

Dimethylated sulfur compounds in symbiotic protists: A potentially significant source for marine DMS(P)

Andres Gutierrez-Rodriguez, Loic Pillet, Tristan Biard, Ward Said-Ahmad, Alon Amrani, Rafel Simó, Fabrice Not

► To cite this version:

Andres Gutierrez-Rodriguez, Loic Pillet, Tristan Biard, Ward Said-Ahmad, Alon Amrani, et al.. Dimethylated sulfur compounds in symbiotic protists: A potentially significant source for marine DMS(P). Limnology and Oceanography, 2017, 62 (3), pp.1139-1154 10.1002/lno.10491. hal-01480231

HAL Id: hal-01480231 https://hal.sorbonne-universite.fr/hal-01480231v1

Submitted on 1 Mar 2017

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

1	DIMETHYLATED SULFUR COMPOUNDS IN SYMBIOTIC PROTISTS: A
2	POTENTIALLY SIGNIFICANT SOURCE FOR MARINE DMS(P)
3	
4	Andres Gutierrez-Rodriguez ^{1,4,*} , Loic Pillet ^{1,5} , Tristan Biard ¹ , Ward Said-Ahmad ² ,
5	Alon Amrani ² , Rafel Simó ³ and Fabrice Not ¹
6	
7	¹ Sorbonne Universités, UPMC Université Paris 06, CNRS, Laboratoire Adaptation et
8	Diversité en Milieu Marin UMR7144, Station Biologique de Roscoff, 29680 Roscoff, France
9	² The Institute of Earth Sciences, The Hebrew University, Jerusalem 91904, Israel
10	³ Department of Marine Biology and Oceanography, Institut de Ciencies del Mar
11	(CSIC), Pg. Maritim de la Barceloneta 37-49,08003 Barcelona, Catalonia, Spain
12	⁴ Current address: National Institute of Water and Atmospheric Research, Private Bag
13	14-901, Wellington 6241, New Zealand
14	⁵ Current address: Department of Genetics and Evolution, University of Geneva, 30,
15	qui Ernest-Ansermer, 1211 Geneva 11, Switzerland.
16	
17	
	*Corresponding author email: <u>Andres.gutierrez@niwa.co.nz</u>
18	
19	Running head: Plankton symbiosis and DMS(P) cycling
20	Key words: Radiolaria, Symbiosis, DMS(P)
21	
22	

23 ABSTRACT

Symbiosis with micro-algae (photosymbiosis) is a common feature among marine planktonic 24 protists, but very little is known about the physiology and ecological significance of these 25 associations. High concentrations of dimethylsulfoniopropionate (DMSP), a metabolite 26 produced by marine microalgae, are commonly observed in coral-microalgae symbiosis, 27 where DMS(P) is involved in multiple physiological functions. Knowledge on concentration 28 29 and role of DMSP in analogous photosymbiosis in plankton is lacking. Here, we assess the 30 total DMSP (DMSPt=DMSP+DMS) concentration and natural stable isotopes of sulfur across ecologically relevant symbiotic plankton groups, the Radiolaria and Foraminifera. We found 31 that intracellular DMSPt concentrations in microalgal symbionts were among the highest 32 recorded (range=170-702 mmol L⁻¹), while lower concentrations (range=0.1-23 mmol L⁻¹) 33 were characteristic of the holobiont (i.e. host-microalgae). The contribution of symbiotic 34 35 Radiolaria to the water column particulate DMSPt concentration ranged 0.1-8%. Sulfur isotopic composition (³⁴S) of DMSPt in the Collodaria holobionts was significantly higher 36 than their symbiotic microalgae isolated in culture. Despite their high intracellular DMSPt 37 content, SO_4^{2-} uptake in these holobionts throughout 3-day incubations was not detected. We 38 observed a systematic ³⁴S depletion (~1.5‰) of DMS relative to DMSP in experimental 39 incubations containing filtered seawater, which we hypothesize is related to the bacterial 40 preference for the uptake of ³⁴S-depleted DMS. Overall, the results indicate that plankton 41 symbiosis can, at times, represent a potentially important source of DMS(P). Specific 42 differences in ³⁴S provided new insights into sulfur isotopic fractionation associated with 43 DMS(P) biotransformation processes, with potential implications for current interpretations of 44 isotopically tracked biogenic sources of marine aerosols. 45

