1 Coccolithophore biodiversity controls carbonate export in the Southern Ocean

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21 Abstract

22 Southern Ocean waters are projected to undergo profound changes in their physical and chemical properties in the coming decades. Coccolithophore blooms in the 23 24 Southern Ocean are thought to account for a major fraction of the global marine calcium carbonate (CaCO₃) production and export to the deep sea. Therefore, changes in the 25 composition and abundance of Southern Ocean coccolithophore populations are likely to 26 27 alter the marine carbon cycle, with feedbacks to the rate of global climate change. However, the contribution of coccolithophores to CaCO₃ export in the Southern Ocean is 28 29 uncertain, particularly in the circumpolar Subantarctic Zone that represents about half of 30 the areal extent of the Southern Ocean and where coccolithophores are most abundant. 31 Here, we present measurements of annual CaCO₃ flux and quantitatively partition them 32 amongst coccolithophore species and heterotrophic calcifiers at two sites representative

of a large portion of the Subantarctic Zone. We find that coccolithophores account for a 33 34 major fraction of the annual CaCO₃ export with highest contributions in waters with low algal biomass accumulations. Notably, our analysis reveals that although Emiliania 35 huxleyi is an important vector for CaCO₃ export to the deep sea, less abundant but larger 36 species account for most of the annual coccolithophore CaCO₃ flux. This observation 37 contrasts with the generally accepted notion that high PIC accumulations during the 38 austral summer in the subantarctic Southern Ocean are mainly caused by E. huxleyi 39 40 blooms. It appears likely that the climate-induced migration of oceanic fronts will initially result in the poleward expansion of large coccolithophore species increasing CaCO₃ 41 production. However, subantarctic coccolithophore populations will eventually diminish 42 as acidification overwhelms those changes. Overall, our analysis emphasizes the need for 43 species-centred studies to improve our ability to project future changes in phytoplankton 44 45 communities and their influence on marine biogeochemical cycles.

46

47 <u>1. Introduction</u>

The emissions of carbon dioxide (CO₂) into the atmosphere by anthropogenic 48 industrial activities over the past 200 years are inducing a wide range of alterations in the 49 marine environment (Pachauri et al., 2014). These include ocean warming, shallowing 50 51 of mixed layer depths, changes in nutrient supply to the photic zone, and decreasing carbonate-ion concentrations and pH of the surface ocean, a process known as ocean 52 acidification (Rost and Riebesell, 2004; Stocker et al., 2014). Substantial evidence from 53 CO₂ manipulation experiments indicates that many species of corals, pteropods, 54 planktonic foraminifera and coccolithophores will reduce their calcification rates under 55 56 future ocean acidification scenarios (Bijma et al., 2002; Langdon and Atkinson, 2005 among others; Orr et al., 2005; Bach et al., 2015; Meyer and Riebesell, 2015). Owing to 57 58 their moderate alkalinity and cold temperatures, Southern Ocean waters are projected to become undersaturated with respect to aragonite no later than 2040 and to calcite by the 59 end of the century (Cao and Caldeira, 2008; McNeil and Matear, 2008; Shadwick et al., 60 2013). This decline in the saturation state of carbonate, together with other changes in 61 carbonate chemistry speciation, will enhance dissolution of both aragonite and calcite 62 shells and will make the biological precipitation of carbonate difficult in some marine 63 64 calcifying organisms (Fabry et al., 2008; Gattuso and Hansson, 2011). Since such 65 thresholds will be reached sooner in polar regions, Southern Ocean ecosystems have been

proposed as bellwethers for prospective impacts of ocean acidification on marineorganisms at mid and low latitudes (Fabry et al., 2009).

68 Coccolithophores are a major component of phytoplankton communities in the 69 Southern Ocean, particularly in its northern-most province, the Subantarctic Zone, where 70 they often exhibit maximum abundances and diversity (e.g. Gravalosa et al., 2008; 71 Saavedra-Pellitero et al., 2014; Malinverno et al., 2015; Charalampopoulou et al., 2016). 72 Coccolithophores play an important and complex role in the Southern Ocean carbon cycle (Salter et al., 2014). On the one hand, the production of calcite platelets (termed 73 74 coccoliths) decreases the alkalinity of surface waters thereby reducing the atmospheric uptake of CO_2 from the atmosphere into the surface ocean. On the other hand, the 75 76 production of organic matter through photosynthesis, and its subsequent transport to 77 depth in settling particles, enhances carbon sequestration via the biological carbon pump (Volk and Hoffert, 1985). Additionally, due to their high density and slow dissolution, 78 coccoliths act as an effective ballast for organic matter, increasing organic carbon 79 80 sequestration depths (Buitenhuis et al., 2001; Boyd and Trull, 2007; Ziveri et al., 2007). Therefore, changes in the abundance, composition and distribution of coccolithophores 81 could have an extensive impact on ocean nutrient stoichiometry, carbon sequestration, 82 83 and nutrition for higher trophic levels in the Southern Ocean (Deppeler and Davidson, 2017). 84

85 The remoteness and vastness of the Southern Ocean, together with the inherent 86 temporal and spatial variability of pelagic ecosystems, hampers accurate characterization and quantification of Southern Ocean phytoplankton communities. Advances in satellite 87 88 technology and modelling algorithms have allowed a circumpolar and year-round coverage of the seasonal evolution of major phytoplankton functional groups within the 89 Southern Ocean (e.g. Alvain et al., 2013; Hopkins et al., 2015; Rousseaux and Gregg, 90 2015). In particular, ocean-colour satellite reflectance observations have been used to 91 92 quantitatively estimate coccolithophore Particulate Inorganic Carbon (PIC) concentrations throughout the Southern Ocean (Gordon et al., 2001; Balch et al., 2005b). 93 94 These satellite estimates suggest apparent high PIC values during summer near the major 95 Southern Ocean fronts attributed to coccolithophores (Balch et al., 2011; Balch et al., 96 2016). This band of elevated reflectance and PIC that encircles the entire Southern Ocean was termed the "Great Calcite Belt" by these authors. However, comparison of satellite 97 98 remote-sensing data with ship-based observations (Holligan et al., 2010; Trull et al.,

2018) indicate that satellite ocean-colour-based PIC estimates could be unreliable, 99 100 particularly in Antarctic waters where they erroneously suggests high PIC abundances. 101 Shipboard observations, on the other hand, provide a detailed picture of phytoplankton community composition and structure, but are dispersed, both temporally and 102 geographically, and provide rather heterogenous data in terms of taxonomic groups 103 104 investigated, and the sampling scales and methodologies used (e.g. Kopczynska et al., 2001; de Salas et al., 2011; Poulton et al., 2013; Patil et al., 2017, among others). In situ 105 106 year-round monitoring of key strategic regions is critically needed to establish baselines 107 of phytoplankton community composition and abundance and to validate and improve 108 ocean biogeochemical models (Rintoul et al., 2012). This information is also essential if 109 we are to detect possible climate-driven changes in the structure of phytoplankton 110 communities that could influence the efficiency of the biological carbon pump, with 111 consequent feedbacks to the rate of deep-water carbon sequestration and global climate change (Le Quéré et al., 2007; Deppeler and Davidson, 2017). 112

113 Here, we document coccolithophore and carbonate particle fluxes collected over a year by four sediment trap records deployed at two strategic locations of the Australia 114 and New Zealand sectors of the Southern Ocean considered representative of a large 115 portion of the SAZ (See section 2.2 for further details). Our measurements provide 116 117 coccolith mass estimates of the main coccolithophore species and quantitatively partition annual carbonate fluxes amongst coccolithophore species and heterotrophic calcifiers. 118 119 We find that coccolithophores are a major vector for CaCO₃ export out of the mixed layer and that the largest contribution to CaCO₃ export is not from the most abundant species 120 121 Emiliania huxleyi but rather from larger coccolithophores species with substantially 122 different physiological traits (e.g. Calcidiscus leptoporus). Our results emphasize the urgent need for diagnostic fitness response experiments on other coccolithophore species 123 124 aside from E. huxleyi (e.g. Feng et al., 2017) in order to be able to be able to predict the 125 impacts of anthropogenically induced changes in Southern Ocean ecosystems and 126 biological carbon uptake mechanisms.

