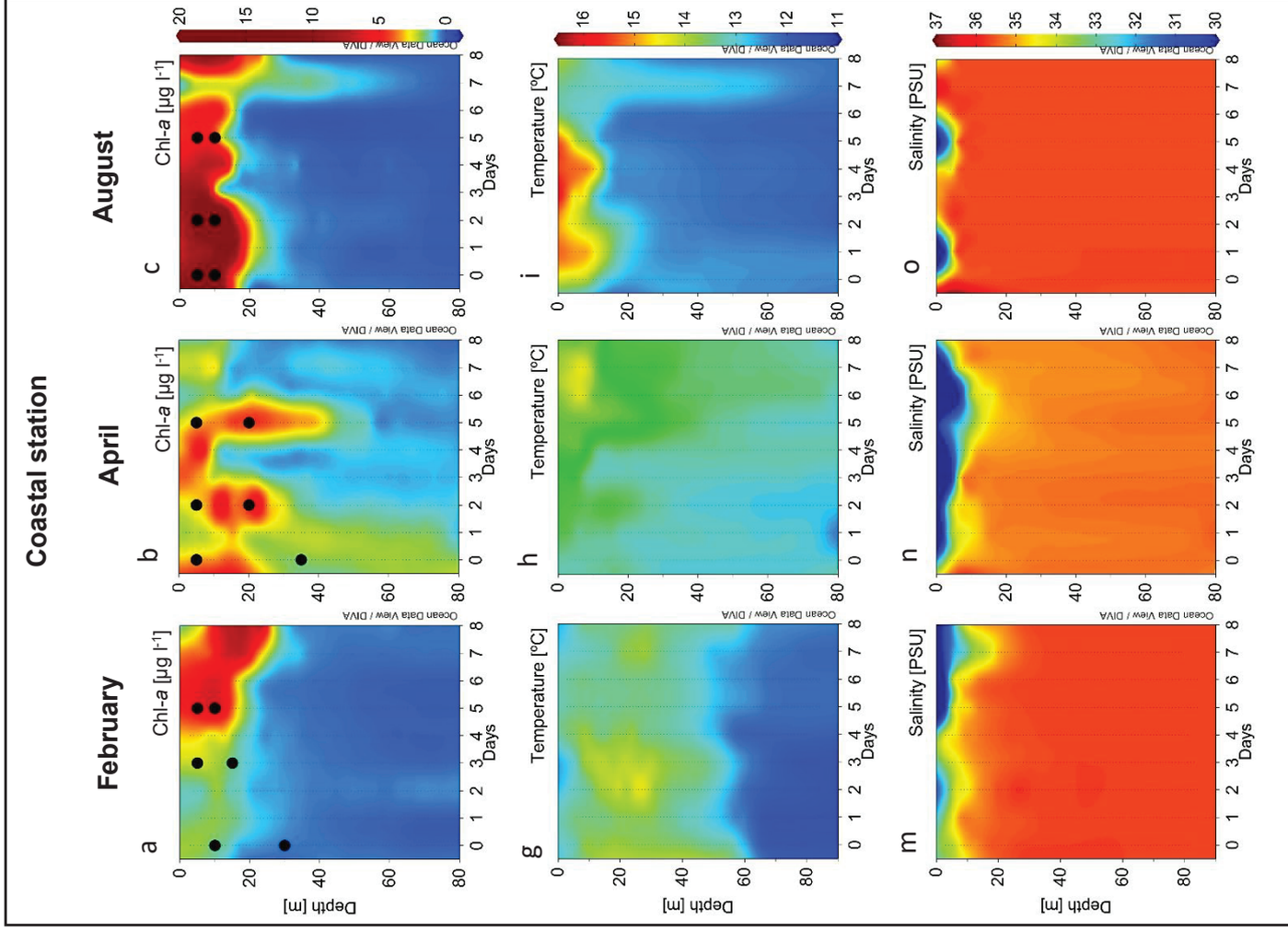


Fig. 03