46 INTRODUCTION

Dimethylsulfoniopropionate (DMSP) is a widespread metabolite in marine 47 ecosystems, mainly produced by marine microalgae (Keller et al. 1989; Blunden et al. 1992; 48 Raina et al. 2013). It is involved in multiple cross-scale processes, from cell physiology to 49 ecosystem functioning. Physiologically, DMSP has been demonstrated to play an important 50 role in osmotic acclimation (Vairavamurthy et al. 1985; Kirst 1990; Lyon et al. 2016) and 51 cryoprotection in polar algae (Kirst et al. 1991; Karsten et al. 1996). DMSP production and 52 breakdown have been also hypothesized to act as an overflow mechanism to get rid of excess 53 reduced sulfur (Stefels 2000), to confer antioxidant protection by scavenging intracellular 54 55 hydroxyl radical (Sunda et al. 2002; Bucciarelli et al. 2013; Deschaseaux et al. 2014), and to participate in the regulation of phytoplankton buoyancy through replacement of other organic 56 solutes (Lavoie et al 2015, 2016). Beyond cellular limits, recent work suggests that DMSP 57 and its cleavage product dimethylsulfide (DMS) act as efficient info-chemicals among 58 plankton microorganisms (Seymour et al. 2010; Garces et al. 2013) and with higher trophic 59 levels (Savoca and Nevitt 2014). DMS is a volatile compound that ventilates to the 60 atmosphere and represents the main global biogenic source of atmospheric sulfur (Bates et al. 61 1992; Simó 2001). Several oxidation processes in the atmosphere transform DMS into 62 sulfuric and methanesulfonic acids, which are key participants of cloud formation (Andreae 63 1997). Three decades after it was postulated, the hypothesis of a feedback loop between 64 phytoplankton DMS production, cloud formation and climate regulation (Charlson et al. 65 1987), remains controversial. Issues like the relative contribution of DMS oxidation products, 66 sea salt and organics to the number of cloud condensation nuclei are yet to be resolved (Quinn 67 68 and Bates 2011; Lana et al. 2012).

In addition to phytoplankton, heterotrophic organisms containing endosymbiotic 69 microalgae (i.e. photosymbiotic holobionts) can be an important source of DMS(P) (Hill et al. 70 2000; Broadbent et al. 2002; Van Alstyne et al. 2006). Indeed, coral reefs hold the highest 71 natural concentrations of oceanic DMS(P) reported to date (Broadbent and Jones 2004). The 72 production of DMSP in corals is seemingly associated with the endosymbiotic dinoflagellate 73 Symbiodinium sp. (Van Alstyne et al. 2009), although a recent study showed capacity for the 74 juvenile animal hosts to produce DMSP as well (Raina et al. 2013). Less studied than that of 75 corals, photosymbiosis is a common feature in the planktonic realm, particularly among 76 protists belonging to the Rhizaria eukaryotic super-group (e.g. Foraminifera and Radiolaria) 77 (Stoecker et al. 2009; Nowack and Melkonian 2010; Decelle et al. 2015). Radiolaria are 78 amoeboid protists exhibiting mineral skeletons that are abundant and widespread in modern 79 oceans (Anderson 1983; Stemmann et al. 2008; Not et al. 2009; Biard et al. 2016). They 80 81 include five major groups - Collodaria, Nassellaria, Spumellaria, Taxopodia and Acantharia spanning a large size range, from a few micrometers for small solitary cells up to several 82 centimeters for colonial forms of the Collodaria (Dennett et al. 2002; Caron et al. 2012; 83 Suzuki and Not 2015), although the bulk of cell sizes range between 200-500 µm (Michaels 84 1988; Caron and Swanberg 1990). They are active predators, but many species dwelling in 85 the surface layers harbor endosymbiotic microalgae in their cytoplasm, which allows them to 86 thrive in ecological niches that otherwise would be less favorable (Decelle et al. 2015). 87

These symbiotic relationships involve essentially dinoflagellate microalgae such as *Brandtodinium nutricula* (Probert et al. 2014) and *Gymnoxanthella radiolariae* (Yuasa et al. 2016) in the case of Radiolaria, or *Pelagodinium beii* (Siano et al. 2010) in the case of Foraminifera. The Prymnesiophyceae species *Phaeocystis* sp., an abundant and widespread microalgal genus, has been recently described in symbiosis with acantharians (Decelle et al.