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- 128 **2. Material and methods**
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130 2.1 Oceanographic setting

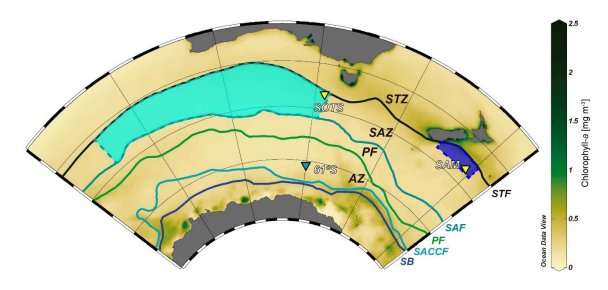
The SAZ alone accounts for more than half of the Southern Ocean area (Orsi et 131 132 al., 1995) and represents a transitional boundary between the warm, oligotrophic waters of the subtropical gyres to the north and the cold, silicate-rich waters south of the Polar 133 Front (PF). The SAZ is arguably the largest high nutrient, low chlorophyll (HNLC) 134 province in the world's ocean and is central to the linkages between the ocean-135 atmosphere CO₂ exchange and climate. The deep winter convection in the SAZ, which 136 exceeds 400 m, results in the formation of a high-oxygen water masses known as 137 Subantarctic Mode and Antarctic Intermediate Waters that connect the upper and lower 138 139 limbs of the global overturning circulation (Sloyan and Rintoul, 2001a, b). The formation of these water masses are responsible for the sequestration of a large fraction of 140 anthropogenic CO₂ (Sabine et al., 2004), with an estimated 1 Gt C yr⁻¹ transported to 141 intermediate depths annually (Metzl et al., 1999). Macronutrient concentrations display 142 143 pronounced seasonal changes in the SAZ with fully replete levels during winter to substantial depletion during summer, particularly for silicate (Dugdale et al., 1995; 144 145 Rintoul and Trull, 2001; Bowie et al., 2011). Phytoplankton community in the 146 subantarctic zone is dominated by pico- and nanoplankton including cyanobacteria, 147 coccolithophores and autotrophic flagellates with lower abundances of diatoms than polar waters south the Polar Front (Chang and Gall, 1998; Kopczynska et al., 2001; de Salas et 148 al., 2011; Rigual-Hernández et al., 2015b; Eriksen et al., 2018). 149

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151 **2.2 Field experiments**

152 Here we report on the coccolithophore and biogeochemical fluxes collected over a year at the Australian Southern Ocean Time Series (SOTS) observatory (Trull et al., 153 154 2010) and the New Zealand Subantarctic Mooring (SAM) site (Nodder et al., 2016) (Fig. 1). The SOTS observatory is located in the abyssal plane of the central SAZ 155 156 approximately 530 km southwest of Tasmania (46° 56' S, 142° 15' E) within an anti-157 cyclonic gyre in a region characterized by weak circulation (Trull et al., 2001; Herraiz-Borreguero and Rintoul, 2011). SOTS was equipped with three vertically moored, conical 158 159 time-series sediment traps (McLane Parflux Mk 7G-21) placed at ~1000, 2000 and 3800 160 m depth between August 2011 until July 2012. The physical, chemical and biological 161 parameters of SOTS site are regarded as representative for large portion of the Indian and Australian sectors of the SAZ (~90°E and 140°E; Trull et al., 2001). The SAM site is 162 163 located in the Bounty Trough in in the subantarctic waters south east of New Zealand

(46°40'S, 178' 30°E) and was equipped with conical, time-incremental sediment trap 164 165 (McLane PARFLUX Mk7G-21) placed at 1500 m depth, with samples used in the present study collected between November 2009 until November 2010. The SAM site is 166 167 considered to be representative of a wide area of the northern sector of the SAZ off eastern New Zealand, approximately 171°E to 179°W and 45 to 47°S (Law et al., 2014; Fig. 1). 168 Full details of the field experiments from these two localities in the Australian and New 169 Zealand sectors of the SAZ can be found in Trull et al. (2001) and Nodder et al. (2016), 170 171 respectively.



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Figure 1: Chlorophyll-a composite map of the Australian-New Zealand sector of the 173 Southern Ocean (July 2002 to July 2012) from the MODIS Aqua Sensor showing the 174 location of the sediment trap moorings sites: SOTS, 61°S and SAM. The regions for 175 176 which the SOTS and SAM sites are representative are marked with light and dark blue 177 areas, respectively. Abbreviations: Subtropical Zone - STZ, Subtropical front - STF, Subantarctic Zone - SAZ, Subantarctic Front - SAF, Polar Frontal Zone - PFZ, Polar 178 Front - PF, Antarctic Zone - AZ, Southern Antarctic Circumpolar Current Front -179 SACCF, southern boundary of the ACC – SB. Oceanic fronts after Orsi et al. (1995). 180

181 **2.3 Sample processing**

In short, the recovered trap bottles were refrigerated upon recovery and then allowed to settle. The sample slurry was then wet-sieved through a 1 mm screen in the case of SOTS (no attempt to extract zooplankton "swimmers" was made for the <1 mm fraction analysed here) and through a 200 μ m sieve to remove "swimmers" for the SAM site. The remaining fraction was then split using a McLane wet sample divider; the SOTS

samples were subdivided into one tenth aliquots while one fifth splits were made for the 187 188 SAM samples. For the SOTS samples, a total of 55 samples were processed for calcareous nannoplankton analysis. The one-tenth splits dedicated to phytoplankton analysis were 189 190 further subdivided into four aliquots with the McLane splitter. One aliquot was used for calcareous nannoplankton analysis and the remaining three were kept refrigerated for 191 192 biomarker and non-calcareous microplankton analyses. In the case of the SAM samples, 193 the one-fifth aliquots were further subdivided into five subsplits, and one of those was 194 used for calcareous nannoplankton analysis. Two different types of glass slides per 195 sample were prepared. The first preparation was used for the estimation of coccosphere 196 and calcareous dinocyst (calcispheres of thoracosphaerids) fluxes and for coccolith 197 imaging. A volume ranging between 1000 and 5000 µl of the raw sample was mounted 198 on a glass slide using Canada balsam following Flores and Sierro (1997). This technique 199 produces random settling of the coccoliths for microscopic identification and 200 enumeration. The second type of glass slide was prepared following a modified protocol 201 for non-destructive disintegration of aggregates modified from Bairbakhish et al. (1999). The objective of this chemical treatment is to reduce biases in the coccolith flux 202 203 estimations associated with the presence of different types of aggregates and 204 coccospheres (Bairbakhish et al., 1999). In brief, 2000 µl were extracted from the aliquot 205 for calcareous nannoplankton analysis and then treated with a solution comprising 900 µl 206 sodium carbonate and sodium hydrogen carbonate, 100 µl ammonia (25%) and 2000 µl 207 hydrogen peroxide (25%). The sample was agitated for 10 seconds every 10 minutes and 208 this process was repeated over an hour. Then, the reaction was stopped with catalase 209 enzyme and samples were allowed to settle for at least 48 hours before preparation on 210 microscope slides. pH controls indicate that the solution kept pH levels near 9, therefore 211 precluding coccolith dissolution. Finally, trap samples were mounted on microscope slides following the same decantation method as used for the first type of glass slides (i.e. 212 213 Flores and Sierro, 1997).

214 **2.4 Determination of CaCO₃ fluxes**

A detailed description of the geochemical analytical procedures for the SOTS samples is provided in Trull et al. (2001) and Rigual-Hernández et al. (2015a) while more detailed procedures of the SAM trap can be found in Nodder et al. (2016). In short, for the SOTS site three of the one tenth splits were filtered onto 0.45 pore size filters. Then the material was removed from the filter as a wet cake of material, dried at 60°C, and

ground in an agate mortar. This material was used to determine the total mass and 220 221 composition of the major components of the flux. Particulate inorganic Carbon (PIC) 222 content was measured by closed system acidification with phosphoric acid and 223 coulometry. For the SAM site, one-fifth split was analysed for elemental calcium (Ca) 224 concentration using ICP-MS techniques. The samples were oven-dried, digested in 225 nitric/hydrochloric acid and then analysed according to the methods under US EPA 200.2. Ca was used to estimate CaCO₃ content in the samples assuming a 1:1 molar ratio in 226 227 CaCO₃.

