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4 **Seasonal cycling of zinc and cobalt in the Southeast Atlantic along the**  
5 **GEOTRACES GA10 section.**

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20

21 **Abstract**

22 We report the distributions and stoichiometry of dissolved zinc (dZn) and cobalt (dCo) in sub-  
23 tropical and sub-Antarctic waters of the Southeast Atlantic Ocean during austral spring 2010  
24 and summer 2011/12. In sub-tropical surface waters, mixed-layer dZn and dCo concentrations  
25 during early spring were  $1.60 \pm 2.58$  nM and  $30 \pm 11$  pM, respectively, compared with summer

26 values of  $0.14 \pm 0.08$  nM and  $24 \pm 6$  pM. The elevated spring dZn concentrations resulted from  
27 an apparent offshore transport of elevated dZn at depths between 20 – 55 m, derived from from  
28 the Agulhas Bank. In contrast, open-ocean sub-Antarctic surface waters displayed largely  
29 consistent inter-seasonal mixed-layer dZn and dCo concentrations of  $0.10 \pm 0.07$  nM and  $11 \pm$   
30  $5$  pM, respectively. Trace metal stoichiometry, calculated from concentration inventories,  
31 suggest a greater overall removal for dZn relative to dCo in the upper water column of the  
32 Southeast Atlantic with an inter-seasonally decreasing dZn/dCo inventory ratios of 19 to 5 mol  
33  $\text{mol}^{-1}$  and 13 to 7 mol  $\text{mol}^{-1}$  for sub-tropical surface water and sub-Antarctic surface water,  
34 respectively. In this paper, we investigate how the seasonal influences of external input and  
35 phytoplankton succession may relate to the distribution of dZn and dCo, and variation in  
36 dZn/dCo stoichiometry, across these two distinct ecological regimes in the Southeast Atlantic.

37

## 38 **1. Introduction**

39 The trace metal micronutrients zinc (Zn) and cobalt (Co) play an important role in the  
40 productivity of the oceans as key requirements in marine phytoplankton metabolism (Morel,  
41 2008; Twining and Baines, 2013). Zinc is required for the acquisition of inorganic carbon and  
42 organic phosphorus via the carbonic anhydrase and alkaline phosphatase metalloenzymes,  
43 respectively (Morel et al., 1994; Shaked et al., 2006; Cox and Saito, 2013). The requirement  
44 for Co stems from its obligation in the biosynthesis of vitamin B<sub>12</sub> (Raux et al., 2000; Rodionov  
45 et al., 2003) and, like Zn, its potential roles as a metal cofactor in carbonic anhydrase and  
46 alkaline phosphatase (Morel et al., 1994; Jakuba et al., 2008; Saito et al., 2017). Significantly,  
47 both dissolved Zn (dZn) and Co (dCo) are often scarce in surface seawater with mean  
48 concentrations that are often similar to, or relatively depleted, compared with typical cellular  
49 requirements of phytoplankton (Moore et al., 2013; Moore, 2016). Hence, dZn and dCo  
50 availability have the potential to regulate phytoplankton metabolism and growth rates in some

51 ocean regions (Sunda and Huntsman, 1992; Saito et al., 2002; Franck et al., 2003; Shaked et  
52 al., 2006; Bertrand et al., 2007; Jakuba et al., 2012; Mahaffey et al., 2014; Chappell et al., 2016;  
53 Browning et al., 2017).

54 The role for Zn and Co in carbonic anhydrase establishes an interaction between their ocean  
55 cycles, whereby biochemical substitutions between the enzyme-bound metals enables a  
56 stoichiometric plasticity in their cellular requirements that can negate the effect of limited  
57 availability. For example, a number of eukaryotic algae can substitute Zn for Co, as well as  
58 cadmium (Cd), in carbonic anhydrase when seawater dZn concentrations are low (Price and  
59 Morel, 1990; Sunda and Huntsman, 1995; Lane and Morel, 2000; Xu et al., 2007; Saito and  
60 Goepfert, 2008; Kellogg et al., 2020). In contrast, the prokaryotic picocyanobacteria  
61 *Synechococcus* and *Prochlorococcus* appear to have an absolute Co requirement (Sunda and  
62 Huntsman, 1995; Saito et al., 2002; Hawco and Saito, 2018). The availability and stoichiometry  
63 of dZn and dCo may therefore also exert a key control on phytoplankton community structure  
64 in some ocean regions (Leblanc et al., 2005; Saito et al., 2010; Chappell et al., 2016).

65 With the arrival of GEOTRACES research cruises, a number of studies have provided  
66 comprehensive data on the basin-scale distributions of Zn and Co in the Atlantic Ocean (e.g.  
67 Bown et al., 2011; Noble et al., 2012, 2017; Wyatt et al., 2014; Roshan et al., 2015; Middag et  
68 al., 2018). Such efforts have transformed our understanding of the biogeochemical processes  
69 associated with Zn and Co cycling (Saito et al., 2017; Vance et al., 2017; Weber et al., 2018;  
70 Tagliabue et al., 2018; Roshan et al., 2018) yet there are still geographically important regions  
71 of the Atlantic that remain largely understudied, including the Southeast Atlantic.

72 The Sub-Tropical Front (STF) of the Southeast Atlantic represents the convergence of warm,  
73 predominately macronutrient-limited Sub-Tropical Surface Water (STSW) and cold, iron-  
74 limited but macronutrient enriched sub-Antarctic Surface Water (SASW), creating one of the  
75 most dynamic nutrient regimes in the oceans (Ito et al., 2005; Browning et al., 2014; Moore,

76 2016). Here, the relative supply and availability of macronutrients and iron (Fe) exert an  
77 important control in maintaining the elevated phytoplankton stock and productivity that is  
78 typical of this frontal region, particularly during austral spring and summer (Moore and Abbott,  
79 2000; Ito et al., 2005; Browning et al., 2014). Dissolved Zn is also depleted in SASW that flows  
80 northwards to converge with STSW at the STF (Wyatt et al., 2014). However, the potential  
81 role for Zn in the mediation of phytoplankton distribution and community structure in this  
82 region is currently unclear.

83 Using data from two UK-GEOTRACES cruises (transect GA10) this study examines the  
84 seasonal availability and ecological stoichiometry of dZn and dCo, by analysis of their  
85 relationships with phosphate, in upper ocean waters of the Southeast Atlantic. These data,  
86 together with measurements of phytoplankton pigment biomass and community structure, offer  
87 an improved knowledge of the seasonal influences of external input and phytoplankton  
88 succession on the distribution and cycling of Zn and Co in these dynamic waters.

89

## 90 **2. Methods**

### 91 **2.1. Sampling methods**

92 Seawater samples were collected during two UK-GEOTRACES cruises in the South Atlantic  
93 Ocean (GA10, Fig. 1). The first cruise (D357) took place during austral spring 2010 (18th  
94 October to 22nd November 2010), sampling the Southeast Atlantic on-board the *RSS*  
95 *Discovery*. During D357, two transects were completed between Cape Town and the zero  
96 meridian that represent early austral spring (D357-1) and late austral spring (D357-2),  
97 respectively. The second cruise (JC068) took place during austral summer 2011/2012 (24th  
98 December 2011 to 27th January 2012), along the same transect of the first cruise and continuing  
99 along 40°S between Cape Town and Montevideo, Uruguay, on-board the *RSS James Cook*. For  
100 JC068, we present here only the repeat transect data between Cape Town and 13°W that

101 represents the Southeast Atlantic aspect of this transect. The stations occupied during the three  
102 transects were not identical, but rather represent a coverage of the Southern Ocean and sub-  
103 tropical waters present. Where stations were reoccupied during one or more transects, they  
104 have the same station number.

105 All sampling bottles were cleaned according to the procedures detailed in the GEOTRACES  
106 sample handling protocols (Cutter et al., 2010). Seawater and particulate samples below 15 m  
107 depth were collected using a titanium-frame CTD with 24 trace metal clean 10 L Teflon-coated  
108 OTE (Ocean Test Equipment) Niskin bottles deployed on a plasma rope. Sub-samples for  
109 dissolved trace metal analysis were filtered through 0.8/0.2  $\mu\text{m}$  cartridge filters (AcroPak™  
110 500, Pall) into 125 mL low density polyethylene bottles inside a class 1000 clean air container.  
111 Each sub-sample was acidified to pH 1.7 (0.024 M) by addition of 12 M hydrochloric acid  
112 (HCl, UpA, Romil) under a class 100 laminar flow hood. Vertical sampling for dissolved trace  
113 metals was augmented by surface samples collected at each station using a towed ‘fish’  
114 positioned at approximately 3-5 m depth. Fish samples were filtered in-line and acidified as  
115 described for samples collected from the titanium sampling system. Particulate samples were  
116 collected onto acid clean 25 mm, 0.45  $\mu\text{m}$ , polyethersulfone membrane disc filters (Supor®,  
117 Pall) and stored frozen (-20°C) until shore-based analysis.