2012). Both Dinophyceae and Prymnesiophyceae classes are typical major DMSP producers 93 (Keller et al. 1989; Caruana and Malin 2014), and include keystone species for the 94 biogeochemical sulfur and carbon cycles (Malin and Steinke 2004; Schoemann et al. 2005). 95 Acantharia-*Phaeocystis* sp. holobionts exhibit extremely high DMSP cellular content, with 96 values significantly higher than those expected if all DMSP was contained in the 97 endosymbiotic microalgae (Decelle et al. 2012). Should elevated cellular content of DMSP 98 be a common feature not only in benthic, but also in planktonic photosymbiosis, this 99 widespread but traditionally overlooked functional group of plankton (i.e. photosymbiotic) 100 may constitute a relevant source of biogenic DMS(P) previously unaccounted by the standard 101 microplankton oriented (*i.e.* <200 µm) sampling procedures. 102

The complexity of marine biogeochemistry makes it difficult to identify and assess the 103 multiple biological sources and flows of DMS(P) from and through the different 104 compartments of the pelagic ecosystem, and their links to the lower atmosphere (Simó, 2001). 105 In this context, the sulfur isotope ratio (${}^{34}S/{}^{32}S$; *i.e.* $\delta^{34}S$) in aerosol sulfate has been used to 106 assess the contribution of different sources, mainly anthropogenic vs marine (Kaye 1987; 107 Norman et al. 1999; Patris et al. 2002). Recent pioneering measurements in macroalgae and 108 natural phytoplankton assemblages have provided similar δ^{34} S values for DMSP but have 109 110 contradictory views of the isotopic fractionation associated with biotransformation processes in the formation of DMS (Oduro et al. 2012; Amrani et al. 2013). In natural planktonic 111 systems, phytoplankton cells are often too small and too intermixed with other organisms to 112 be isolated in sufficient quantities for analysis of sulfur isotopic composition of DMSP by 113 conventional isotope ratio analysis methods. This hampers the characterization of the isotopic 114 composition of specific phytoplankton taxa or even functional groups in the field, and little is 115 known about the contribution of the different phytoplanktonic components to the bulk 116

community isotopic composition of DMSP. The new method of Compound Specific Sulfur 117 Isotope Analysis (CSSIA) enables sub-nanogram level sensitivity (Amrani et al. 2009; Said-118 Ahmad and Amrani 2013) and opens the door for such single-cell level studies. The present 119 study aims at opening this black box and refining our understanding of the different biogenic 120 sources and transformations of dimethyl sulfur compounds in planktonic systems, with 121 particular attention to the contribution of widespread photosymbiotic organisms. We have 122 combined CSSIA with culture and field-based approaches for free-living phytoplankton and 123 single-celled symbiotic Rhizaria (Radiolaria and Foraminifera) collected from different 124 environments. The specific objectives of our study were to assess i) the cellular DMSP 125 content, ii) the S-isotopic composition of DMSP and iii) the potential fractionation associated 126 with DMSP biosynthesis and degradation to DMS in photosymbiotic Rhizaria and 127 phytoplankton. 128