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229 **2.5** Quantification and characterization of coccolithophore sinking assemblages

230 Qualitative and quantitative analyses of coccospheres and coccoliths were 231 performed using a Nikon Eclipse 80i polarised light microscope at 1000 x magnification. 232 The taxonomic concepts of Young et al. (2003) and the Nannotax website (Young et al., 233 2019) were used. A target of 100 coccospheres and 300 coccoliths was established; 234 however, owing to the pronounced seasonality in coccolithophore export, there were 235 some periods with very low abundance of coccospheres in the samples and therefore the 236 target of 100 coccospheres was not always met. Coccosphere and coccolith species counts were then transformed into relative abundances and daily fluxes using the following 237 formula: 238

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- 240

 $F = \frac{N \times \frac{A}{n \times a} \times V \times S}{d \times T}$

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where F = coccolith flux, N = number of coccoliths, A = area of the Petri dish, n= number of fields of view, a = area of a field of view, V = dilution volume, S = sample split, d = number of days of collection and T = sediment trap aperture area.

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246 **2.6 Determination of coccolith mass and size**

Birefringence and morphometric methods are the two most commonly used approaches for estimating the calcite content of isolated coccoliths. The circularlypolarized light-microscopy-based technique (Fuertes et al., 2014) is based on the systematic relationship between the thickness of a given calcite particle (in the thickness range of 0 - 1.55 mm) and the first-order polarization colours that it displays under

polarized light (Beaufort, 2005; Beaufort et al., 2014; Bolton et al., 2016). The advantages 252 253 of this approach are that: (i) it directly measures complete coccoliths with no assumptions 254 regarding their shape or thickness and (ii) it allows for quantification of calcite losses 255 associated with missing parts or etching of the coccoliths. Disadvantages of this technique 256 are the errors associated with the coccolith-calcite calibration and their consequent effect 257 on the coccolith mass estimates (Fuertes et al., 2014; González Lemos et al., 2018). The 258 morphometric approach, on the other hand, allows better taxonomic identification of the 259 coccoliths and has smaller errors in the length measurements (~0.1 to 0.2 μ m; Poulton et 260 al. 2011). However, this method does not allow direct measurement of coccolith thickness 261 and assumes identical shape and width proportions for all specimens of the same species, 262 among other uncertainties (see Young and Ziveri, 2000 for a review). Since the two 263 methods have different associated errors (Poulton et al., 2011), we applied both 264 approaches to our coccolith flux data in order to obtain two independent estimates of the 265 fractional contribution of coccolithophores species to total carbonate export in the SAZ.

266 For the birefringence-based approach, a minimum of 50 coccoliths of each of the 267 main coccolithophore species were imaged using a Nikon Eclipse LV100 POL light 268 microscope equipped with circular polarisation and a digital camera (Nikon DS-Fi1 8-bit colour). The only exception was *E. huxleyi* for which coccolith mass values had already 269 270 been estimated in all the same samples at high resolution by Rigual-Hernández et al. (under review). For the minor components of the flux assemblage, a lower number of 271 272 coccoliths were measured (Table 1). A photograph of the same apical rhabdolith of the 273 genus Acanthoica was taken and used for calibration at the beginning of each imagining 274 session during which microscopy light and camera settings were kept constant. A 275 different number of fields of view of multiple samples representative of different seasons were photographed until the target number of coccoliths for each species was reached. 276 277 Photographs were then analysed by the image processing software C-Calcita. The output 278 files for single coccoliths were visually selected and classified into the lowest possible 279 taxonomic level. Length and weight measurements were automatically determined by C-280 Calcita software. Morphometric measurements of all the species are summarized in Table 281 1. For further methodological details see Fuertes et al. (2014) and Bolton et al. (2016).

The second approach consisted of performing morphometric measurements on the coccoliths followed by the estimation of their coccolith mass assuming a systematic relation between length and thickness (Young and Ziveri, 2000). Young and Ziveri (2000) proposed that the calcite content of a given coccolith could be estimated using thefollowing formula:

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Coccolith calcite (pg) = $2.7 \text{ x } \text{k}_{\text{s}} \text{ x } \text{l}^{3}$

where 2.7 is the density of calcite (CaCO₃; pg μ m³), "k_s" is a shape constant that varies between species and morphotypes and whose value is based on the reconstruction of coccolith cross profiles and "l" is the distal shield length (DSL). In order to undertake coccolith measurements on the same coccoliths used for the birefringence-based approach, we employed the distal shield length values measured by C-Calcita using circularly polarized light instead of morphometric measurements on Scanning Electron Micrographs (SEM) as made in Young and Ziveri (2000).

295 Since coccolith distal shield length (DSL) has been reported to be systematically 296 underestimated using cross-polarized light microscopy (e.g. D'Amario et al., 2018), we 297 evaluated the possible errors in the DSL measurements made by C-Calcita. For this 298 assessment, we measured 40 detached coccoliths of C. leptoporus under the SEM from 299 samples of the SOTS sediment traps using the image processing software Image-J. 300 Average DSL measurements under the SEM were then compared with those made by C-301 Calcita on 40 randomly selected C. leptoporus coccoliths. The average coccolith length 302 obtained with the SEM analysis $(6.37 \pm 1.02, n = 40)$ was ~ 4% shorter than that estimated 303 with C-Calcita (6.62 \pm 1.47, n = 40). Therefore, we assumed the error for the DSL 304 measurements with circularly polarized light is < 5%. Given the low numbers of the rest 305 of species in the samples we considered that this error is applicable for the rest of the taxa 306 measured in the current study. The subtle differences in coccolith distal length 307 measurements between techniques are most likely due to the fact that the peripheral limit 308 of the coccolith shield under the circulary-polarized light microscope (LM) is not as sharp as is the case for SEM images. It follows that differences in DSL measurements between 309 310 SEM and LM techniques will be likely similar or smaller in the case of larger species. 311 Since the majority of coccolith species identified in the current study display a similar 312 (e.g. Gephyrocapsa oceanica, Syracosphaera pulchra, Umbellosphaera tenuis and 313 Umbilicosphaera sibogae) or larger size (e.g. Coccolithus pelagicus and Helicosphaera 314 *carteri*) than *C. leptoporus*, it could be assumed that the <5% error on DSL estimates for 315 C. leptoporus is applicable for the rest of the species found in the current study. For the k_s value of each taxa, data from the literature was employed (Table 1). E. huxleyi 316 assemblages in the SAZ are composed of a mixture of five different morphotypes: A, A 317

- overcalcified, B, B/C and C, each of which is characterized by different shape factors (k_s) .
- Since k_s is not available for all the morphotypes found in the SAZ and it is not possible
- to differentiate between morphotypes in our light microscopy images, we used the mean
- shape factor constant for *E. huxleyi* (i.e. $k_s = 0.0275$) to provide a range of coccolith mass
- estimates for this species (Table 1 and Fig. 4).
- 323 2.7 Calculation of annual estimates

Since the trap collection periods encompassed a period shorter than a calendar 324 325 year, annual estimates of coccolith and CaCO₃ fluxes and species relative abundances had 326 to be estimated. For the SOTS site, a total of 336 days were sampled for the 1000 and 2000 m traps and 338 days for the 3800 m. Since the unobserved interval occurred in 327 winter, the missing sampling period was filled using an average flux value of the winter 328 329 cups (first and last trap bottles). In the case of the SAM trap, the number of samples available for CaCO₃ and calcareous nannoplankton analyses was different, covering a 330 period of 313 and 191 days respectively. Since gaps were quasi-equally distributed along 331 the time series, annual fluxes were estimated by filling the gaps in the record with average 332 333 fluxes calculated from the available data. The estimated range of the annual contribution of coccolithophores to total CaCO₃ export at the SOTS and SAM traps was calculated by 334 335 multiplying the coccolith flux of each species in each sampling interval by its average 336 coccolith weight values obtained with the birefringence and morphometric techniques.