118

## 119 **2.2. Trace metal analysis**

120 Dissolved Co was determined in the ISO accredited clean room facility (ISO 9001) at the  
121 University of Plymouth (UK) using flow injection with chemiluminescence detection,  
122 modified from the method of Cannizzaro et al. (1999) as described by Shelley et al. (2010).  
123 Briefly, dCo was determined in UV-irradiated samples using the reaction between pyrogallol  
124 (1,2,3-trihydrobenzene) and hydrogen peroxide formed in the presence of Co. Standards (20 –  
125 120 pM Co) were prepared in 0.2  $\mu\text{m}$  filtered low-dCo seawater ( $16.5 \pm 5.2$  pM,  $n = 15$ ) by

126 serial dilution of a 1000 ppm Co ICP-MS standard (Romil, UK). The accuracy of the analytical  
127 method was validated by quantification of dCo in SAFe (S and D2) and GEOTRACES (GD)  
128 reference seawater (Table 1). There was no detectable analytical dCo blank and the limit of  
129 detection ( $3\sigma$  of the lowest concentration standard) was  $1.98 \pm 0.87$  pM ( $n = 15$ ).

130 Dissolved Zn was determined using flow injection coupled with fluorescence detection,  
131 modified from the method of Nowicki et al. (1994) and described previously for this  
132 GEOTRACES section by Wyatt et al. (2014). The accuracy of the analytical method was  
133 validated by quantification of dZn in SAFe (S and D2) reference seawater (Table 1). The blank  
134 for dZn FIA was  $0.14 \pm 0.13$  nM and the limit of detection ( $3\sigma$  of the lowest concentration  
135 standard) was  $0.01 \pm 0.01$  nM ( $n = 15$ ).

136 Measurement uncertainties were estimated after the Nordtest approach (Worsfold et al., 2019)  
137 where a combined uncertainty ( $u_c$ ) is computed from day-to-day within-lab reproducibility and  
138 uncertainties associated with the determination of reference materials (Table 1). This approach  
139 creates higher uncertainties than those previously published for dZn and dCo analyses but  
140 provides a more realistic estimation of analytical uncertainty. During this study, the  $u_c$  for dZn  
141 and dCo analysis was 22 and 19 %, respectively, similar to the 13 – 25 % reported by Rapp et  
142 al. (2017) for the determination of trace metals, including dZn and dCo, by on-line pre-  
143 concentration and high-resolution sector field ICP-MS detection. The elevated  $u_c$  within our  
144 data results from the greater uncertainty surrounding the very low dZn and dCo concentration  
145 SAFe S reference sample whereas the dZn and dCo  $u_c$  using only the Safe D2 are <5 %.

146 Hereafter, when presenting low dZn and dCo concentrations for comparison with  
147 phytoplankton biological requirements (Section 3.5), we apply a fixed  $u_c$  of 20 % to our data.

148 Total particulate trace metals (i.e. pZn, pCo, pTi) were determined using inductively coupled  
149 plasma-mass spectrometry (Thermo Fisher XSeries-2) following a sequential acid digestion  
150 modified from Ohnemus et al. (2014). Potential interferences (e.g.  $^{40}\text{Ar}^{16}\text{O}$  on  $^{56}\text{Fe}$ ) were

151 minimized through the use of a collision/reaction cell utilizing 7 % H in He and evaluation of  
152 efficiency and accuracy assessed using Certified Reference Material (CRM). Full details of the  
153 method and CRM results can be found in Milne et al. (2017).

154

### 155 **2.3. Nutrients, phytoplankton, temperature and salinity**

156 The dissolved macronutrients phosphate ( $\text{PO}_4^{3-}$ ), silicic acid ( $\text{Si}(\text{OH})_4$  but referred to as Si  
157 hereafter) and nitrate (determined as nitrate + nitrite,  $\text{NO}_3^-$ ) were determined in all samples for  
158 which trace metals were determined, in addition to samples collected from a stainless steel  
159 rosette. Macronutrients were determined using an AA III segmented-flow AutoAnalyzer (Bran  
160 & Luebbe) following colorimetric procedures (Woodward and Rees, 2001). Salinity,  
161 temperature and depth were measured using a CTD system (Seabird 911+) whilst dissolved  $\text{O}_2$   
162 was determined using a Seabird SBE 43  $\text{O}_2$  sensor. Salinity was calibrated on-board using  
163 discrete samples taken from the OTE bottles and an Autosal 8400B salinometer (Guildline)  
164 whilst dissolved  $\text{O}_2$  was calibrated using a photometric automated Winkler titration system  
165 (Carritt and Carpenter, 1966). Mixed-layer depths (MLD) were calculated using the threshold  
166 method of de Boyer Montégut et al. (2014), where MLD is identified from a linear interpolation  
167 between near-surface density and the depth at which density changes by a threshold value  
168 ( $0.125 \text{ kg m}^{-3}$ ).

169 Measurements of phytoplankton pigment biomass and community structure were made on  
170 discrete samples collected using a 24 position stainless-steel CTD rosette equipped with 20 L  
171 OTE Niskin bottles. For chlorophyll-*a* analysis, samples were filtered ( $0.7 \mu\text{m}$  Whatman GF/F)  
172 and then the filters extracted overnight in 90 % acetone (Holm-Hansen et al., 1965). The  
173 chlorophyll-*a* extract was measured on a pre-calibrated (spinach chlorophyll-*a* standard,  
174 Sigma) Turner Designs Trilogy fluorometer. High performance liquid chromatography  
175 (HPLC) samples ( $0.5 - 2 \text{ L}$ ) for accessory pigment analyses were filtered ( $0.7 \mu\text{m}$  Whatman

176 GF/F), flash frozen in liquid nitrogen and stored at -80 °C prior to analysis using a Thermo  
177 HPLC system. The matrix factorization program CHEMTAX was used to estimate the  
178 contribution of taxonomic groups to total chlorophyll-*a* (Mackey et al., 1996). Concentrations  
179 of nanophytoplankton, *Synechococcus*, *Prochlorococcus* and photosynthetic picoeukaryotes  
180 were analysed by analytical flow cytometry (AFC) using a FACSort flow cytometer (Becton  
181 Dickenson, Oxford, UK) according to the methods described in Davey et al. (2008) and Zubkov  
182 et al. (2003).

183

### 184 **3. Results and Discussion**

#### 185 **3.1. Hydrographic setting and macronutrient distributions**

186 The prominent waters masses along the D357 and JC068 transects (Fig. 2) were identified by  
187 their characteristic thermohaline and macronutrient properties (Sarmiento et al., 2004; Ansoerge  
188 et al., 2005; Browning et al., 2014). Wyatt et al. (2014) provide a more detailed description of  
189 the JC068 hydrography along the entire GA10 section. Whilst we aim to compare the nearshore  
190 versus offshore distributions of micro- and macronutrients, note that sub-Antarctic mode water  
191 was not sampled for trace metals during the D357-2 late spring transect, and therefore only the  
192 early spring and summer values are discussed for SASW hereafter.

193

#### 194 ***Surface mixed-layer***

195 During all three transects the STF was identified by a sharp potential temperature ( $\theta$ ) gradient  
196 in the upper 200 m with the  $\theta$  15°C isotherm corresponding well to changes in macronutrient  
197 concentrations between STSW and SASW. North of the STF, mixed-layer macronutrient  
198 concentrations (Table 2) decreased in STSW between the three occupations of the transect. The  
199 largest relative depletion observed was for  $\text{NO}_3^-$  with a ~2.7-fold reduction in mean inventory  
200 concentration from 870 to 326  $\mu\text{mol m}^{-3}$  between early spring and summer, whilst  $\text{PO}_4^{3-}$  and



201 Si concentrations were reduced 1.5- and 1.4-fold, respectively. The largest absolute depletion  
202 was observed for Si with a reduction of  $848 \mu\text{mol m}^{-3}$  between early spring and summer.  
203 Conversely, summer SASW mixed-layer mean concentrations of  $\text{NO}_3^-$ ,  $\text{PO}_4^{3-}$  and Si were  
204 relatively 1.6, 1.4 and 2.1-fold lower than early spring, respectively, whilst the largest absolute  
205 depletion of  $1912 \mu\text{mol m}^{-3}$  was observed for  $\text{NO}_3^-$ . SASW mixed-layer concentrations of  $\text{NO}_3^-$   
206 and  $\text{PO}_4^{3-}$  were at least 2.1-fold higher than for STSW during the study, whilst the Si  
207 concentration was at least 1.5-fold lower, highlighting the relative deficiencies in major  
208 nutrients between high and low latitude derived surface waters (Sarmiento et al., 2004; Moore,  
209 2016).

210

### 211 *Sub-surface waters*

212 The Southern Ocean derived Sub-Antarctic Mode Water (SAMW) and underlying Antarctic  
213 Intermediate Water (AAIW) were identified using their characteristic core potential density  
214 ( $\sigma_\theta 26.8 \text{ kg m}^{-3}$ ) (Sarmiento et al., 2004; Palter et al., 2010) and thermohaline ( $S < 34.4$ ,  $\theta$   
215  $> 2.8^\circ\text{C}$ ) properties (Fig. 2). Wyatt et al. (2014) have identified these water masses along this  
216 section between 200 and 500 m. During all three transects, low sub-surface (50 – 500 m)  
217 macronutrient concentrations were observed between 13 and 16°E, associated with a salinity  
218 maxima. The feature conforms to the mean locality and depth range of Agulhas water  
219 (Duncombe Rae, 1991), clearly highlighting the penetration of Indian Ocean water into  
220 northward flowing SAMW.