129 **METHODS**

Cultures of free-living microalgae and field sampling --- Monoclonal cultures of 130 131 free-living strains of symbiotic dinoflagellates Brandtodinium nutricula (RCC3468) (Probert et al. 2014), Gymnoxanthella radiolariae (RCC3507) (Yuasa et al. 2016), and Pelagodinium 132 beii (RCC1491) (Siano et al. 2010) have been previously obtained through single-cell 133 134 isolation from their radiolarian and foraminiferan hosts, respectively. Cultures of Phaeocystis strain RCC1383, found in association with symbiotic Acantharia (Decelle et al. 2012), but 135 originally isolated in its free-living stage, were chosen to represent the free-living symbiotic 136 algae of the ubiquitous Acantharia-Phaeocystis symbiotic association. All cultures were 137 maintained in 0.22 µm filter-sterilized (Stericup-GP, Millipore) seawater with K/2 (-Tris,-Si) 138 medium supplements (Keller et al. 1987) at 18° C, ~ 80 µmol photon m⁻² s⁻¹ light intensity and 139 14:10 light:dark cycle in the lab. Samples for DMSP analysis, cell counts and image-based 140 6

analysis of biovolume (see below) were taken for cultures at exponential and stationary phase,
during night and daytime (4 hours into each cycle; Supplementary information).

Field samples of symbiotic Radiolaria and Foraminifera were collected in coastal 143 144 waters of the Red Sea in Eilat (29°33'N, 34°57'E) and the western Mediterranean in Villefranche-sur-Mer (43°42'N, 7°18'E) during March and June of 2014, respectively. 145 Plankton community was sampled using plankton net with 220 µm mesh size towed obliquely 146 (0-30 m) for 10 minutes from a boat or swimming at surface (0-5 m) for approximately 200 147 m. Collected samples were immediately diluted in buckets with freshly collected surface 148 seawater, protected from direct sunlight, and transported to the lab within less than an hour. 149 Individual specimens were then manually sorted under a stereomicroscope using a 150 151 micropipette and transferred to Petri dishes in the case of single-celled acantharians and foraminiferans, and to larger beakers in the case of collodarians, where they were rinsed with 152 surface 0.22 µm filtered seawater (fsw) before following DMSP/image analysis or 153 experimental procedure. This single-cell approach allowed us to assess intracellular DMSP 154 concentration and isotopic composition (δ^{34} S-DMSP) in ecologically relevant, uncultured 155 specific symbiotic taxa. While large collodarians could be identified to genus level based on 156 morphological characteristics, identification was not so reliable for Acantharia, and so related 157 158 morphotypes of different species were used in subsequent measurements (Fig. 1). Samples manipulation and experimental work was conducted in the laboratory facilities of the 159 Interuniversity Institute for Marine Sciences in Eilat (Israel) and the Observatoire 160 161 Oceanologique de Villefranche-sur-Mer (France).

Image analysis and cellular biovolume assessment and microalgae cell counts --- Cultured and uncultured organisms were imaged using a digital camera (Canon EOS 5D)
 coupled to an optical direct microscope (Olympus BX51 and Nikon Eclipse) or stereoscope

(Zeiss Stereo discovery V200). Microalgae and rhizarian cell and colonial size dimensions
were assessed with the ImageJ open source image processing software

(https://imagej.nih.gov/ij/) and organismal biovolume derived from minimum and maximum 167 length dimensions and the formula of a prolate sphere, as described in Biard et al. (2016). For 168 single-celled amoeboid acantharians and Foraminifera with highly variable space occupation 169 of the cytoplasm the perimeter drawn to estimate the min/max lengths included the skeleton 170 axis regardless of the position of the cytoplasm; while for colonial collodarians, the perimeter 171 was given by the contour of the colony. Samples of cultured phytoplankton were fixed with 172 0.1 % glutaraldehyde (final concentration), for 15 min at room temperature in the dark, flash-173 freeze in liquid nitrogen and stored at -80 °C (Marie et al. 1997) until analysis. Cell 174 abundance was determined using a FACSAria (Becton Dickinson, San José, CA, USA) Flow 175 Cytometer and raw files analyzed with the FlowJo software (TreeStar Data Analysis 176 177 Software).