337 **2.8 Remotely sensed chlorophyll-***a* and PIC concentrations

Weekly Chlorophyll-a and PIC concentrations for the sampling intervals at the SOTS and SAM sites were derived from Giovanni online data system, developed and maintained by the NASA Goddard Earth Sciences Data Active Archive Center (Acker and Leptoukh, 2007). Each value is a weekly value is produced by computing spatial averages within the area 48.5-45.5°S and 130-150°E for the SOTS site and 47-45° S and 171°E-179°W for the SAM site (Fig. 5).

344

345 **3. RESULTS**

346 **3.1 Magnitude and seasonality of coccolithophore and CaCO₃ fluxes**

Annualized coccolith fluxes were similar at the SOTS three trap depths, with 8.6, 7.3 and 8.6 x 10^{11} liths m⁻² yr⁻¹ at 1000, 2000 and 3800 m respectively, and about three times larger than those of the SAM site $(3.0 \times 10^{11} \text{ liths m}^{-2} \text{ yr}^{-1})$. The contribution of intact coccospheres to the total coccolith export was low at both sites, with annual coccosphere fluxes two orders of magnitude lower than coccolith fluxes at SOTS (3.5, 3.3 and 1.8 x 10⁹ coccospheres m⁻² yr⁻¹ at 1000, 2000 and 3800 m, respectively) and SAM (2.2 x10⁹ coccospheres m⁻² yr⁻¹). Annualized CaCO₃ export was similar at both sites with 14.6, 16.2 and 17.1 g m⁻² yr⁻¹ at 1000, 2000 and 3800 m at the SOTS site and 13.9 g m⁻² yr⁻¹ at the SAM sediment trap (1500 m).

Both coccolith and coccosphere fluxes displayed a marked seasonality that 356 357 followed the general trend of algal biomass accumulation in the surface waters at the SOTS and SAM sites (Fig. 2). Coccolith fluxes at 1000 m started to increase in early 358 October and remained above the threshold of 1 x 10^9 coccoliths m² d⁻¹ until mid-April 359 (Fig. 2). Three maxima were recorded during the period of high coccolith export: 360 October-early November 2011 (4 x 10⁹ coccoliths m² d⁻¹), late December 2011 (9 x 10⁹ 361 coccoliths $m^2 d^{-1}$) and March 2012 (4 x 10⁹ coccoliths $m^2 d^{-1}$). Coccolith fluxes of the 362 main coccolithophore species generally followed the similar seasonal pattern to that of 363 364 the total coccolith flux (Supplementary figure 1) and are not discussed further. Coccolithophore fluxes registered by the 2000 and 3800 m sediment traps followed a 365 generally similar seasonal pattern to those of the shallower trap at the SOTS site (Fig. 2). 366 At SAM, coccolith fluxes exhibited a strong seasonality with peak fluxes in early January 367 2010 (up to 6 x 10^9 coccoliths m² d⁻¹) and a secondary peak in August 2010 (3 x 10^9 368 coccoliths m² d⁻¹). Coccosphere fluxes at both sites displayed maximum fluxes during the 369 370 austral summer and minima during winter; however maximum coccosphere export peaks 371 did not always match those of coccolith export (Fig. 2). The seasonality of total CaCO₃ 372 followed a similar pattern to coccolith fluxes with peak values in the spring-summer and minima during winter at both study sites. 373

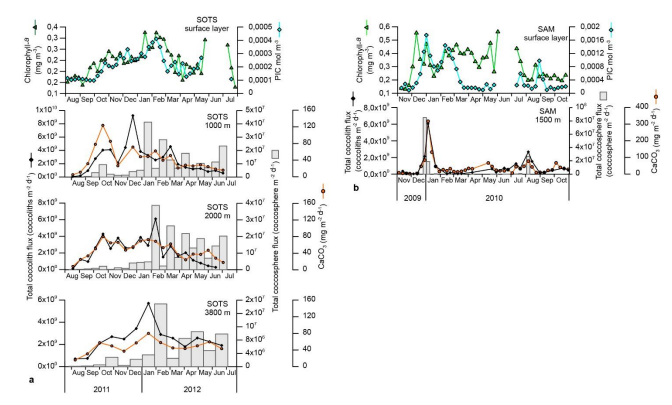


Figure 2: Ocean-colour satellite-derived chlorophyll-*a* and Particulate Inorganic Carbon
(PIC) concentration in the surface layer and total CaCO₃, coccolith and coccosphere
fluxes registered by the sediment traps at the SOTS (a) and SAM (b) sites.

378 **3.2.** Coccolithophore assemblage composition

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Coccolith sinking assemblages were overwhelmingly dominated by *Emiliania* 379 huxleyi for all sediment trap records analysed (Fig. 3a). At the SOTS site, the annualized 380 flux-weighted relative contribution of E. huxleyi decreased slightly with depth, 381 382 comprising 88% of the total coccolithophore assemblage at 1000 m, 82% at 2000 m and 383 80% at 3800 m. Secondary components of the coccolith sinking assemblage were Calcidiscus leptoporus (sensu lato) (6.8, 10.1 and 9.6% at 1000, 2000 and 3900 m, 384 385 respectively), *Heliscosphaera carteri* (1.4, 2 and 1.3%) and small *Gephyrocapsa* spp. (< 3 µm) (1.4, 1.5 and 4.7%). Background concentrations ($\leq 1\%$) of *Calciosolenia* spp., 386 387 Coccolithus pelagicus, Gephyrocapsa muellerae, *Gephyrocapsa* oceanica, 388 Gephyrocapsa spp. (> 3 μ m), Syracosphaera pulchra, Syracosphaera spp., 389 Umbellosphaera tenuis (sensu lato), and Umbilicosphaera sibogae were also registered. At the SAM site, E. huxleyi accounted for 83% of the annualized coccolith flux, with 390 391 subordinate contributions of C. leptoporus (12.2%) and Gephyrocapsa spp. (< $3 \mu m$) (1.5%). Background concentrations (< 1%) of *Calciosolenia* spp., *Coccolithus pelagicus*, 392

393 *G. oceanica, Gephyrocapsa muellerae, Gephyrocapsa* spp. (> 3 μm), *H. carteri*,
394 *Syracosphaera pulchra, Syracosphaera* spp., *U. sibogae* and *U. tenuis* were observed.

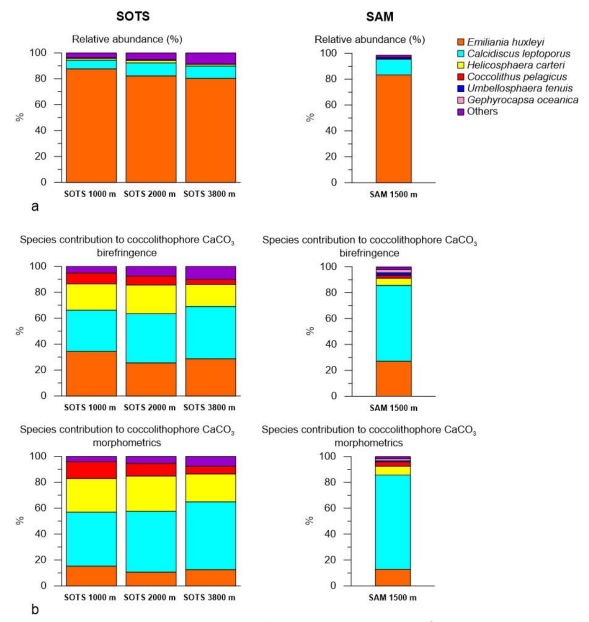


Figure 3: a. Annualized integrated relative abundance of the most important coccolithophore species in the SOTS and SAM sediment trap records. **b.** Fractional contribution of coccolithophore species to total coccolithophore CaCO₃ export in the SOTS and SAM sediment traps.

399 3.3 Calcite content per species

400 Coccolith length and mass for all species measured using birefringence and 401 morphometric techniques are provided in Table 1. Overall, the average coccolith mass 402 estimates for the coccolithophore species at SOTS and SAM sites using both approaches are within the range of values in the published literature. The Noelaerhabdaceae family
members, *G. oceanica* and *Gephyrocapsa* spp., display almost identical mass values with
both approaches (Fig. 4). In contrast, substantial discrepancies are identifiable for *C. pelagicus*, *C. leptoporus*, *H. carteri* and *U. sibogae*, for which coccolith mass estimates
are about two-fold greater using morphometrics than with the birefringence approach.
The range of annual contributions of coccolithophores to carbonate is illustrated in Figure
5.