221

## 222 **3.2. Zn and Co distributions of the Southeast Atlantic Ocean**

### 223 *Surface mixed-layer*

224 Figure 3 shows the dZn and dCo distributions for the upper 500 m of the Southeast Atlantic for  
225 the D357 and JCO68 transects. For full-depth dZn distributions along JC068 refer to Wyatt et

226 al. (2014). In the surface mixed-layer, dZn and dCo concentrations ranged from 0.01 to 4.57  
227 nM and 1 to 50 pM, respectively. The large range in dZn concentrations resulted from an  
228 apparent offshore transport of elevated dZn within STSW between 20 – 50 m during early  
229 spring (1.48 – 4.57 nM; Stns. 1 – 2) that was reduced by late spring (0.48 – 1.76 nM; Stns. 0.5  
230 – 1.5) and was absent during summer (0.01 – 0.13 nM; Stns. 1 – 2). Similarly, but to a lesser  
231 extent, elevated dCo concentrations were observed in STSW between 10 and 50 m during early  
232 and late spring (15 – 50 pM), compared with summer (18 – 33 pM). Our findings are consistent  
233 with previous observations of elevated dissolved and particulate trace metals over the same  
234 depth range in waters close to South Africa, including Co, Fe, Mn, and Pb (Chever et al., 2010;  
235 Bown et al., 2011; Boye et al., 2012; Paul et al., 2015). We postulate that these trace metal  
236 enrichments can arise from either atmospheric inputs, and/or from the lateral advection of  
237 metal-enriched waters from the Agulhas Current (AC) and/or South African continental shelf,  
238 and discuss this further in Sect. 3.3. In SASW, mixed-layer dZn and dCo concentrations ranged  
239 from 0.01 to 0.25 nM and 3 to 18 pM, respectively, during the study, significantly lower than  
240 STSW values, with the lowest concentrations observed during the summer transect (Table 2).

241

#### 242 *Sub-surface waters*

243 During the early spring D357-1 transect, elevated dZn and dCo concentrations were observed  
244 between the surface mixed-layer and 500 m (1.48–3.85 nM and 39–62 pM, respectively) at the  
245 station closest the South African continent (Stn. 1). Here, the highest dZn concentrations were  
246 associated with the dZn-enriched waters (20–55 m) described above for the surface mixed-  
247 layer. During the late spring D357-2 transect, the near-shore (Stns. 0.5–1) dZn concentrations  
248 were lower (0.31–1.76 nM) whilst dCo remained similar to early spring values (27–57 pM).  
249 During summer, near-shore (Stn. 1) sub-surface dZn concentrations were markedly lower  
250 (0.03–0.50 nM) than spring values whilst dCo concentrations (17–52 pM) were only

251 marginally lower. In offshore waters, sub-surface dZn concentrations ranged from 0.01 to 1.01  
252 nM across all three transects with extremely low values in the upper 400 m ( $0.22 \pm 0.21$  nM)  
253 and the highest values between 400 and 500 m. The absence of a significant return path for  
254 dZn with SAMW to waters above 400 m at this latitude (Wyatt et al., 2014; Vance et al., 2017)  
255 is likely an important control on dZn distributions across all three transects. In contrast, dCo  
256 concentrations were depleted in the upper 200 m (1–35 pM) and elevated in SAMW (23–56  
257 pM) suggesting that these Southern Ocean derived waters also play an important role in upper  
258 water column dCo distributions of the South Atlantic.

259 To assess whether seasonal changes in subsurface supply could influence dissolved Zn and Co  
260 concentrations in the upper water column of the Southeast Atlantic, we examined the metal  
261 versus  $\text{PO}_4^{3-}$  distributions of underlying SAMW and AAIW. Throughout this paper metal: $\text{PO}_4^{3-}$   
262 will be used to indicate an uptake remineralisation ratio derived from a regression slope, whilst  
263 metal/ $\text{PO}_4^{3-}$  will denote a concentration ratio. Figure 4 and supplementary table 1 show how  
264 the dZn: $\text{PO}_4^{3-}$  regression slope for SAMW and AAIW varied little between the three transects.  
265 These slopes are a function of the pre-formed micro- and macronutrient concentrations and the  
266 uptake/remineralisation ratio of the sources waters, as well as mixing during advection between  
267 the Southern Ocean and South Atlantic (Vance et al., 2017; Middag et al., 2018). The dZn: $\text{PO}_4^{3-}$   
268 slopes steepen with the introduction of AAIW with higher dZn/ $\text{PO}_4^{3-}$  concentration ratios, yet  
269 it is the relatively shallow slopes of overlying SAMW that imply a low, and relatively  
270 consistent, subsurface supply of dZn to STSW and SASW of the South Atlantic (Wyatt et al.,  
271 2014). The shallower waters overlying SAMW clearly show elevated dZn concentration,  
272 specifically during the spring transects, compared with what could be delivered if subsurface  
273 supply was the dominant source governing dZn availability in surface waters (Fig. 4). It is  
274 therefore unlikely that a change in subsurface supply from underlying SAMW is responsible  
275 for the change in dZn inventories of STSW and SASW between the three transects.

276 Similarly, the  $d\text{Co}:\text{PO}_4^{3-}$  regression slope varied little between the three transects (Fig. 4 and  
277 Supp. Table 1). In  $d\text{Co}:\text{PO}_4^{3-}$  space, a single slope can be fit to SAMW and AAIW with no net  
278 scavenging effect on dCo distribution over the upper 1000 m. Like dZn, the waters overlying  
279 SAMW displayed spring dCo concentrations elevated above that potentially delivered via  
280 SAMW supply. During summer however, SAMW may provide a subsurface source of dCo  
281 (Fig. 4c) to overlying waters highlighting how Southern Ocean derived waters may play  
282 important, yet different, roles in upper water column metal distributions of the Southeast  
283 Atlantic.

284

### 285 **3.3. Shelf derived sources of Zn and Co**

286 Potential sources of trace metals to surface waters of the Southeast Atlantic include  
287 atmospheric inputs from South Africa and Patagonia (Chance et al., 2015; Menzel Barraqueta  
288 et al., 2019) as well as interactions with shelf and slope waters of the Agulhas Bank (Bown et  
289 al., 2011; Boye et al., 2012; Paul et al., 2015). During the D357 spring transects, elevated  
290 mixed-layer dZn and dCo concentrations (up to 4.57 nM and 50 pM, respectively; Sect. 3.2)  
291 were observed at stations closest the Agulhas Bank shelf and slope (Stns. 0.5, 1, 1.5 and 2).  
292 Here, we compare these metal elevations with respect to the aforementioned sources. Firstly,  
293 we encountered only brief, light rain during the study, thus minimal wet deposition of  
294 atmospheric aerosol. By combining the median atmospheric dry deposition flux for soluble Zn  
295 and Co for the Southeast Atlantic (Zn 6.0 and Co 0.05  $\text{nmol m}^{-2} \text{d}^{-1}$ ; Chance et al., 2015) with  
296 the mean mixed-layer depth (34 m) for STSW during D357, dust dissolution is estimated to  
297 add approximately 5.5 and 0.05  $\text{nmol m}^{-3}$  dZn and dCo, respectively, over a one month period.  
298 These inputs are low compared with the mixed-layer metal inventories, representing <1 % of  
299 dZn and dCo concentration in STSW during the D357 transects (Table 2), and would not be  
300 sufficient to generate distinct mixed-layer maxima. It is likely, therefore, that the dZn and dCo

301 elevations originated from the advection of metal-enriched waters from the western Agulhas  
302 Bank, a region identified as a distinct source of both dissolved and particulate trace metals to  
303 the Southeast Atlantic (Chever et al., 2010; Bown et al., 2011; Boye et al., 2012; Paul et al.,  
304 2015), and/or from the leakage of Indian Ocean water into the Southeast Atlantic via the AC.  
305 The detachment of Agulhas rings and filaments from the AC during its retroflection back  
306 towards the Indian Ocean constitutes a source of Pb to the surface Southeast Atlantic along the  
307 D357 transects (Paul et al., 2015). Whilst we observed elevated mixed-layer dZn and dCo at  
308 ~15°E during both D357 transects, the absence of metal enrichment across the depth of the AC  
309 salinity maxima (Figs. 2 and 3) suggests that the signal must be entrained from elsewhere.  
310 Furthermore, dZn concentrations from the AC along the east coast of South Africa do not  
311 exceed 0.5 nM in the upper 200 m (Gosnell et al., 2012). It is therefore likely that the dZn and  
312 dCo enrichment was derived from the Agulhas Bank. The AC has been shown to meander over,  
313 and interact with, the Agulhas Bank, forming eddies and filaments on the shoreward edge of  
314 the AC proper, that tend to move northwards along the western shelf edge and into the  
315 Southeast Atlantic (Lutjeharms and Cooper, 1996; Lutjeharms, 2007), potentially delivering  
316 shelf-derived sedimentary material. We found no evidence of a fluvial signature in our data,  
317 and no significant fluvial source for trace elements to our study region has been reported in the  
318 literature. We focus here on the more likely scenario of sedimentary inputs as the driver of  
319 mixed-layer dZn and dCo elevations at the shelf and slope stations during D357. Despite no  
320 available particulate trace metal data for the D357-1 early spring transect for direct comparison  
321 with the highest dZn and dCo elevations, we observed elevated mixed-layer particulate Zn  
322 (pZn; 0.08–1.40 nM) and Co (pCo; 8–49 pM) at stations closest South Africa during the D357-  
323 2 late spring transect (Stns. 0.5, 1 and 1.5, Fig. S1), coincident with elevated dZn (0.05–1.82  
324 nM) and dCo (1–43 pM). Furthermore, for the upper 500 m at stations 0.5 and 1, we found  
325 strong positive correlations between particulate aluminium and titanium (pAl:pTi, slope 41.7