Dimethyl sulfur compounds concentration analysis --- DMSP was measured as the 178 DMS evolved by alkaline hydrolysis using purge and trap coupled to gas chromatography 179 (Shimadzu GC14A) with flame photometric detection (FPD) as described in Galí et al. 180 (2011). We therefore measured total DMSP (hereafter DMSPt) that comprises mainly 181 particulate DMSP+DMS in the case of individual symbiotic specimens, while DMSPt 182 measurements in cultured microalgae comprise both particulate and dissolved forms. For 183 cultured phytoplankton DMSP analysis, an aliquot of 0.5-1 mL from the culture was 184 dispensed into 13 mL gas-tight vials previously filled with MilliQ water. For single-celled 185 Radiolaria and Foraminifera, 2-4 specimens were transferred from the containers where they 186 had been rinsed with fsw, to a 13 mL vial previously filled with MilliQ. For colonial 187 Radiolaria, 1-2 specimens were transferred to the analytical vial, which was filled with fsw 188

instead, to minimize the potential contamination by DMSO in the MilliQ water system. We 189 then added two pellets of NaOH before sealing the vial with Teflon-capped lids. Samples 190 were stored at room temperature in the dark until analysis, within the next 2 months. To 191 192 account for dissolved DMSPt in solution, we preserved and analyzed blanks of the solutions where the organisms were analyzed (MilliQ for microalgae and single-celled 193 Radiolaria/Foraminifera, and fsw for colonial Radiolaria). Dissolved DMSP measured in the 194 blanks was then subtracted from total DMSP in the corresponding organism samples to yield 195 the particulate total DMSP values presented in this study. The contribution of dissolved 196 DMSP to the total DMSP was minor (<5%). 197

Samples were sparged with 40 mL min⁻¹ of high-purity helium, with the volatiles 198 trapped in a Teflon loop tube submersed in liquid nitrogen for 3-5 min before re-volatizing 199 them by placing the Teflon tube in hot water. Sulfur compounds were separated using a 200 packed Carbopack® 60/80 mesh column (Sigma-Aldrich) maintained at 170°C. Intracellular 201 content and cell concentration of cultured microalgae and freshly-collected Radiolaria or 202 203 Foraminifera holobionts were assessed from the concentration of DMSPt measured in the sample vial minus the corresponding blank, and normalized by the number and biovolume of 204 the specimens fixed in the vial. DMSP cell content in symbiotic microalgae within the 205 holobiont - in hospite - were calculated by dividing the DMSPt measured for the holobiont 206 by the mean number of host cells per colony surface, and also the mean number of microalgae 207 cells per host obtained from image analysis, assuming that the DMSP measured in the 208 holobiont was entirely confined in the symbionts. 209

210

211

212

Analysis of sulfur isotopic composition of sulfate and dimethyl sulfur compounds ---The samples for sulfur isotope analysis of DMSP were prepared similarly to the concentration measurements and therefore, they refer to DMSPt as well. For sulfur isotopic composition of

DMS, non-fixed samples were analyzed within 36 hours of collection (see below). Sulfur 213 isotopic composition of DMS and cellular DMSP were measured by purge and trap system 214 that was connected to a gas chromatograph (GC) coupled to a multicollector inductively 215 coupled plasma mass spectrometer (MC-ICPMS) (Amrani et al. 2009; Said-Ahmad and 216 Amrani 2013). Details for this method can be found in Said-Ahmad and Amrani (2013). 217 Briefly, seawater samples diluted in anoxic MilliQ or fsw were collected from the original 218 13ml vial using a syringe with minimal disturbance, and injected gently into a new 40 mL 219 220 sparging vial equipped with a Teflon septum. The vial was then sparged with He (40 mL/min) for 12 minutes. Water vapor was removed by a Nafion-membrane dryer (Perma pure LLC, 221 NJ, USA) using dry N₂ as the counter flow. A Teflon sample loop was inserted in a dewar of 222 liquid N₂ to trap DMS. After sparging, the 6-port valve (Valco Instrument Co, TX, USA; 223 heated to 80°C) was turned to the inject position, and the sample loop transferred quickly 224 225 from the liquid N₂ to hot water so that the trapped gases were injected into a Agilent J&W capillary column (DB-1, 60m x 0.32mm ID x 1.0µm), connected directly to the 6-port valve. 226 227 At the same time the GC (Perkin Elmer 580) and the MC-ICPMS (Neptune Plus, 228 ThermoFischer Scientific) were started. A standard DMS sample was introduced to the system for calibration every 3-4 samples and we used a bracketing technique to correct for 229 instrumental mass bias and calibration of the SF6 internal standard (Said-Ahmad and Amrani 230 2013). The results are expressed in conventional δ^{34} S notation as a per mil (‰) deviation from 231 the international standard V-CDT (Vienna Canyon Diablo Troilite) according to the equation 232 below. 233