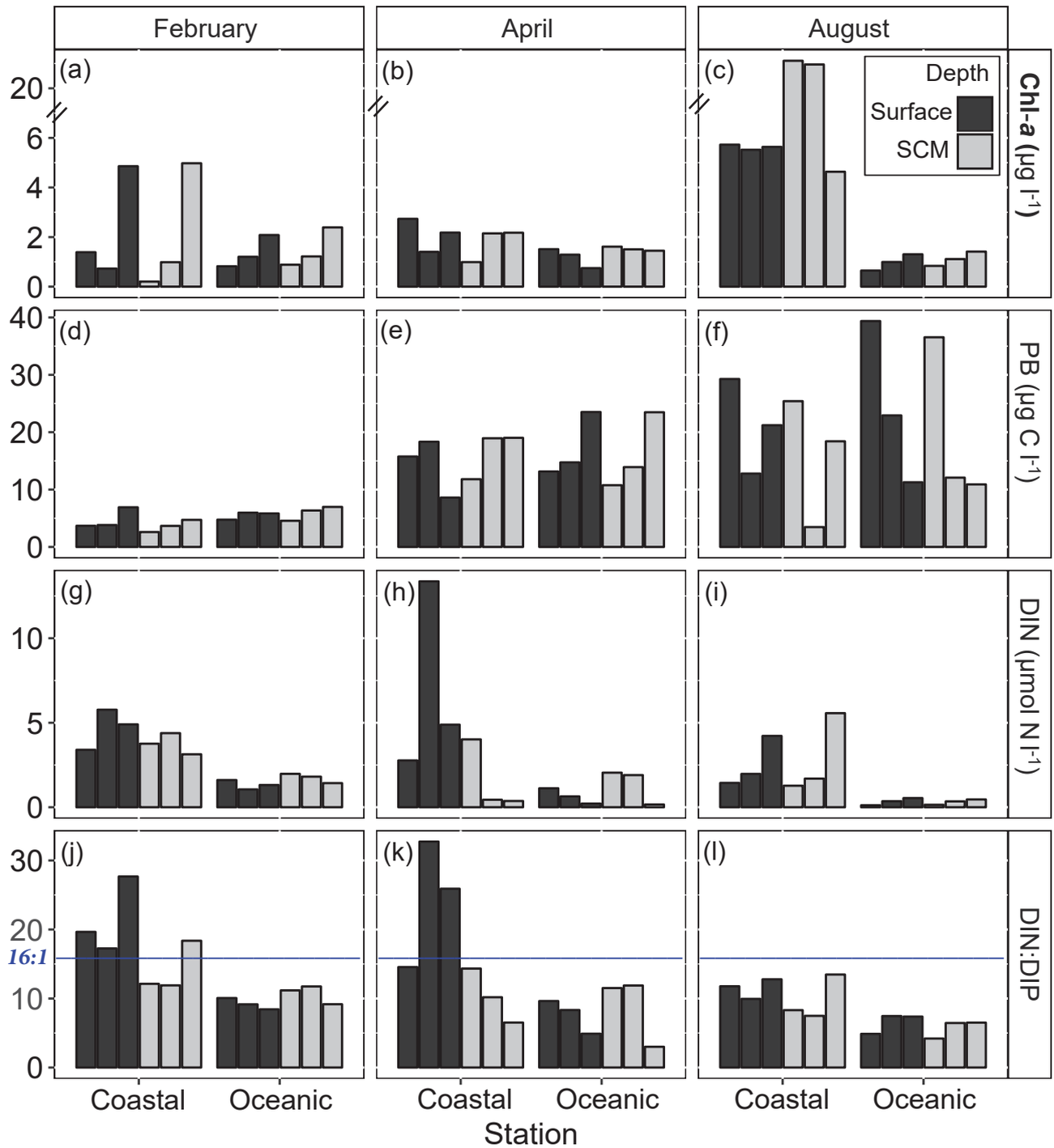
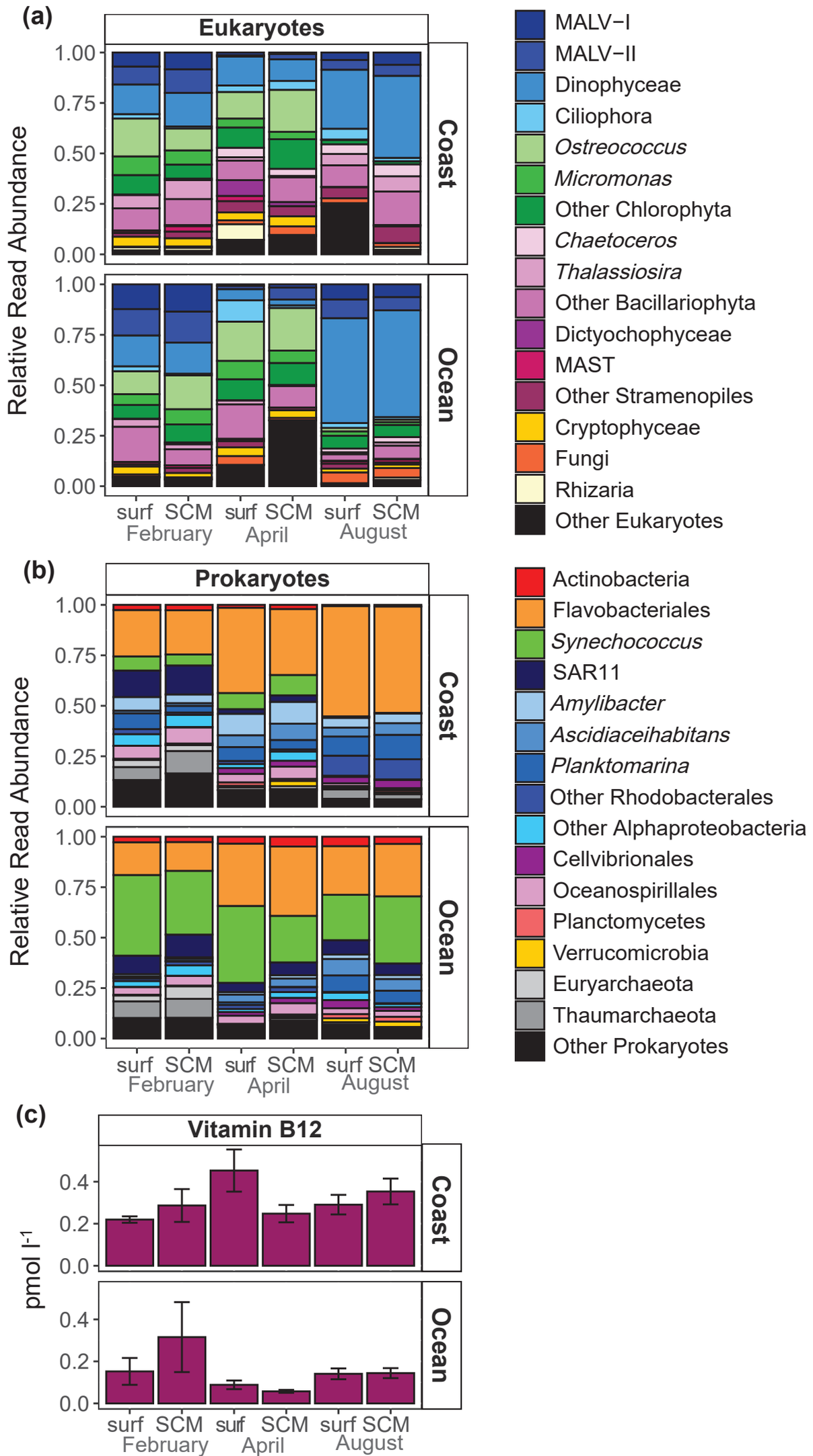