Species and morphotypes	Type of measurement	n	Length (µm)		Mass CaCO ₃ (pg)		1.	Crystal units	D.6
			Average	SD	Average	SD	k,	types	References
Calcidiscus leptoporus	Birefrigence	210	6.39	1.49	33.65	21.11	-	V and R	
	Morphometrics	210	6.39	1.49	66.23	53.28	0.080		1
	Literature estimates	-	4.3-9.6		22.6-125.2		0.061-0.105		1,2
Coccolithus pelagicus	Birefrigence	54	13.28	1.14	170.90	32.33	-	V and R	
	Morphometrics	54	13.28	1.14	387.96	99.64	0.060		1
	Literature estimates	-	8.5-13.5		99.5-398.6		0.051 - 0.060		1,2,3
Emiliania huxleyi	Birefrigence	12842	2.78	0.57	2.64	1.43	-	R	
	Morphometrics	12842	2.78	0.57	0.99-2.64 (1.81)*	0.60-1.60	0.015-0.04 (0.0275)*	(V-unit vestigial)	
<i>E. huxleyi</i> type A	Literature estimates	-	3-4		1.50 - 3.50		0.02		1,4,5
<i>E. huxleyi</i> type A o/c	Literature estimates	-	3.5		4.6		0.04		1
<i>E. huxleyi</i> type B/C	Literature estimates	-	1.8-5.5		0.3-3.5		0.015		5,6,7
<i>E. huxleyi</i> type B	Literature estimates	-	3.5-5		2.30 - 6.81		0.02		1,5
Gephyrocapsa oceanica	Birefrigence	51	5.87	0.62	26.70	5.64	-	R	
	Morphometrics	51	5.87	0.62	28.14	8.97	0.050	(V-unit vestigial)	
	Literature estimates	-	5-5.35		16.9-25.7		0.050-0.062		1,2
Gephyrocapsa spp.	Birefrigence	10	4.03	0.59	9.00	2.51	-	R	
	Morphometrics	10	4.03	0.59	9.33	3.84	0.050	(V-unit vestigial)	1
	Literature estimates	-	-	-	-		-		
Helicosphaera carteri	Birefrigence	64	11.20	1.12	100.10	20.34	-	V and R	
	Morphometrics	64	11.20	1.12	194.95	56.45	0.050		1
	Literature estimates	-	9.1-10		135-142.8		0.050-0.070		1,2
Syracosphaera pulchra	Birefrigence	81	6.77	1.09	17.77	6.80	-	V, R and T	
	Morphometrics	81	6.77	1.09	26.94	11.16	0.030		1
	Literature estimates	-	2.7-6		13.5-16.5		0.027-0.083		1,2,4
Umbellosphaera tenuis	Birefrigence	54	6.42	0.99	15.69	5.02	-	R	
	Morphometrics	54	6.42	0.99	11.45	4.61	0.015		1
	Literature estimates	-	5-6		8.7-23.9		0.015-0.071		1,2
Umbilicosphaera sibogae Birefrigence		6	7.76	1.81	27.14	11.07	-	V and R	
	Morphometrics	6	7.76	1.81	78.93	51.38	0.055		1
	Literature estimates	-	4.1-6	-	16-35		0.055-0.086		1,2

 Table 1: Coccolith mass estimates of the main coccolithophore species found at the SOTS
 410 and SAM sites using birefringence (C-Calcita) and morphometrics. Additionally, length 411 and mass estimates from the literature are also listed for most species. References: (1) 412 413 Young and Ziveri (2000), (2) Beaufort and Heussner (1999), (3) Samtleben and Bickert (1990), (4) Poulton et al. (2010), (5) Poulton et al. (2011), (6) Holligan et al. (2010) and 414 415 (7) Charalampopoulou et al. (2016). * coccolith mass range obtained applying the 416 minimum and maximum k_s values for *E. huxleyi* found in the literature (i.e. 0.015 and 417 0.04, respectively).

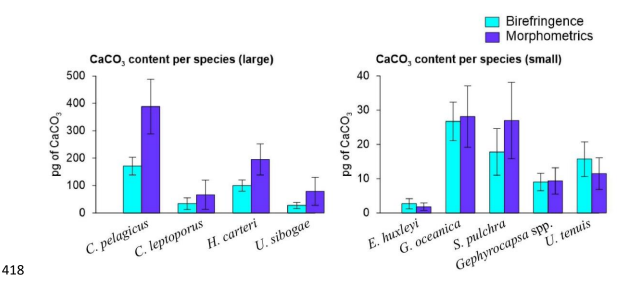
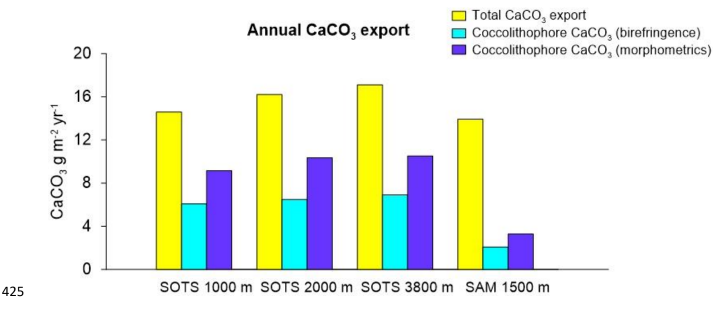
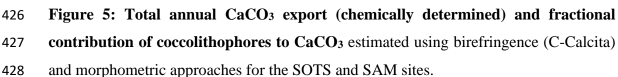


Figure 4: Average and standard deviation of the coccolith mass estimates of the most important coccolithophore species captured by the SOTS and SAM sediment traps using birefringence (C-Calcita) and morphometric approaches. For *E. huxleyi*, the morphometric-based coccolith mass estimate was calculated by applying a mean shape factor constant (k_s) value estimated from the range of all the morphotypes found at the SAZ (i.e. $k_s = 0.0275$, Table 1).





429 4. Discussion



Total coccolith flux seasonality at the SOTS site shows good congruence with satellite-431 432 derived PIC in the surface layer, with both parameters suggesting enhanced coccolithophore productivity between October and March (austral mid-spring to early 433 autumn; Fig. 2a). Interestingly, substantial coccosphere export (> 1×10^7 coccospheres 434 $m^2 d^{-1}$) does not occur until January indicating that coccolith and coccosphere export are 435 436 not tightly coupled in the subantarctic waters south of Australia. Two different processes could be invoked to explain the mismatch between coccolith and coccosphere fluxes at 437 438 this site. Firstly, E. huxleyi, the dominant coccolithophore species in the Southern Ocean, 439 is able to produce coccoliths rapidly (up to three coccoliths per hour; Paasche, 1962; 440 Balch et al., 1996) and shed the excess of coccoliths into the surrounding water under 441 certain environmental conditions (Paasche, 2002). Although the coccolith shedding rate 442 of E. huxleyi increases linearly with cellular growth rate (Fritz and Balch, 1996; Fritz, 443 1999), the tiny size and low weight of detached coccoliths allow them to remain in the upper water column long after cell numbers have begun to decline. It follows that high 444 445 concentrations of detached coccoliths do not necessary imply a proportional abundance 446 of coccospheres in the surface layer (Tyrrell and Merico, 2004; Poulton et al., 2013) or in 447 the traps. Additionally, a substantial fraction of the coccospheres produced in the surface layer may experience substantial mechanical breakage by zooplankton before reaching 448 the trap depths. Indeed, microzooplankton grazing pressure can remove up to 82% of 449 primary production in mid-summer in the subantarctic waters south of Tasmania 450 (Ebersbach et al., 2011) and about 60% of the daily coccolithophore growth in the North 451 452 Atlantic (Mayers et al., 2019), therefore suggesting a strong top-down control on coccolithophore populations. Additionally, a polyacrylamide gel sediment traps study in 453 454 the subantarctic waters south of Tasmania by Ebersbach et al. (2011) revealed that most 455 of the particles exported out the mixed layer during the productive period occur in the form of faecal aggregates. Therefore, it is highly likely that: (i) the intensity of 456 457 coccosphere export registered by the traps is influenced by grazing pressure in the surface 458 layer, and (ii) that the impact of grazing on coccolithophores varies throughout the year (Calbet et al., 2008; Lawerence and Menden-Deuer, 2012; Quéguiner, 2013). 459

In contrast, seasonal variations in satellite-derived PIC concentration and coccolith fluxes at SAM show some discrepancies not observed at SOTS. While maximum PIC concentrations in the surface layer and coccolith and coccosphere fluxes co-occur in December and January (austral early to mid-summer), satellite-derived PIC suggests a secondary maximum in February-early-March not recorded by the trap (Fig.