 $\partial \mathcal{S} = \left(R^{4} \right)^{3} R^{4}$

where ³⁴R is the integrated ³⁴S/³²S ion-current ratio of the sample and standard peaks.
Analytical precision of analysis of DMS and DMSP standards was usually in the range of 0.1-10

0.4% (1 σ standard deviation). The precision of sulfur isotope analysis for duplicate or triplicate samples of seawater and organism samples were usually less than 1‰. Accuracy as calculated by standards was in the range of 0.2 ‰ (Said-Ahmad and Amrani 2013) and estimated to be better than 1‰ for the seawater samples.

Dissolved sulfate (SO_4^{2-}) sulfur isotope analysis was performed by a conventional 241 elemental analyzer (EA) coupled to isotope ratio mass spectrometer (IRMS) method 242 (Giesemann et al. 1994) using Delta Plus (Thermo) IRMS. Samples of BaSO₄ were prepared 243 from diluted seawater samples by addition of 5% BaCl₂ solution and then analyzed for their 244 δ^{34} S values by EA-IRMS. The sulfur isotope reference materials NBS-127 (BaSO₄; δ^{34} S = 245 21.1‰), IAEA-S-1 (Ag₂S; -0.3‰), and IAEA-SO-6 (BaSO₄; -34.1‰) were purchased from 246 the National Institute of Standards and Technology (NIST) and used for calibration. Precision 247 of this method for duplicates/triplicates was usually better than 0.3%. 248

Sulfate assimilation and DMSP synthesis experiment --- For the biological sulfate 249 assimilation experiment we incubated two cultured strains of microalgae 250 251 (Phaeocystis RCC1383 and B.nutricula RCC3468) and freshly-collected specimens of 252 solitary symbiotic *Thalasicolla* sp. (Collodaria-*Brandtodinium* holobiont) during three days in two different types of K/2 culture media prepared with isotopically distinct sulfate (Fig. 2a). 253 254 One media ('Heavy') was prepared with filtered seawater amended with standard K/2 supplements to yield an average seawater sulfate δ^{34} S of 21.5 ± 0.5‰ at the beginning of the 255 incubations. The second media ('Light') consisted of artificial seawater (Berges et al. 2001) 256 prepared using isotopically depleted δ^{34} S-Sulfate relative to seawater sulfate, and mixed in 1:1 257 ratio with K/2 standard media, as described above, to give a sulfate δ^{34} S value of 7.6 ± 0.4‰. 258 This procedure resulted in two 2-L batches of media with very distinct δ^{34} S-Sulfate 259 composition but virtually identical nutrient concentrations. Experimental incubations for 260 11

each microalgae species were prepared by inoculating a 50 mL aliquot from a culture that had 261 been maintained in K/2-based standard media for at least 50 generations, into 1 L of 'Heavy' 262 and 'Light' media. After gentle homogenization initial samples (T_0) were taken from the two 263 cultures and the remaining volume was subsequently aliquoted into 8 replicated 70 mL sterile 264 tissue culture vessels (Fig. 2a). For incubations of Radiolaria holobionts freshly-collected 265 specimens, collected and maintained in 0.2 um fsw since the previous day, were transferred (2 266 267 ind/vessel) into 8 replicated tissue culture vessels previously filled with 'Heavy' or 'Light' media. Experimental design included therefore, three organismal types - two symbiotic 268 microalgae strains in free-living stage, and one Radiolaria holobiont - each distributed on a 269 270 series of replicated 8 vessels, half of which had been filled with 'Heavy' or 'Light' media, 271 respectively (Fig. 2a). Organisms were incubated in parallel under constant temperature (19-20°C) and light (200 μ mol photon m⁻² s⁻¹, 14:10 light:dark cycle) conditions in the lab. 272 Replicated incubations from each organism and media type were sampled at time 0 and after 273 274 22, 46 and 70 hours of incubations for analysis of DMSPt concentration and sulfur isotopic composition. 275