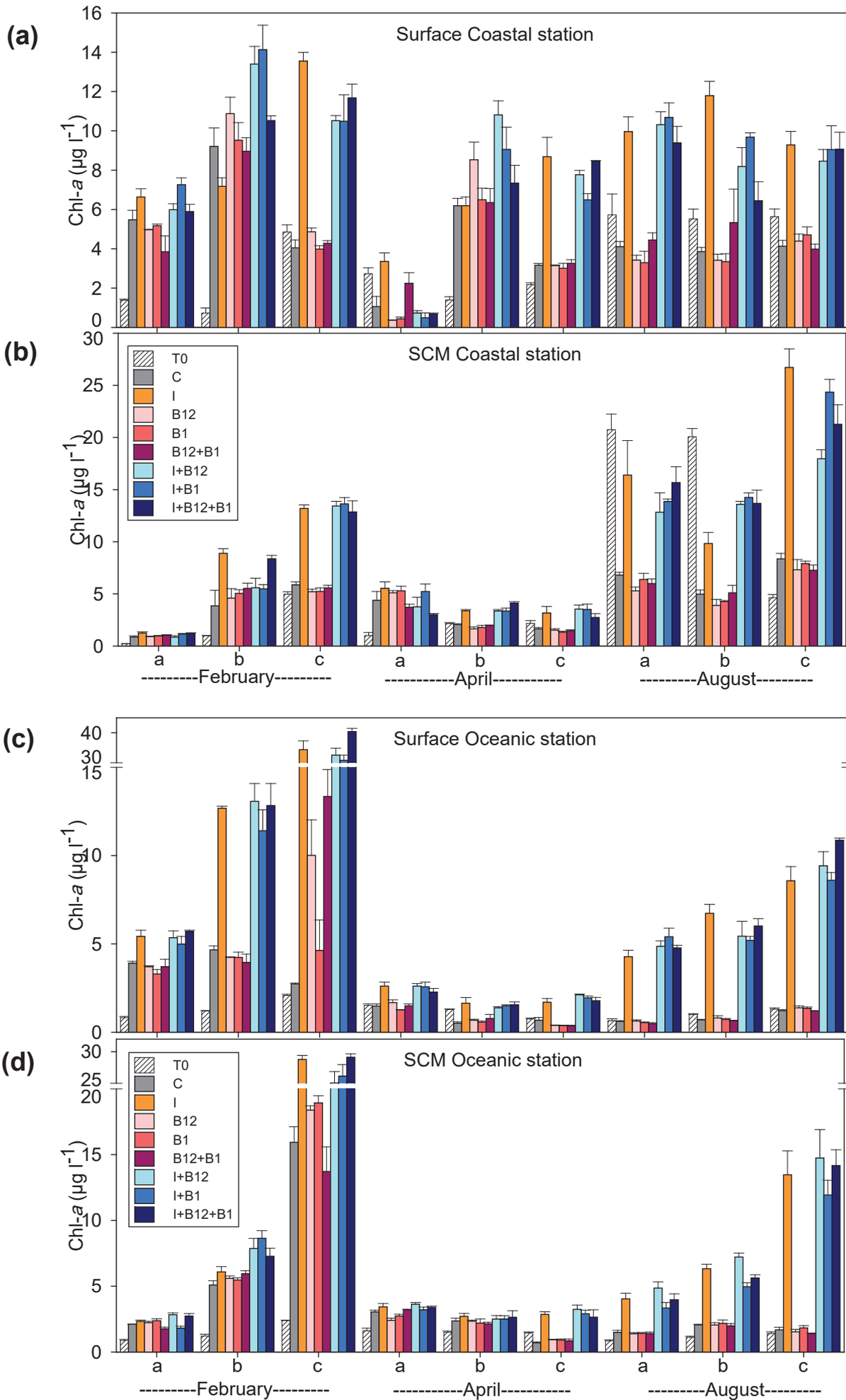