2b). One possibility is that the satellite secondary maximum is not coccoliths. The higher 465 466 chlorophyll-a levels at the SAM site (Fig. 2) suggests that other phytoplankton groups, such as diatoms, are more abundant than in the subantarctic waters south of Tasmania. 467 468 Empty and broken diatom valves have been suggested to display similar spectral characteristics than those of coccolithophore blooms (Broerse et al., 2003; Tyrrell and 469 470 Merico, 2004; Winter et al., 2014). Therefore, the second peak in satellite-derived PIC could have been caused by a senescent diatom bloom. This hypothesis is likely since 471 472 diatom blooms in the SAZ are known to develop early in the productive season (Rigual-473 Hernández et al., 2015b) and rapidly decay following the depletion of silicate and/or iron 474 stocks in the surface layer (Lannuzel et al., 2011). However, no secondary late summer 475 maximum was observed in biogenic silica fluxes in the SAM. Another possible 476 explanation is a contribution to the satellite record from lithogenic material or storm-477 induced microbubble injection (Zhang et al., 2002). Fully resolving causes of mismatches 478 between in-situ and satellite PIC estimates is not achievable for the SAM site (nor more 479 broadly for the Southern Ocean; Trull et al., 2018).

480 A second difference between the SAM and SOTS sites is that maximum annual 481 coccosphere export occurred one week earlier than maximum coccolith fluxes at SAM, (Fig. 2). The different seasonalities between the sites suggest that different export 482 483 mechanisms may operate. The formation of rapidly sinking algal aggregates by diatoms is known to scavenge particles they have collided with and increase particle sinking 484 (Alldredge and McGillivary, 1991; Passow and De La Rocha, 2006), thus the formation 485 of such rapid-sinking aggregates could potentially facilitate the preservation of 486 coccospheres early in the productive season at the SAM site. However, the lack of 487 488 accompanying in situ information on plankton community structure in the study region 489 precludes the assessment of these hypotheses.

Despite the uncertainties involved in our interpretations, the overall picture that emerges from our comparison of satellite and sediment trap flux data is that the duration of the coccolithophore bloom based on ocean-colour-based PIC concentrations most likely provides an over-estimation of the coccolithophore productive season. Our observations motivate caution in describing coccolithophore phenology solely based on satellite-derived PIC concentrations (e.g. Hopkins et al., 2015).

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497 **4.2 Magnitude and composition of subantarctic coccolithophore assemblages**

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498 Annual coccolith export across the major zonal systems of the Australian sector 499 of the Southern Ocean exhibits a clear latitudinal gradient, with maximum fluxes at the SAZ (8.6 x 10^{11} liths m⁻² yr⁻¹) and eight-fold lower fluxes in the polar waters of the AZ 500 (1.0 x 10¹¹ liths m⁻² vr⁻¹; Rigual Hernández et al., 2018). Coccolithophore species 501 502 occurrence documented by our subantarctic sediments traps are consistent with previous 503 reports on coccolithophore assemblage compositions in the surface layer (Findlay and Giraudeau, 2000; Saavedra-Pellitero et al., 2014; Malinverno et al., 2015; Chang and 504 Northcote, 2016) and sediments (Findlay and Giraudeau, 2000; Saavedra-Pellitero and 505 506 Baumann, 2015) and are more diverse than those found in the AZ (Rigual Hernández et 507 al., 2018). The southward decline in coccolithophore abundance and diversity is most 508 likely due to the decrease in sea-surface temperature (SST) and light availability moving 509 poleward (Charalampopoulou et al., 2016; Trull et al., 2018). In particular, the close 510 relationship between temperature and growth rates has been demonstrated in a range of coccolithophore species and strains (Buitenhuis et al., 2008), and seems to be a critical, 511 512 if not the most important, control on the biogeographical distribution of coccolithophore 513 species in the Southern Ocean (Trull et al., 2018). This pronounced latitudinal change in 514 coccolithophore assemblage composition contrasts with the little longitudinal variability 515 between the subantarctic SOTS and SAM sites (Fig. 3). These observations underscore 516 the role of circumpolar fronts as natural physical barriers for plankton species distribution in the Southern Ocean (Medlin et al., 1994; Boyd, 2002; Cook et al., 2013). 517

Notably, the rare occurrence of the cold-water species Coccolithus pelagicus at 518 519 the SOTS and SAM sites contrasts with the high contribution of C. pelagicus to the living 520 coccolithophore communities in the subpolar and polar waters of the North Atlantic and 521 North Pacific oceans, where it is often the second most abundant species after E. huxleyi 522 (McIntyre and Bé, 1967; Baumann et al., 2000; Broerse et al., 2000a; Broerse et al., 2000b; Ziveri et al., 2000). This important difference in species composition between 523 Northern and Southern hemisphere subpolar ecosystems could have important 524 525 implications in the calibration of the satellite PIC signal in the Southern Ocean. Previous research in the Southern Ocean comparing satellite and shipboard observations identified 526 527 a substantial over-estimation of coccolithophore PIC in the Southern Ocean waters by 528 satellite ocean-colour-based PIC algorithms (Holligan et al., 2010; Trull et al., 2018). 529 Since satellite reflectance observations are mainly calibrated against Northern Hemisphere PIC results (Balch et al., 2011; Balch et al., 2016), the lower the calcite 530 531 content of dominant E. huxleyi morphotypes (B/C) in the Southern Ocean compared to

their northern hemispheric counterparts has been suggested as a possible factor causing 532 533 the over-estimation of PIC concentrations in the Southern Ocean. Following this reasoning, we speculate that differences in other components of the coccolithophore 534 535 assemblages, and particularly, differences in C. pelagicus numbers, could contribute to the over-estimation of PIC concentrations by the satellite PIC algorithm in the Southern 536 537 Ocean. Indeed, the scaling of reflectance (in satellite images) to PIC (in ocean) is very dependent on coccolith area:mass ratios (Gordon and Du, 2001; Balch et al., 2005a). 538 Coccolithus pelagicus has remarkably heavier and thicker coccoliths (100-400 pg per 539 540 coccolith; Table 1) than E. huxleyi (~3 pg per coccolith), i.e. about 100 times heavier. 541 However, the average coccolith area of C. pelagicus is only about ten times greater than 542 that of E. huxleyi. Thus, this lack of proportional relationship between area and mass 543 between these species should be taken into consideration when calibrating the satellite 544 signals of coccolithophore-related PIC in the Southern Ocean. However, it should be 545 noted that this is only one possible factor contributing to the overestimation of PIC 546 concentrations in Southern Ocean waters. Other factors such as the presence of 547 microbubbles -- that are a source of enhanced reflectance -- must also play an important 548 role (Balch et al., 2011).