DMSP \rightarrow **DMS cleavage fractionation experiment** --- To assess potential differences 276 in the isotopic signature of the DMS generated by symbiotic Radiolaria from that of the 277 microbial community we determined the sulfur isotope composition of DMS resulting from 278 the cleavage of DMSP produced during experimental incubations containing different 279 280 planktonic biota (Fig. 2b). This was achieved by incubating a) the natural microbial assemblage $< 200 \,\mu\text{m}$ (*i.e.* surface whole sea water, (*wsw*)), b) a heterogeneous assemblage of 281 Acantharia-microalgae holobionts (100 ind.) in 0.2 µm-filtered seawater obtained with sterile 282 Stericup filtration device (Stericup-GP, Millipore) (fsw+rads), and a third control treatment 283 containing only the same filtered seawater (fsw-only) (Fig. 2b). Organisms were manipulated 284

using sterile micropipette and petri dishes on the laboratory bench and incubations were 285 prepared in 70 mL sterile polystyrene tissue culture vessels. Triplicates of each assemblage 286 type were incubated for 4 hours (13:30-17:30) under ~30% incident light. Samples from wsw 287 for DMSP were preserved with NaOH in gas-tight vials, while those for DMS were 288 maintained in the dark at similar temperature to that of the surface water, until analysis. In the 289 "fsw+rads" treatments, 5 individuals from each incubation were sorted, imaged and preserved 290 for DMSP concentration. The remaining ~95 specimens were transferred into 40 mL gas-tight 291 292 vials with the same fsw where they were incubated and kept at room temperature until analysis. 293

Statistical analysis --- The statistical significance of the difference between organism
groups in DMSP cell content, DMSP and DMS sulfur isotopic composition was tested with
one-way ANOVA followed by Tukey's post hoc test. All statistical analyses were conducted
on experimental results using GraphPad 5.0 software (GraphPad Software, Inc, La Jolla,
USA).

299 **RESULTS**

Intracellular DMSPt in cultured phytoplankton and freshly collected photosymbiotic 300 organisms --- Table 1 summarizes cell size, intracellular DMSPt content and concentration in 301 cultures of free-living phytoplankton and field-collected symbiotic Radiolaria and 302 Foraminifera holobionts (Fig. 1). The highest values were observed in the large Collodaria-303 304 *Brandtodinium* holobiont with average DMSPt cellular content ranging from 2757 ± 750 to 3652 ± 2008 pmol specimen⁻¹ (Table 1). Colonial forms contained hundreds of radiolarian 305 cells (i.e., estimated by the number of central capsules forming the colonies) per specimen 306 $(N_{Sphaerozoum} = 166 \pm 65; N_{Collozoum} = 509 \pm 72 \text{ central capsules colony}^{-1})$. Each central capsule 307 harbored a few endosymbiont cells (N_{Collozoum} = 9.8 ± 3.5 , N_{Sphaerozoum} = 9.1 ± 2.6 symbionts 308 13