326 mol mol<sup>-1</sup>, Pearson's  $r$  0.99,  $n = 15$ ), as well as particulate Fe and titanium (pFe:pTi, slope 10.2  
327 mol mol<sup>-1</sup>, Pearson's  $r$  0.99,  $n = 15$ ), indicative of a strong lithogenic source. Whilst there are  
328 presently no South African sedimentary data against which we can compare our water column  
329 values, our pAl:pTi and pFe:pTi slope ratios are in excess of upper crustal mole ratios (34.1  
330 and 7.3 mol mol<sup>-1</sup>, respectively; McLennan, 2001). These 500 m ratios are also steeper than the  
331 aggregate slopes for the full depth Atlantic Ocean away from hydrothermal sources (32.1 and  
332 7.4 mol mol<sup>-1</sup>, Pearson's  $r > 0.97$ ,  $n = 593$ , Schlitzer, 2018). Given the refractory nature of  
333 lithogenic pTi across diverse oceanic environments (Ohnemus and Lam, 2015), this may  
334 suggest the resuspension and dissolution of Agulhas Bank sediments enriched in dAl and dFe,  
335 followed by westward offshore transport, a common feature of the Bank's physical circulation  
336 during spring and summer (Largier et al., 1992). Such processes may in turn provide an  
337 additional source of dZn and dCo to STSW of the Southeast Atlantic. For example, Little et al.  
338 (2016) proposed that oxygen-deficient, organic-rich, continental margin sediments may  
339 constitute a significant global sink within the marine Zn cycle. These sediments could  
340 additionally provide a local source of dZn following remineralisation. Recent model outputs  
341 have likewise highlighted oxygen-deficient, boundary sediments as a dominant external source  
342 of Co to the oceans (Tagliabue et al., 2018). Given that oxygen depleted (<45  $\mu$ M) bottom  
343 waters are prevalent across the western Agulhas Bank (Chapman and Shannon, 1987;  
344 Chapman, 1988), considered to arise from high organic matter input to sediments and its  
345 bacterial decomposition, a sedimentary source of dZn and dCo appears likely.

346

#### 347 **3.4. Trace metal stoichiometry of the upper Southeast Atlantic**

348 In addition to seasonal variations in the lateral advection of continentally derived trace metals,  
349 the lower dZn and dCo concentrations in STSW during summer, compared with spring, likely  
350 reflect differences in biological utilisation. Here, we examine the micro- and macronutrient

351 concentration inventories to assess the trace metal stoichiometry of the Southeast Atlantic over  
352 seasonal timescales. The data were grouped into STSW and SASW regimes, with STSW  
353 defined by  $\theta \geq 15^\circ\text{C}$ . This isotherm was located at a mean depth of  $144 \pm 96$  m across the study,  
354 compared with a mean mixed-layer depth of  $39 \pm 10$  m, and thus the inventories for SASW  
355 were determined over this depth for comparison with STSW (Table 2). Early and late spring  
356 STSW samples in the depth range 20 - 55 m that clearly exhibited continentally derived  
357 elevated dZn and dCo were removed from the analysis in order to compare stoichiometry with  
358 respect to biological processes. For SASW, micronutrient sampling did not occur during late  
359 spring and therefore only early spring and summer values are compared.

360 Distinct temporal trends in the stoichiometric relationship with  $\text{PO}_4^{3-}$  were evident for both dZn  
361 and dCo (Fig. 4). Within STSW, the dZn/ $\text{PO}_4^{3-}$  inventory ratio ranged from 699 to 1876  $\mu\text{mol}$   
362  $\text{mol}^{-1}$  (Table 2) with the highest value observed during early spring and the lowest during  
363 summer. Combined with summer dZn concentrations 4-fold lower than early spring, this  
364 suggests strong biological uptake of dZn alongside  $\text{PO}_4^{3-}$  between seasons. In contrast, lower  
365 dZn/ $\text{PO}_4^{3-}$  ratios of 372 and 188  $\mu\text{mol mol}^{-1}$  were observed in SASW during early spring and  
366 summer, respectively. Here, the absolute change in dZn concentration between spring and  
367 summer was lower than for STSW, but was greater for  $\text{PO}_4^{3-}$ , likely reflecting the increased  
368 availability of  $\text{PO}_4^{3-}$  in these Southern Ocean derived waters (Table 2) and an open-ocean  
369 phytoplankton community that have lower trace metal requirements than their counterparts  
370 north of the STF. Such dZn/ $\text{PO}_4^{3-}$  ratios sit at the lower end of cellular Zn/P reported for the  
371 diatom and haptophyte-type phytoplankton typical of this region ( $\sim 100 - 1100 \mu\text{mol mol}^{-1}$ ;  
372 Twining and Baines, 2013 and refs. therein), highlighting the importance of micronutrient  
373 processes with respect to Zn availability.

374 In contrast to dZn, the spatiotemporal variation observed for STSW dCo/ $\text{PO}_4^{3-}$  was small with  
375 ratios ranging from 82 to 129  $\mu\text{mol mol}^{-1}$  (Table 2), likely reflecting external inputs to the

376 oceans and biological Co requirements that are typically 4-fold less than for Zn (Ho et al.,  
377 2003; Roshan et al., 2016; Hawco et al., 2018). The STSW  $d\text{Co}/\text{PO}_4^{3-}$  ratio decreased between  
378 early and late spring transects, potentially in part due to the westward expansion of STSW  
379 during late spring (Fig. 2) and subsequent mixing with SASW depleted in dCo relative to  $\text{PO}_4^{3-}$   
380 (Fig. 3). This dilution is likely also true of dZn and Si, yet their STSW concentration inventories  
381 may be sufficiently high as to mask this effect. Unfortunately, an insufficient quantity of late  
382 spring SASW data are available with which to affirm this postulation. The highest  $d\text{Co}/\text{PO}_4^{3-}$   
383 ratio was observed during summer due to the preferential biological removal of  $\text{PO}_4^{3-}$  relative  
384 to dCo.

385 In SASW,  $d\text{Co}/\text{PO}_4^{3-}$  was consistently low with ratios of 23 and 26  $\mu\text{mol mol}^{-1}$  for early spring  
386 and summer, respectively. Much higher inventory ratios of  $\sim 580 \mu\text{mol mol}^{-1}$  can be calculated  
387 over similar depths for open-ocean North Atlantic waters (GA03 Stns. 11-20, Schlitzer et al.,  
388 2018), likely reflecting an elevated atmospheric Co input and/or an extremely low surface  $\text{PO}_4^{3-}$   
389 inventory (Wu et al., 2000; Martiny et al., 2019).

390 Our results provide evidence for the greater availability and preferential removal of dZn  
391 relative to dCo in the upper water column the Southeast Atlantic based on STSW dZn/dCo  
392 inventory ratios of 19, 17 and 5  $\text{mol mol}^{-1}$  for the three transects and SASW ratios of 13 and 7  
393  $\text{mol mol}^{-1}$  for early spring and summer, respectively (Table 2). With relatively consistent inter-  
394 seasonal dCo inventories for STSW and SASW, indicating a more balanced ecophysiological  
395 regime with regard to dCo organisation, the change in dZn/dCo inventory stoichiometries  
396 principally reflects changes in dZn concentration. We postulate that the inter-seasonal  
397 variations in dZn and dCo availability and stoichiometry of the Southeast Atlantic reflect  
398 changes in the relative nutritional requirement of resident phytoplankton and/or biochemical  
399 substitution of Zn and Co to meet nutritional demand.

400



### 401 3.5. Phytoplankton controls on trace metal ecological stoichiometry

402 Here we discuss the principle phenomena that together likely explain our observations of  
403 seasonally decreasing dZn/dCo inventory stoichiometries in STSW and SASW of the  
404 Southeast Atlantic: i.e. the preferential removal of dZn, relative to dCo, leading to low dZn  
405 availability, and differences in phytoplankton assemblages with different cellular metal  
406 requirements.

407 Satellite images show elevated surface chlorophyll concentrations across the Southeast Atlantic  
408 STF, compared with waters further north and south, with peak concentrations observed during  
409 summer in January 2012 (Fig. 1). Profiles of total chlorophyll-*a* concentration (Fig. S2) also  
410 show maximum summer values in the upper water column of STSW (1.02 mg m<sup>-3</sup>) and SASW  
411 (0.49 mg m<sup>-3</sup>) compared with spring values (<0.61 and <0.36 mg m<sup>-3</sup>, respectively). This is  
412 consistent with the hypothesis that increasing irradiance, coupled with shallower mixed-layer  
413 depths (de Boyer Montégut et al., 2004), result in enhanced growth conditions across the STF  
414 between September and February (Browning et al., 2014). Diagnostic pigment analyses (Fig.  
415 5a) indicated that eukaryotic nanophytoplankton, specifically *Phaeocystis*-type haptophytes,  
416 dominated the early spring STSW chlorophyll-*a* content (73 %) but with a reduced contribution  
417 during summer (20 %). Maximum growth rates for cultured *Phaeocystis antarctica* have been  
418 achieved under elevated Zn concentrations (Saito and Goepfert, 2008), and thus, the dominance  
419 of this haptophyte would likely contribute to the removal of dZn between spring and summer.  
420 Furthermore, an increased summer diatom contribution (13 % chlorophyll-*a* compared with  
421 near zero during spring transects) would have further reduced the dZn inventory, with diatoms  
422 having at least 4-fold higher cellular Zn/P ratios than co-occurring cell types (Twining and  
423 Baines, 2013).