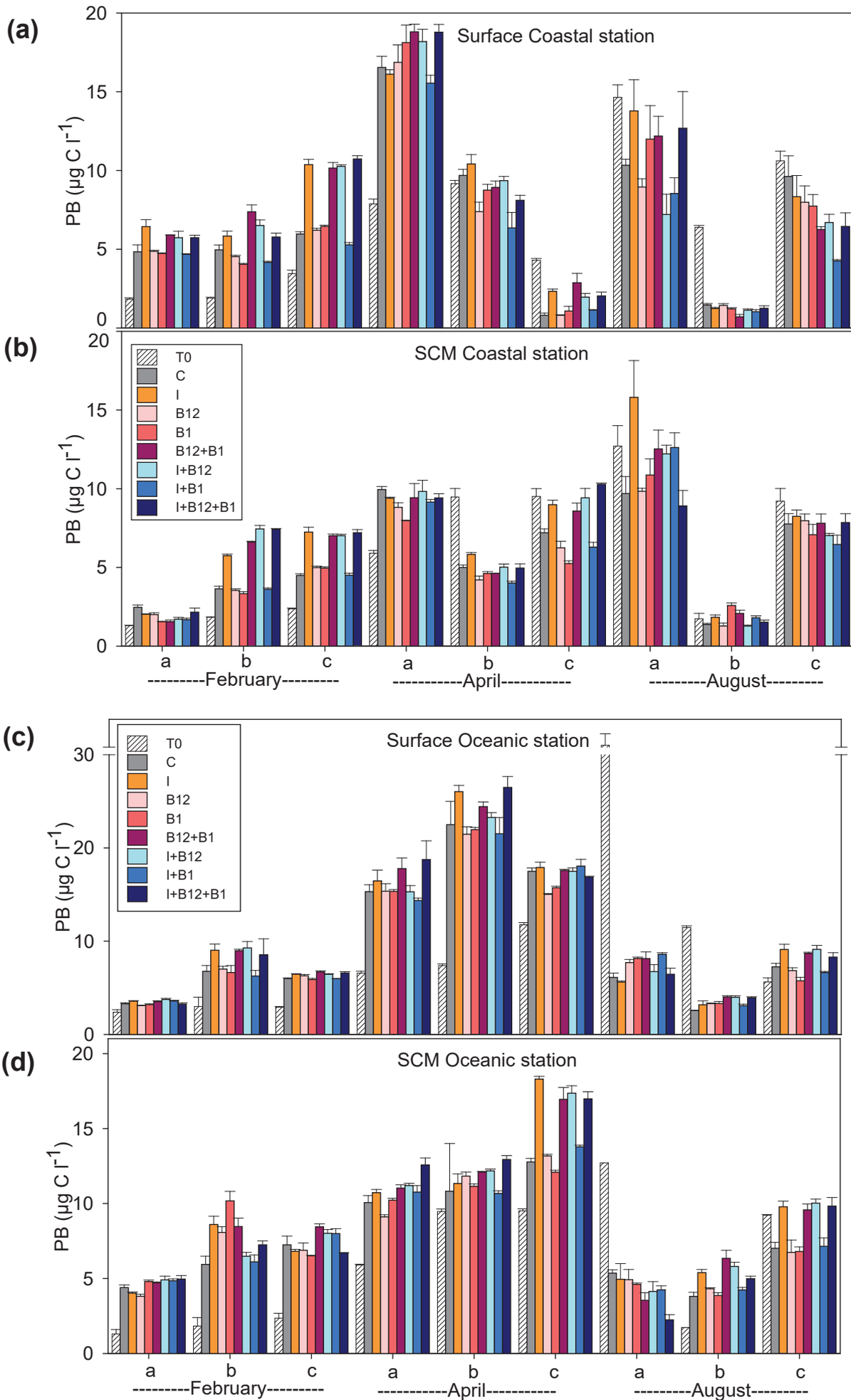