549

550 **4.3** Coccolith calcite content of subantarctic coccolithophore species

551 Multiple methodological biases associated with each of the methods used for 552 estimating coccolith calcite content (i.e. birefringence, morphometrics) could be invoked 553 to explain the different estimates observed for some of the species (see Young and Ziveri, 2000; Fuertes et al., 2014 and references therein). However, the fact that these 554 555 discrepancies vary greatly across species suggests that the composition of the crystalunits of the coccoliths could be the most important factor causing these differences. All 556 557 the heterococcoliths of the species analysed are mainly composed of either V- or R-558 calcite crystal units or a combination of both (Young et al., 2003; Table 1). R units are 559 characterized by sub-radial c-axes that are reasonably well measured by the birefringence 560 technique, but, the almost vertical optical axes of the V units (Young, 1992; Young et al., 561 1999) make the same thickness less birefringent (Fuertes et al., 2014). Thus, it is likely 562 that differences in the birefringence properties of the R and V units could be responsible for the different estimates provided by the two approaches. This is supported by our 563 results which show coccolith mass estimates of those species composed of R units, such 564

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as G. oceanica and Gephyrocapsa spp. exhibit almost identical values with both 565 566 techniques (Table 1). In contrast, those species with coccoliths composed by a combination of R and V units, such as C. pelagicus, C. leptoporus, H. carteri and U. 567 sibogae, display divergent mass estimates between approaches. The case of E. huxleyi is 568 more complex due to the large intraspecific genetic variability that results in substantial 569 570 differences in the profile and degree of calcification between specimens (Young and 571 Ziveri, 2000). Our birefringence mass estimate for *E. huxleyi* (2.67 ± 1.49 pg) is less than 572 one picogram lower than the mean range value calculated with the morphometric 573 technique (i.e. 1.81 ± 1.10 pg with an average k_s value of all the morphotypes found at 574 the SAZ, i.e. $k_s = 0.0275$), but identical to the maximum (2.64 ± 1.60 pg; using $k_s = 0.04$). 575 These results suggest a reasonably good consistency between techniques for E. huxleyi.

Taking into consideration all the above, it is likely that the coccolith mass of some species is underestimated by the birefringence technique, and therefore, the fractional contribution of coccolithophores to total PIC using this approach should be taken as a conservative estimate. Since both methods for estimating calcite content have inherent uncertainties, the range of values provided by both techniques is used here as an approximation of the fractional contribution of coccolithophores to total annual CaCO₃ export to the deep sea in the Australian and New Zealand sectors of the SAZ.

4.4 Contribution of coccolithophores to carbonate export in the Australian-New Zealand sectors of the Southern Ocean

The magnitude of the total PIC export in the subantarctic waters was similar 585 between the SOTS and SAM sites (range 14-17 g m⁻² yr⁻¹), and thus slightly above the 586 global average (11 g m⁻² yr⁻¹; Honjo et al., 2008). Our estimates indicate that 587 coccolithophores are major contributors to CaCO₃ export in the Australian and New 588 Zealand waters of the SAZ, accounting for 40-60% and 15-25% of the annual CaCO₃ 589 590 export, respectively (Fig. 5). Heterotrophic calcifiers, mainly planktonic foraminifera 591 (Salter et al., 2014), must therefore account for the remainder of the annual CaCO₃ export 592 at both sites. Previous work on foraminifera fluxes in our study regions allows an 593 approximate estimate of the contribution of foraminifera to total $CaCO_3$ flux that can be 594 used to assess the validity of our estimates. Combining counts of foraminifera 595 shells (King and Howard, 2003) with estimates of their average shell weights (20-40 µg per shell depending on size; Moy et al., 2009) suggests contributions of 1/3 to 2/3 of 596 597 planktonic foraminifera to the total CaCO₃ flux in the Australian SAZ (Trull et al., 2018).

In the subantarctic waters south of New Zealand, Northcote and Neil (2005) estimated 598 599 that planktonic foraminifera accounted for about 78-97% of the total CaCO₃. Thus, 600 estimations of the contribution of heterotrophic calcifiers to total carbonate in both study 601 regions are in reasonable agreement with our coccolithophore CaCO₃ estimates at both 602 sites. The lower contribution of coccolithophores to CaCO₃ export at the SAM site in 603 comparison with that of SOTS may be explained by differences in the ecosystem structure between sites. Algal biomass accumulation in the surface waters of the SAM region 604 (average chlorophyll-a concentration between 2002 and 2018 is 0.31 mg m⁻³) is 605 substantially higher than that at SOTS (0.23 mg m^{-3}) . We speculate that the higher 606 abundance of non-calcareous phytoplankton (e.g. diatoms) in the subantarctic waters 607 608 south of New Zealand could simultaneously reduce coccolithophore biomass through 609 resource competition (Quéré et al., 2005; Sinha et al., 2010) while stimulating 610 foraminifera growth (Schiebel et al., 2017). The combination of both factors could be 611 responsible for the lower coccolithophore productivity at the SAM site despite similar 612 total CaCO₃ export. Assuming that both the SOTS and SAM sites can be considered 613 representative of a broad longitudinal swath of the SAZ south of Australia and New 614 Zealand (ca. 1% of areal extent of the global ocean), the coccolithophore CaCO₃ export in these two regions together account for approximately 0.4 Tmol C_{inorg} yr⁻¹. This value 615 represents approximately 1% of the global annual PIC export to the deep ocean (Honjo et 616 617 al., 2008) and underscores the notion that the high nutrient low-chlorophyll waters of the circumpolar SAZ should not be taken as indicative of low biological activity or export. 618

Our results indicate that although E. huxleyi overwhelmingly dominates the 619 620 coccolithophore sinking assemblages at both study sites, other species with lower relative 621 contribution but substantially heavier coccoliths are more important contributors to the annual coccolithophore CaCO₃ export budget (Fig. 3). Particularly relevant is the case of 622 623 C. leptoporus that despite its relatively low abundance (~ 10% of the annual assemblage 624 at both sites; Fig. 3), it accounts for between 30-50% and 60-70% of the annual 625 coccolithophore-CaCO₃ export at the SOTS and SAM sites, respectively (Fig. 3). 626 Similarly, other species with heavy coccoliths, such as *H. carteri* and *C. pelagicus*, are 627 important contributors to the annual coccolithophore PIC export to the deep sea (up to ~30% and ~10% of the annual coccolithophore PIC, respectively) despite their low annual 628 629 relative abundance (<2% at both sites) (Fig. 3). These results serve as an important reminder that it is often not the most abundant species, but rather the largest 630

631 coccolithophore species that account for the greatest contribution to coccolithophore
632 CaCO₃ production and export (Young and Ziveri, 2000; Baumann et al., 2004; Daniels et
633 al., 2016).

The important contribution made by the coccolithophore community in setting the 634 635 magnitude of carbonate production and export to the deep sea is evidenced when we 636 compare the coccolith and total CaCO₃ fluxes of the SOTS sediment trap with those 637 deployed in the AZ along the 140°E meridian (Fig. 1). Although both total and coccolithophore CaCO₃ export decrease with increasing latitude these changes are largely 638 639 uneven. While total CaCO₃ decreases two-fold from the SAZ to the AZ, coccolithophore 640 $CaCO_3$ export decreases 28-fold (Supplement Figure 2). This lack of proportional 641 latitudinal change can be attributed to two main factors. First, subantarctic coccolithophore populations are diverse and relatively rich in species with large and 642 643 heavy coccoliths such as C. leptoporus or H. carteri that account for a large fraction of 644 the annual carbonate production and export. South of the PF, assemblages become 645 monospecific, or nearly monospecific, dominated by the small and relatively lightly 646 calcified E. huxleyi. Second, latitudinal variations in the abundance of heterotrophic 647 calcifiers (mainly foraminifera but also pteropods) must play a major role in modulating the observed variations in CaCO₃ export. In particular, our data suggests that the 648 649 fractional contribution of heterotrophic calcifiers to CaCO₃ production increases from 650 ~40-60 % in the Australian SAZ to up to 95% in the AZ (Rigual Hernández et al., 2018). 651 This pattern is consistent with previous shipboard and sediment trap studies that reported higher abundances of planktonic foraminifera at the PFZ and AZ compared to that of the 652 653 SAZ in the Australian sector (King and Howard, 2003; Trull et al., 2018). Controls on the 654 biogeographic distribution of foraminifera species are complex and beyond the scope of 655 this paper, however, we provide a few observations. Both temperature and diet are critical 656 factors controlling the spatial distribution of planktonic foraminifera species. In 657 particular, the lower temperatures south of the SAF seem to favour the development of 658 *Neogloboquadrina pachyderma* sin. and *Turborotalita quinqueloba* as indicated by the 659 high abundance of these species in the PFZ (> 80% of the annual integrated flux for both 660 species together; King and Howard, 2003). Additionally, the dramatically different algal 661 communities dwelling in each zonal system may also play a role in planktonic 662 foraminifera species distributions. In particular, diatoms can account for a major part of the diet of some foraminifera species, including N. pachyderma (Schiebel and Hemleben, 663

2017). Therefore, it is likely that the preferential grazing on diatoms of some foraminifera
species may play an important role in the increase of foraminifera CaCO₃ production
moving poleward.