424 The fact that both *Phaeocystis* and diatomaceous nanophytoplankton maintain a contribution  
425 to the summer STSW chlorophyll-*a* complement, when dZn availability is low, is intriguing.

426 Both *P. antarctica* and the large, coastal diatoms *Thalassiosira pseudonana* and *Thalassiosira*  
427 *weissflogii* have been shown to be growth limited in culture by free  $Zn^{2+}$  concentrations  $\leq 10$   
428 pM (Sunda and Huntsman, 1992; Saito and Goepfert, 2008). A simple estimate of summer  
429 STSW free  $Zn^{2+}$  availability, based on North Atlantic organic complexation data ( $>96\%$ ;  
430 Ellwood and Van den Berg, 2000), indicated free  $Zn^{2+}$  averaged  $6.3 \pm 5.3 \mu c$  pM, suggesting the  
431 potential for growth limitation of these phytoplankton. In addition, when comparing the  
432 Southeast Atlantic dZn stoichiometry with the cellular requirements of phytoplankton grown  
433 under growth rate limiting conditions (Fig. 6), we found summer STSW dZn/ $PO_4^{3-}$  to be in  
434 deficit of the requirements of coastal *T. pseudonana* but not those of the smaller, open-ocean  
435 diatom *T. oceanica*. The variation in cellular Zn/P between small and large phytoplankton is  
436 related to the higher surface-area-to-volume ratio of smaller cells, and the limitation of  
437 diffusive uptake rates at low  $Zn^{2+}$  concentrations (Sunda and Huntsman, 1995). This would  
438 suggest that the lower dZn availability in summer STSW should influence phytoplankton  
439 species composition by selecting for smaller organisms with lower cellular Zn requirements,  
440 and confirmed by a ratio of picophytoplankton to nanophytoplankton at least 4-fold higher  
441 during summer compared with spring values. The comparison further implies that the presence  
442 of *Phaeocystis* and diatoms in summer STSW may be linked with their metabolic Zn-Co-Cd  
443 substitution capability, potentially allowing them to overcome some portion of their Zn  
444 deficiency. Largely connected to carbonic anhydrase enzymes, several species of eukaryotic  
445 phytoplankton are capable of biochemical substitution of Zn, Co or Cd to maintain optimal  
446 growth rates under low trace metal conditions (Price and Morel, 1990; Sunda and Huntsman,  
447 1995; Lee and Morel, 1995; Lane and Morel, 2000; Xu et al., 2007; Saito and Goepfert, 2008;  
448 Kellogg et al., 2020). For example, metabolic substitution of Co in place of Zn has been  
449 observed to support the growth of *P. antarctica*, *T. pseudonana* and *T. weissflogii* in media  
450 with  $Zn^{2+} < 3$  pM (Sunda and Huntsman, 1995; Saito and Goepfert, 2008; Kellogg et al., 2020).

451 Thus, the lower mixed-layer dCo inventory of summer STSW, relative to early spring, may be  
452 in part related to enhanced dCo uptake through biochemical substitution alongside the growth  
453 of phytoplankton with distinct Co requirements.

454 In contrast to *Phaeocystis*, *E. huxleyi*-type haptophytes were near-absent in spring STSW (<5  
455 % chlorophyll-*a*; Fig. 5a) and increased in contribution during summer (18 %). *Emiliania*  
456 *huxleyi* appear to have a biochemical preference for Co over Zn (Xu et al., 2007), which could  
457 potentially be a contributing factor to the increased fraction of this haptophyte in summer  
458 STSW. Based on Co organic complexation data for Southeast Atlantic STSW (>99 %; Bown  
459 et al., 2012), however, even the maximum dCo concentration of 56 pM (estimated free  $\text{Co}^{2+}$   
460  $0.56 \pm 0.11 \mu\text{c pM}$ ) observed for STSW during this entire study would limit the growth of  
461 cultured *E. huxleyi* in the absence of Zn or Cd (Sunda and Huntsman, 1995; Xu et al., 2007).  
462 This is supported by inter-seasonal dCo/ $\text{PO}_4^{3-}$  stoichiometries in deficit of the cellular  
463 requirements of cultured *E. huxleyi* (Fig. 6). Despite this, Xu et al. (2007) showed that *E.*  
464 *huxleyi* can maintain significant growth at only 0.3 pM  $\text{Co}^{2+}$  in the presence of Zn, with the  
465 limitation by, and substitution of these metals reported to occur over a range of free ion  
466 concentrations (0.2–5 pM) that is relevant to summer conditions of the Southeast Atlantic. This  
467 assessment implies an additional need for Zn in phytoplankton nutrition due to low dCo  
468 availability throughout the Southeast Atlantic, which may accelerate the decrease in dZn/dCo  
469 inventory ratio between seasons.

470 The elevated summer STSW chlorophyll-*a* concentrations were accompanied by increased cell  
471 concentrations of the *Synechococcus* and *Prochlorococcus* (up to 100 and 400 cells  $\mu\text{L}^{-1}$ ,  
472 respectively) relative to early spring abundance (Fig. 5b). This pattern suggests an inter-  
473 seasonal community shift towards smaller picocyanobacterial cells that is coincident with  
474 decreased dZn availability. *Synechococcus* and *Prochlorococcus* are thought to have little or  
475 no Zn requirement and relatively low Co requirements (growth limited by  $\leq 0.2 \text{ pM } \text{Co}^{2+}$ ; Sunda

476 and Huntsman, 1995; Saito et al., 2002). This, alongside their small cell size, hence greater  
477 capacity for acquiring fixed nitrogen under conditions where this nutrient is depleted, may  
478 allow these prokaryotes to flourish following depletion and export of Zn associated with  
479 *Phaeocystis* and diatom blooms. This supposition is supported by a persistently high abundance  
480 of *Synechococcus* and *Prochlorococcus* ( $>1000$  cells  $\mu\text{L}^{-1}$ ), relative to eukaryotic  
481 nanophytoplankton, in the dZn depleted surface waters of the Costa Rica Dome (Saito et al.,  
482 2005; Ahlgren et al., 2014; Chappell et al., 2016). Here, surface dCo concentrations were  
483 maintained above that of surrounding waters by the biological production of Co-binding  
484 ligands (Saito et al., 2005). The increased abundance of these prokaryotic autotrophs in summer  
485 STSW of the Southeast Atlantic may have also contributed to the inter-seasonal decrease in  
486 dCo inventory.

487 In contrast to STSW, cells counts of eukaryotic phytoplankton and prokaryotic cyanobacteria  
488 in SASW varied little between early spring and summer (Fig. 5b), indicative of a more balanced  
489 ecophysiological regime. The fractional contribution of *Phaeocystis* (Fig. 5a), the dominant  
490 contributor to the SASW chlorophyll-*a* complement, was similar between transects at 54 and  
491 44 %, respectively, whilst the contribution of *E. huxleyi* increased from 19 to 33 % between  
492 spring and summer, respectively. Whilst it is proposed that the low Fe supply rate to these  
493 waters provides a dominant control on phytoplankton biomass and composition (Browning et  
494 al., 2014), low dZn and dCo availability may also be important drivers of such change. The  
495 Summer SASW dZn inventory ( $0.08 \pm 0.07u_c$  nM) and stoichiometry with  $\text{PO}_4^{3-}$  (Fig. 6)  
496 indicate growth limiting conditions for *Phaeocystis* and *E. huxleyi* in the absence of  
497 cambialistic metabolism (Sunda and Huntsman., 1995; Saito and Goepfert, 2008; Xu et al.,  
498 2007). The presence of these phytoplankton therefore indicates Zn biochemical substitution  
499 occur in oceanic waters of the Southeast Atlantic. A lower Co half-saturation growth constant  
500 for cultured *P. antarctica* ( $K_m = \sim 0.2$  pM  $\text{Co}^{2+}$ ), compared with *E. huxleyi* ( $K_m = \sim 3.6$  pM

501  $\text{Co}^{2+}$ ), further suggests that *Phaeocystis* species may more effectively occupy low dZn and dCo  
502 environments (Saito and Goepfert, 2008), such as SASW of the South Atlantic.  
503 Conversely, the absence of a significant diatom contribution to summer SASW chlorophyll-*a*  
504 (Fig. 5a), relative to early spring, is surprising as the summer dZn/ $\text{PO}_4^{3-}$  inventory ratio is in  
505 excess of the cellular Zn/P requirements of typical oceanic diatoms such as *T. oceanica* (Fig.  
506 6). Furthermore, whilst the dCo/ $\text{PO}_4^{3-}$  ratio of summer SASW is in deficit of the cellular Co/P  
507 below which growth limitation of *T. oceanica* may occur, this species has been shown to grow  
508 effectively at  $\text{Co}^{2+} < 0.1 \text{ pM}$  in culture in the presence of Zn (Sunda and Huntsman, 1995). The  
509 low diatom fractional contribution to summer SASW may be instead related to low Fe  
510 availability (Browning et al., 2014) and stress-induced Si exhaustion. In support of this, we  
511 calculate summer SASW mixed-layer Si concentrations ( $0.9 \pm 0.3 \text{ }\mu\text{M}$ ) to be 50 % of early  
512 spring values ( $1.8 \pm 0.2 \text{ }\mu\text{M}$ ) with a dissolved  $\text{NO}_3^-/\text{Si}$  stoichiometry of  $3.8 \text{ mol mol}^{-1}$  close to  
513 the  $4 \text{ mol mol}^{-1}$  shown to limit diatom growth in culture (Gilpin et al., 2004), and in contrast to  
514 the  $2.9 \text{ mol mol}^{-1}$  calculated for early spring.