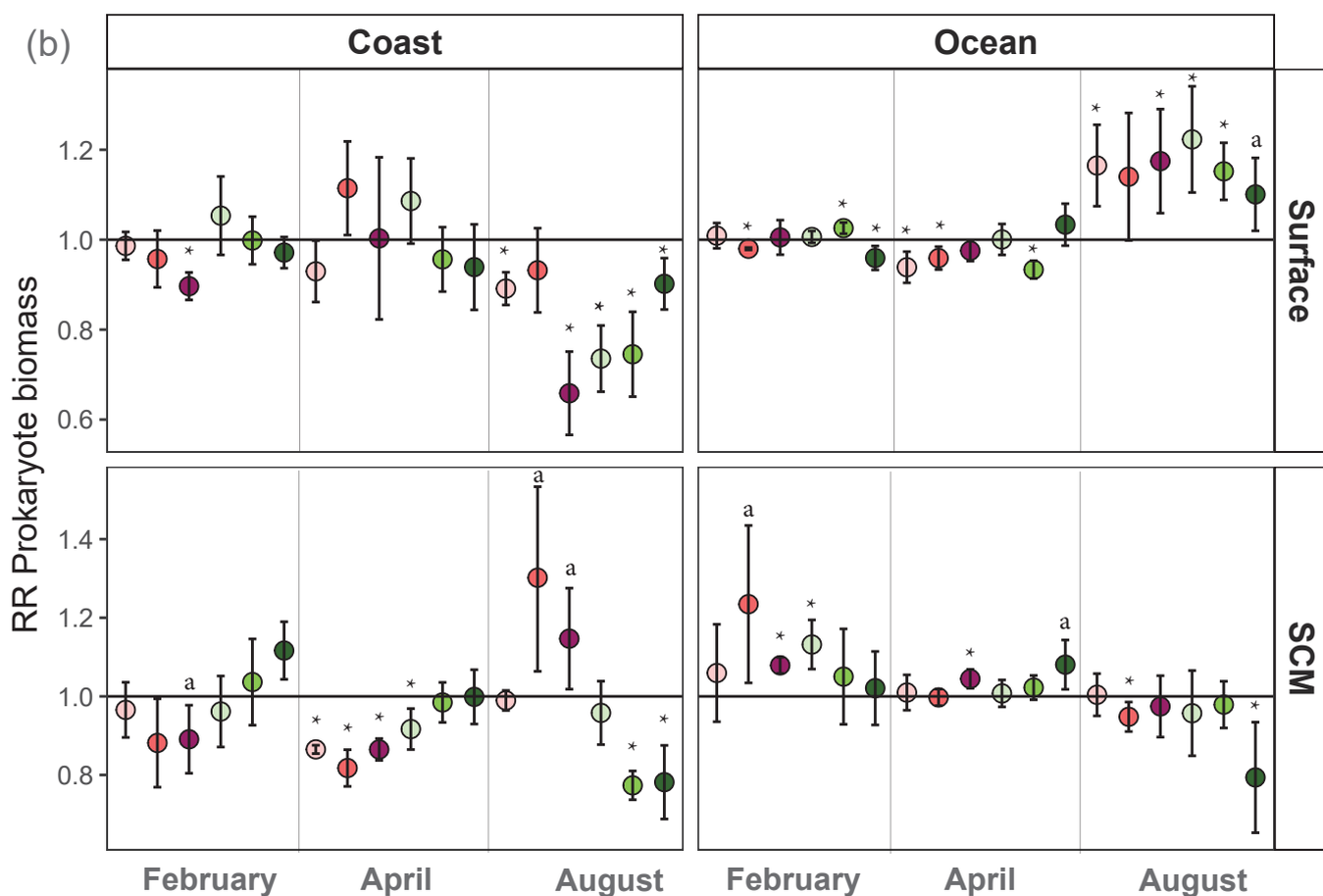
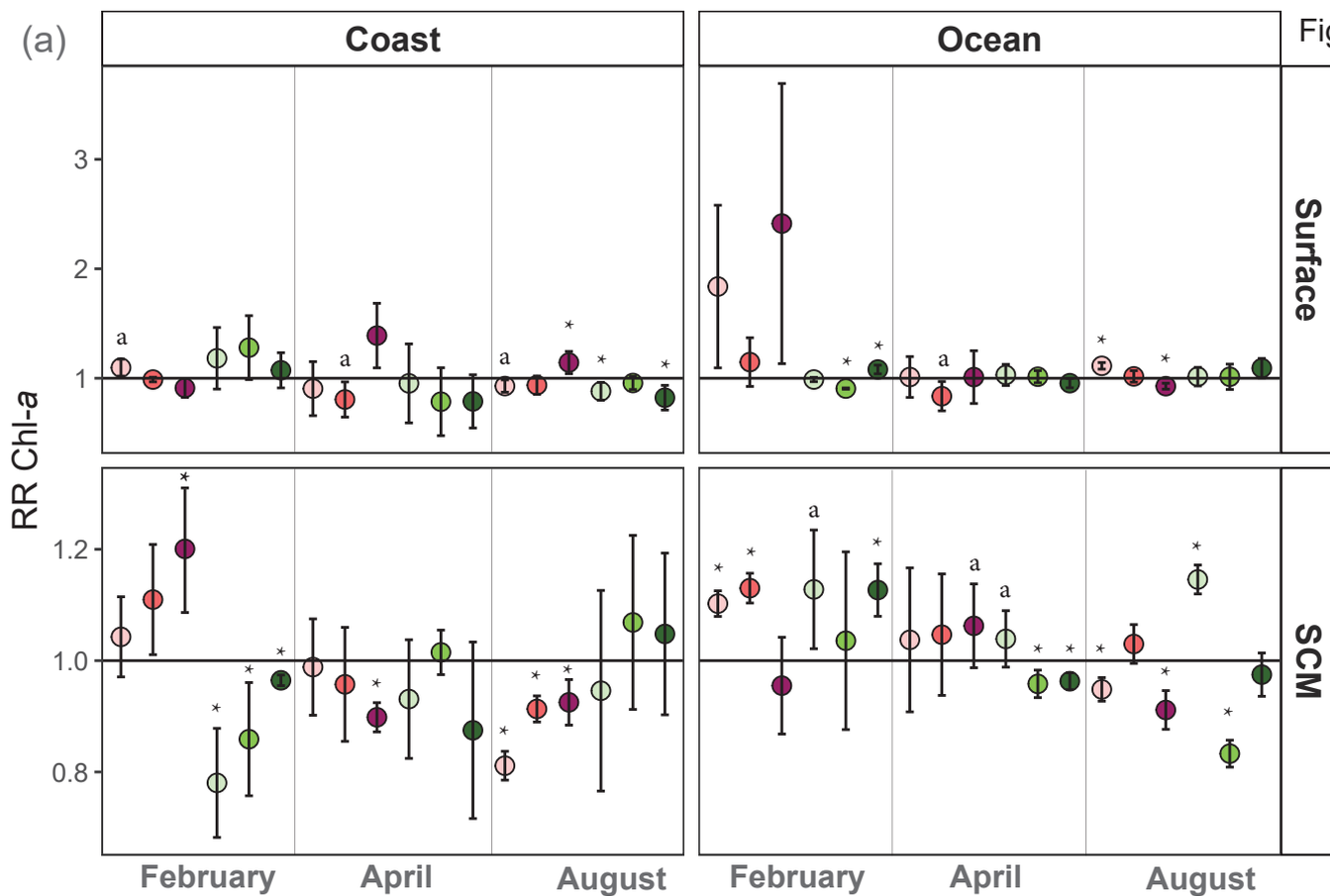


Fig. 04









Treatment

- B12 /C
- I+B12 /I
- B1 /C
- I+B1 /I
- B12+B1 /C
- I+B12+B1 /I

Fig. 08