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4.5 Future predictions of coccolithophore community response to environmental change in the subantarctic zone

670 The response of *E. huxleyi* to environmental change has been extensively studied 671 in laboratory experiments (Meyer and Riebesell, 2015; Müller et al., 2015; Feng et al., 672 2017) and the available information is sufficient to propose possible changes of its niche and calcification in the Southern Ocean, as discussed in detail in Trull et al. (2018) and 673 674 Krumhardt et al. (2017). Due to the ubiquity and abundance of E. huxleyi, the 675 ecophysiology of this species is often regarded as typical of all coccolithophores. 676 However, E. huxleyi is rather different from most other coccolithophore species in that its 677 physiological adaptations place it in the upper limit of the *r*-*K* ecological gradient of these 678 organisms (i.e. an opportunistic species), while most of the other species are located at 679 the opposite end of the spectrum (i.e. conservative or K-selected species) (Probert and Houdan, 2004). Our results demonstrate that E. huxleyi plays an important, but not 680 681 dominant role in CaCO₃ export, with other species such as C. leptoporus, H. carteri or C. 682 pelagicus making a larger contribution to the annual CaCO₃ export in the SAZ (Fig. 3). 683 Therefore, it is of critical importance to evaluate how these other biogeochemically 684 important coccolithophore species will respond to projected climate-induced changes in 685 the Southern Ocean. Here, we now assess the response of large coccolithophore species to projected changes in temperature and carbonate chemistry, that have been highlighted 686 687 among the most important environmental stressors expected to impact Southern Ocean coccolithophore physiological rates (Müller et al., 2015; Charalampopoulou et al., 2016; 688 689 Feng et al., 2017; Trull et al., 2018).

The Southern Ocean is warming rapidly (Gille, 2002; Böning et al., 2008), largely due to the southward migration of the ACC fronts (Sokolov and Rintoul, 2009). Only between 1992 and 2007 the position of Southern Ocean fronts shifted by approximately 60 km to the south (Sokolov and Rintoul, 2009) and this trend may continue throughout the next century. Therefore, it is likely that any further southward migration of ACC fronts will be coupled with an expansion of subantarctic coccolithophore species towards higher latitudes. The poleward expansion of *E. huxleyi* geographic range has already been

suggested in the Southern Ocean (Cubillos et al., 2007; Winter et al., 2014; 697 698 Charalampopoulou et al., 2016) and it also appears to be occurring in the North Atlantic 699 (Rivero-Calle et al., 2015; Neukermans et al., 2018). Given the important contribution of 700 large subantarctic coccolithophore species to CaCO₃ export, the expansion of their 701 ecological niche could result in a substantial increase in CaCO₃ production and export in 702 the Southern Ocean. However, this may not be the future scenario for the SAZ southeast 703 on New Zealand, where bathymetry strongly controls the location of ocean fronts (Fernandez et al., 2014; Chiswell et al., 2015). If the fronts are bathymetrically 'locked', 704 705 then the SAZ will not expand in areal extent, although the region is still predicted to 706 undergo significant physical, biogeochemical and biological changes (Law et al., 2017) 707 that will have likely flow-on effects on coccolithophore productivity and export 708 (Deppeler and Davidson, 2017).

709 The available carbonate chemistry manipulation experiments with C. leptoporus have come to different conclusions. While some studies identified an increase in coccolith 710 711 malformations with increasing CO₂ concentrations (Langer et al., 2006; Langer and Bode, 712 2011; Diner et al., 2015), another study (Fiorini et al., 2011) reported no changes in the 713 calcification of C. leptoporus at elevated pCO₂. Interestingly, C. leptoporus did not 714 experience changes in its photosynthesis rates over the tested CO₂ range in any of the aforementioned studies. The most likely explanation for the different results between the 715 studies is a strain-specific variable responses to changing carbonate chemistry (Diner et 716 717 al., 2015). Strain-specific variability in response to changing carbonate chemistry has 718 been previously reported in other coccolithophores, such as E. huxleyi (Langer et al., 719 2009; Müller et al., 2015), and therefore it is likely that this also occurs in other species. 720 Given the fact that Southern Ocean fronts act as barriers for species distributions and gene 721 flows (Medlin et al., 1994; Patarnello et al., 1996; Thornhill et al., 2008; Cook et al., 722 2013), it is possible that the subantarctic C. leptoporus populations exhibit a different 723 ecophysiology than those used in the above mentioned laboratory experiments. Prediction 724 of the responses of *H. carteri* and *C. pelagicus* is even more challenging due to the lack of experiments testing the response of these species to changing seawater carbonate 725 726 chemistry. The only available insights in the response of one of these species to ocean 727 acidification are found in the fossil record. Both Gibbs et al. (2013) and O'Dea et al. 728 (2014) reconstructed the evolution of C. pelagicus populations during the Palaeocene-729 Eocene Thermal Maximum (PETM), a period arguably regarded as the best geological 730 approximation of the present rapid rise in atmospheric CO₂ levels and temperatures.

These studies concluded that C. pelagicus most likely reduced its growth rates and 731 732 calcification during this period. This limited number of studies suggest that the ongoing ocean acidification in the Southern Ocean could potentially have a negative impact on the 733 physiological rates of C. leptoporus and C. pelagicus while the effect on H. carteri is 734 unknown. Physiological response experiments (e.g. Müller et al., 2015) with Southern 735 736 Ocean strains of C. leptoporus, H. carteri and C. pelagicus are, therefore, urgently needed to be able to quantify the effect of projected changes in oceanic conditions in the SAZ on 737 their physiological rates and consequent effects on carbon cycling in the Southern Ocean. 738

Our synthesis suggests opposing influence of environmental stressors on subantarctic coccolithophore populations. Poleward migration of fronts will likely increase coccolithophore CaCO₃ production in the Southern Ocean, while changes in carbonate chemistry speciation will reduce growth rates of subantarctic coccolithophores. It seems possible that coccolithophores will initially expand southward as waters warm and fronts migrate, but then eventually diminish as acidification overwhelms those changes.

746

747 Acknowledgments

This project has received funding from the European Union's Horizon 2020 research and 748 innovation programme under the Marie Skłodowska-Curie grant agreement number 749 750 748690 - SONAR-CO2 (ARH, JAF and FA). The SOTS mooring work was supported 751 by IMOS, the ACE CRC, and the Australian Marine National Facility. The work at SAM 752 was supported by funding provided by the New Zealand Ministry of Business, Innovation 753 and Employment and previous agencies, and most recently by NIWA's Strategic Science 754 Investment Fund. NIWA is acknowledged for providing capital grants for mooring equipment purchases, and thanks to all the NIWA scientists, technicians and vessels staff, 755 756 who participated in the New Zealand biophysical moorings programme (2000-12). 757 Cathryn Wynn-Edwards (IMAS) provided support in sample splitting/processing and 758 laboratory analysis. Satellite Chlorophyll-a and PIC data sets were produced with the 759 Giovanni online data system, developed and maintained by the NASA GES DISC. We 760 thank Griet Neukermans and Alex Poulton for their constructive comments and 761 suggestions that helped improve and clarify this manuscript.

762

763 Author contributions

- 764 TWT, SDN, DMD and LN planned and performed the field experiment. ARH led the
- coccolithophore study and performed sample processing and microscopy and image
- analyses. AMB and ARH performed SEM analyses. ARH and SN preformed numerical
- analyses. ARH wrote the paper with feedback from all authors.

- 768 **Competing interests**
- 769 The authors declare no competing interests.
- 770

771 Data Availability

- 772 Morphometric data of major coccolithophore species generated during the current study are listed
- in Table 1, while species relative abundance and species fluxes (plotted in Supplement Figure 1)
- can be accessed in the following link:
- 775 https://data.aad.gov.au/metadata/records/Coccolithophore_Fluxes_SAZ_2009-2012.
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