515

### 516 **3.6. Conclusion**

517 We report the distributions of dZn and dCo in the upper water column of sub-tropical and sub-  
518 Antarctic waters of the Southeast Atlantic during austral spring and summer periods. We  
519 identify an apparent continental source of dZn and dCo to sub-tropical waters at depths between  
520 20 – 55 m, derived from sedimentary inputs from the Agulhas Bank. In contrast, open-ocean  
521 sub-Antarctic surface waters displayed largely consistent inter-seasonal mixed-layer dZn and  
522 dCo concentrations indicating a more balanced ecophysiological regime with regard to their  
523 organisation. The vertical distributions of dZn and dCo in the upper water column were similar  
524 to that of  $\text{PO}_4^{3-}$  indicating biological drawdown in surface waters and mixing with underlying  
525 Southern ocean-derived waters travelling equatorward significantly influences their

526 distribution. Absolute trace metal concentrations alongside concentration inventory ratios  
527 suggest the preferential utilization of dZn, relative to dCo, in the Southeast Atlantic with  
528 dZn/dCo decreasing from 19 to 5 mol mol<sup>-1</sup> between early spring and summer in STSW and  
529 from 13 to 7 mol mol<sup>-1</sup> in SASW. This pattern is consistent with our understanding of the  
530 cellular requirement of phytoplankton (Twining and Baines, 2013). The inter-seasonal removal  
531 of dZn results in summer concentrations that are potentially growth limiting for certain  
532 phytoplankton species estimated to be present in these waters by diagnostic pigment analyses.  
533 We therefore suggest cambialistic metabolic substitution between Zn and Co, and potentially  
534 Cd, is an important factor regulating the growth, distribution and diversity of phytoplankton in  
535 the Southeast Atlantic.

536

537 *Data availability.* The trace metal and macronutrient data sets used for analyses in this study  
538 are available at <https://www.bodc.ac.uk/geotraces/data/idp2017/> (GEOTRACES GA10) and  
539 phytoplankton data at <https://www.bodc.ac.uk/>.

540

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

542

543 *Author contribution.* MCL and EPA acquired the funding. NJW, MCL, AM, TJB, EMSW, and  
544 HAB collected samples at sea. NJW conducted the Zn and Co measurements, EMSW the  
545 macronutrient measurements and TJB the phytoplankton measurements. NJW prepared the  
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547

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552

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834

835 Table 1. Analytical validation results for open ocean surface seawater (SAFe S), 1000 m  
836 seawater (SAFe D2) and 2000 m seawater (GEOTRACES GD). All concentrations are in nM  
837 ( $\pm 1$  std. dev.). Consensus value conversion = 1.025 kg/L. ND indicates sample not determined.

838

	SAFe S	SAFe D2	GEOTRACES GD
Zn (FIA)	0.060 (0.020) $n = 7$	7.723 (0.091) $n = 12$	ND
Zn consensus value	0.071 (0.010)	7.616 (0.256)	1.753 (0.123)
Co (FIA)	0.004 (0.001) $n = 3$	0.049 (0.001) $n = 2$	0.073 (0.004) $n = 5$
Co consensus value	0.005 (0.001)	0.047 (0.003)	0.067 (0.001)

839

840

841 Table 2. Southeast Atlantic dissolved micro- and macronutrient mean concentration inventories  
842 for the upper water column during early spring (D357-1), late spring (D357-2) and summer  
843 (JC068) transects. STSW and SASW waters were defined using the  $\theta$  15°C isotherm (Section  
844 3.4) and are compared with total inventories calculated for the shallower mixed layer (in  
845 parenthesis) that include continental inputs of dissolved Zn and Co. Zn/PO<sub>4</sub><sup>3-</sup>, Co/PO<sub>4</sub><sup>3-</sup> and  
846 Zn/Co represent the concentration inventory ratios for STSW and SASW, respectively. STSW  
847 = Sub-Tropical Surface Water, SASW = Sub-Antarctic Surface Water.

848

Oceanographic Regime	Transect	Zn (nmol m <sup>-3</sup> )	Co (nmol m <sup>-3</sup> )	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup> (μmol m <sup>-3</sup> )	Si(OH) <sub>4</sub>	Zn/PO <sub>4</sub> <sup>3-</sup> (μmol mol <sup>-1</sup> )	Co/PO <sub>4</sub> <sup>3-</sup> (μmol mol <sup>-1</sup> )	Zn/Co (mol mol <sup>-1</sup> )
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STSW	Early spring	624 (1597)	32 (30)	2694 (870)	333 (203)	3735 (2790)	1876	97	19
	Late spring	384 (592)	23 (17)	1846 (763)	276 (191)	2781 (2326)	1387	82	17
	Summer	158 (139)	29 (24)	1557 (326)	226 (139)	2711 (1942)	699	129	5
SASW	Early spring	182 (112)	14 (13)	6035 (5300)	615 (566)	1875 (1847)	296	22	13
	Summer	83 (94)	12 (10)	4143 (3388)	439 (400)	1027 (886)	188	26	7

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850

851

852 Figure 1. The Southeast Atlantic stations sampled for dissolved Zn and Co along the GA10  
853 section during UK-GEOTRACES cruises D357 (red circles) and JC068 (black circles),  
854 overlain a VIIRS monthly composite image of chlorophyll-*a* concentrations for January 2012  
855 (<https://oceancolor.gsfc.nasa.gov/>). Two transects were completed during D357 between Cape  
856 Town and the zero meridian that represent early austral spring 2010 (D357-1; Stns. 1, 2, 3, 4,  
857 5 & 6) and late austral spring 2010 (D357-2; Stns. 0.5, 1, 1.5, 2.5, 3.5, 4.5), respectively. JC068  
858 took place during austral summer 2011/12 and we present here only the repeat transect data  
859 between Cape Town and 13°W (Stns. 1, 2, 3, 7, 8, 9, 11). STSW = Sub-Tropical Surface Water,  
860 SASW = Sub-Antarctic Surface Water, AC = Agulhas Current, AR = Agulhas retroflexion.

861

862 Figure 2. Upper 500 m potential temperature ( $\theta$ ) and dissolved  $\text{PO}_4^{3-}$  distributions for the  
863 Southeast Atlantic along early spring (a,b; D357-1), late spring (c,d; D357-2) and summer (e,f;  
864 JC068) transects. The dominant Southern Ocean (SASW & SAMW) and South Atlantic  
865 (STSW) water masses that influence the distribution of nutrients are shown. The  $\theta$  15°C  
866 isotherm (solid contour) represents a practical definition of the STF location, whilst SAMW is  
867 identified by the median potential density ( $\sigma_\theta$ ) isopycnal 26.8 kg m<sup>-3</sup> (dashed contour, see Sect.  
868 4.1.). STSW = Sub-Tropical Surface Water, SAMW = Sub-Antarctic Mode Water, AAIW =  
869 Antarctic Intermediate Water.

870

871 Figure 3. Upper 500 m dissolved Zn and Co distributions for the Southeast Atlantic along early  
872 spring (a,b; D357-1), late spring (c,d; D357-2) and summer (e,f; JC068) transects. The STF is  
873 delineated by  $\theta$  15°C (solid contour), whilst the influence of SAMW is evident by the median  
874 potential density ( $\sigma_\theta$ ) isopycnal 26.8 kg m<sup>-3</sup> (dashed contour, see Section 4.1.). STSW = Sub-  
875 Tropical Surface Water, SAMW = Sub-Antarctic Mode Water, AAIW = Antarctic Intermediate  
876 Water. Note the changing y-axis scales for dZn distribution.

877

878 Figure 4. The dissolved Zn and Co versus  $\text{PO}_4^{3-}$  distribution for the Southeast Atlantic during  
879 early spring (a,b; D357-1), late spring (c,d; D357-2) and summer (e,f; JCO68) transects. The

880 green and red lines indicate the  $dZn:PO_4^{3-}$  regression slopes for SAMW and AAIW,  
881 respectively. The yellow line indicates the  $dCo:PO_4^{3-}$  regression slope for SAMW and AAIW  
882 combined. The equations for regression lines are detailed in Supplementary table 1. SAMW =  
883 Sub-Antarctic Mode Water, AAIW = Antarctic Intermediate Water. The full depth  $dZn:PO_4^{3-}$   
884 relationship along JC068 can be found in Wyatt et al. (2014).

885

886 Figure 5. Seasonal differences in (a) pigment-derived taxonomic contributions to total  
887 chlorophyll-*a* (percentage), and (b) AFC counts of *Synechococcus*, *Prochlorococcus*,  
888 nanophytoplankton (approx.  $>2\mu m$ ) and photosynthetic picoeukaryotes (approx.  $<2\mu m$ ) in the  
889 Southeast Atlantic.

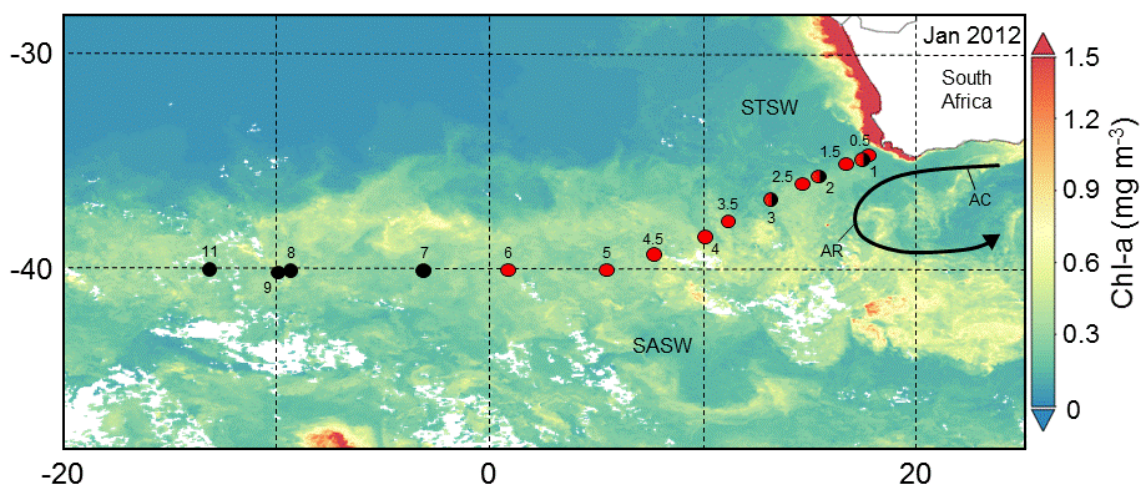
890

891 Figure 6. Metal/ $PO_4^{3-}$  inventory ratios for the upper water column of the Southeast Atlantic  
892 (horizontal bars) compared with laboratory estimates of cellular ratios in eukaryotic  
893 phytoplankton below which growth limitation occurs (solid vertical lines represent Zn:P with  
894 no added Co to media whilst dashed lines represent Co:P with no added Zn; phytoplankton  
895 data from Sunda and Hunstman, 1995). Error bars on inventory ratios represent 20 % combined  
896 uncertainty for  $dZn$  and  $dCo$  analyses (see Section 2.2). This figure is adapted from that in Saito  
897 et al. (2010) and implies that inter-seasonal differences in metal/ $PO_4^{3-}$  stoichiometry could  
898 impact phytoplankton community composition in the Southeast Atlantic.

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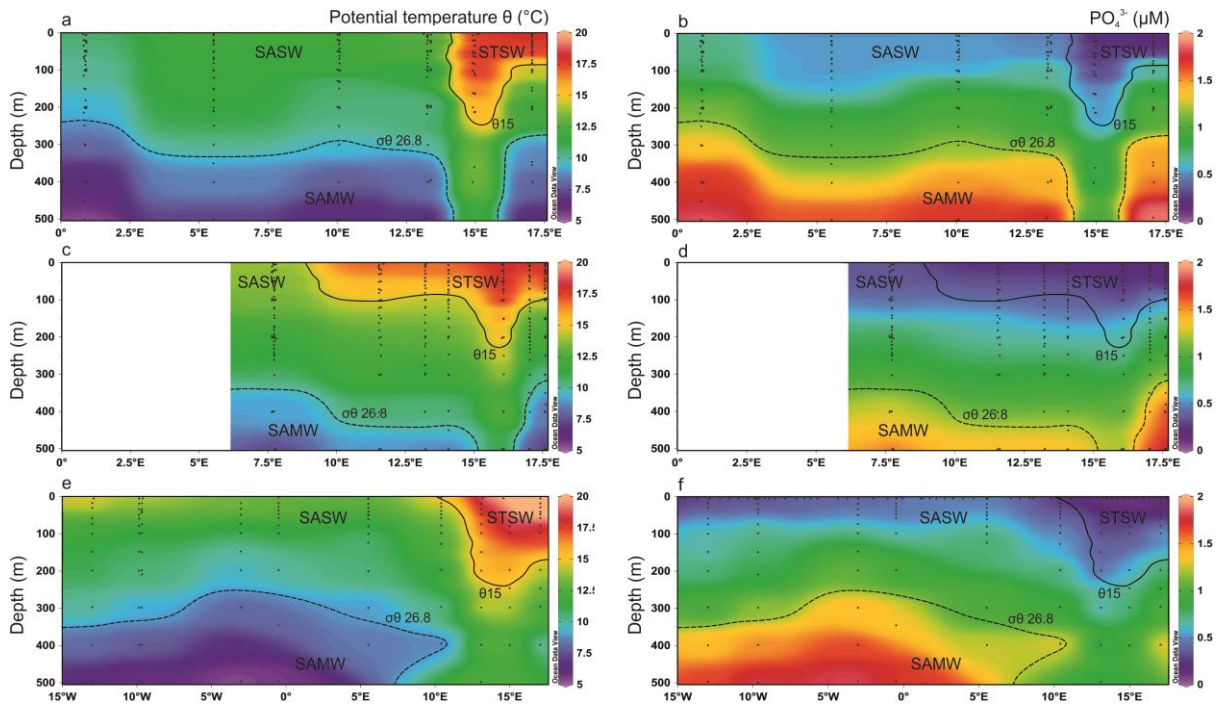


902

903 Figure 1.

904



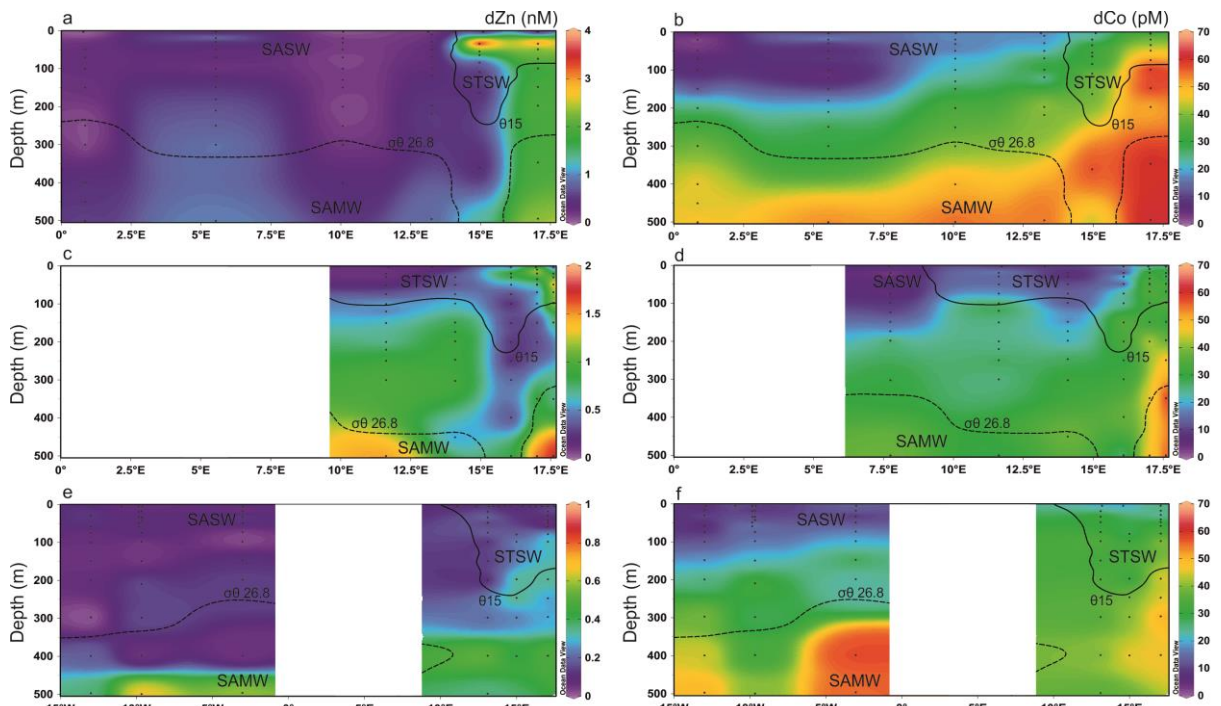


905

906 Figure 2.

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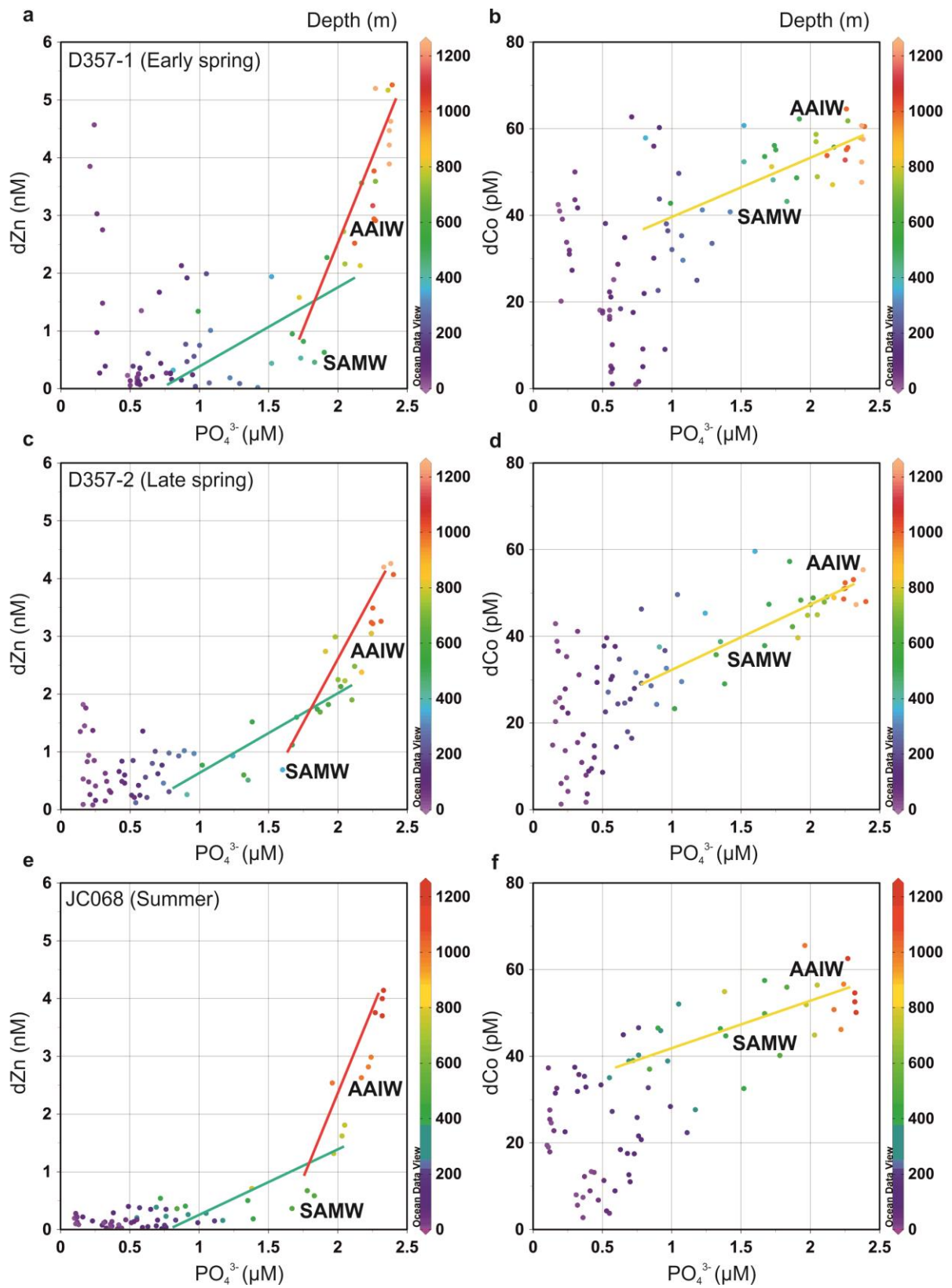
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910 Figure 3.

911

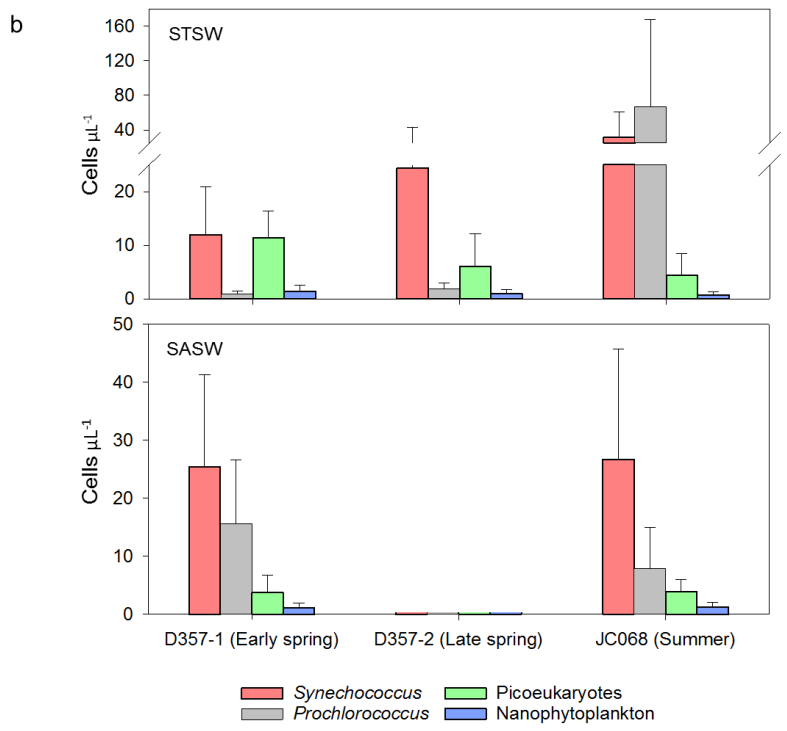
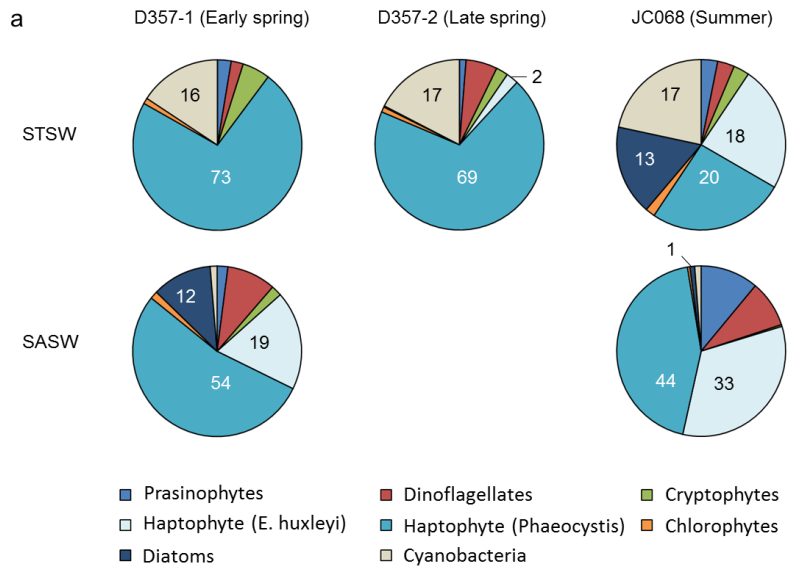


912

913 Figure 4.

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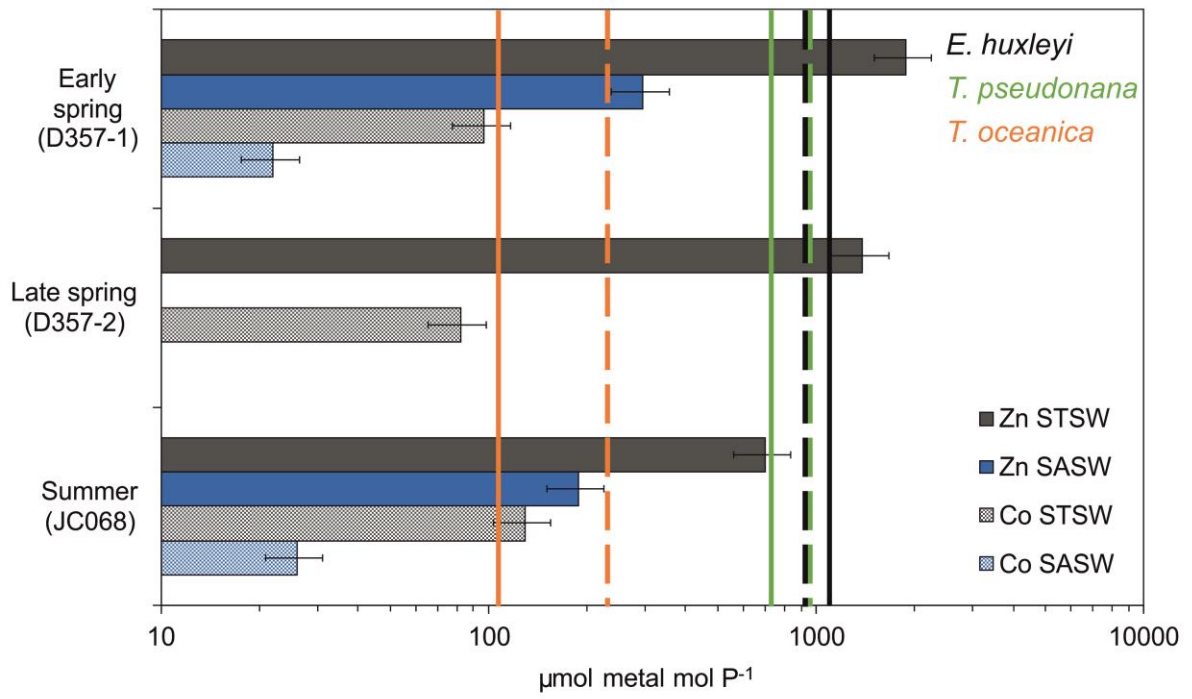
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916

917 Figure 5.

918



919  
 920 Figure 6.  
 921