309	central capsule ⁻¹). Solitary forms, consisting of millimeters-size single host cell, contained
310	hundreds of endosymbiotic cells ($N_{Thalassicolla} = 723 \pm 703$ symbionts radiolarian cell ⁻¹) (Fig. 1,
311	Table 1). DMSPt content in different morphotypes of single-celled Acantharia and planktonic
312	Foraminifera were much lower, according to their lower cell size, and varied within a
313	relatively narrow range (15-38 pmol cell ⁻¹). For a minifera (37.8 ± 15.5 pmol cell ⁻¹) and 'Star'
314	morphotype acantharians $(36.4 \pm 10.4 \text{ pmol cell}^{-1})$ exhibited higher values than A. elongata
315	$(15.2 \pm 5.4 \text{ pmol cell}^{-1})$ and 'Translucid' acantharians $(20.9 \pm 3.6 \text{ pmol cell}^{-1})$, yet differences
316	were only significant between A. elongata and Globigerinella sp. ($p=0.03$, $F_{3.12}=4.1$, one-way
317	ANOVA, Tukey's post hoc test). Analysis of DMSP concentration per cellular biovolume
318	yielded a different picture with A. elongata $(17.1 \pm 6.1 \text{ mmol } \text{L}^{-1})$ having significantly higher
319	concentrations than Foraminifera (7.5 \pm 3.1 mmol L ⁻¹), 'Star' (2.4 \pm 0.7 mmol L ⁻¹) and
320	'Translucid' $(0.3 \pm 0.1 \text{ mmol } L^{-1})$ acantharian morphotypes, while both solitary and colonial
321	collodarian species showed significantly lower concentrations ($p < 0.001$, $F_{6,16} = 14.8$) (Table 1).
322	Cellular DMSPt content in cultures of free-living algae was markedly lower
322	Cellular DMSPt content in cultures of free-living algae was markedly lower
322 323	Cellular DMSPt content in cultures of free-living algae was markedly lower (range=0.01-1.24 pmol cell ⁻¹) according to their smaller size compared to their hosts (Table
322 323 324	Cellular DMSPt content in cultures of free-living algae was markedly lower (range=0.01-1.24 pmol cell ⁻¹) according to their smaller size compared to their hosts (Table 1). Highest mean values were measured in <i>P. beii</i> (0.9 ± 0.2 pmol cell ⁻¹); these were 3-4-fold
322 323 324 325	Cellular DMSPt content in cultures of free-living algae was markedly lower (range=0.01-1.24 pmol cell ⁻¹) according to their smaller size compared to their hosts (Table 1). Highest mean values were measured in <i>P. beii</i> (0.9 ± 0.2 pmol cell ⁻¹); these were 3-4-fold higher than in <i>B. nutricula</i> (0.2 ± 0.1 pmol cell ⁻¹) while <i>G. radiolariae</i> exhibited intermediate
322 323 324 325 326	Cellular DMSPt content in cultures of free-living algae was markedly lower (range=0.01-1.24 pmol cell ⁻¹) according to their smaller size compared to their hosts (Table 1). Highest mean values were measured in <i>P. beii</i> (0.9 ± 0.2 pmol cell ⁻¹); these were 3-4-fold higher than in <i>B. nutricula</i> (0.2 ± 0.1 pmol cell ⁻¹) while <i>G. radiolariae</i> exhibited intermediate average cell DMSPt content (0.1 ± 0.1 pmol cell ⁻¹). DMSPt concentration calculated from
322 323 324 325 326 327	Cellular DMSPt content in cultures of free-living algae was markedly lower (range=0.01-1.24 pmol cell ⁻¹) according to their smaller size compared to their hosts (Table 1). Highest mean values were measured in <i>P. beii</i> (0.9 ± 0.2 pmol cell ⁻¹); these were 3-4-fold higher than in <i>B. nutricula</i> (0.2 ± 0.1 pmol cell ⁻¹) while <i>G. radiolariae</i> exhibited intermediate average cell DMSPt content (0.1 ± 0.1 pmol cell ⁻¹). DMSPt concentration calculated from microscope-based estimates of cellular biovolume in free-living algae was one-to-three orders
322 323 324 325 326 327 328	Cellular DMSPt content in cultures of free-living algae was markedly lower (range=0.01-1.24 pmol cell ⁻¹) according to their smaller size compared to their hosts (Table 1). Highest mean values were measured in <i>P. beii</i> (0.9 ± 0.2 pmol cell ⁻¹); these were 3-4-fold higher than in <i>B. nutricula</i> (0.2 ± 0.1 pmol cell ⁻¹) while <i>G. radiolariae</i> exhibited intermediate average cell DMSPt content (0.1 ± 0.1 pmol cell ⁻¹). DMSPt concentration calculated from microscope-based estimates of cellular biovolume in free-living algae was one-to-three orders of magnitude higher than concentration in the radiolarian and foraminiferan holobionts (Table
322 323 324 325 326 327 328 329	Cellular DMSPt content in cultures of free-living algae was markedly lower (range=0.01-1.24 pmol cell ⁻¹) according to their smaller size compared to their hosts (Table 1). Highest mean values were measured in <i>P. beii</i> (0.9 ± 0.2 pmol cell ⁻¹); these were 3-4-fold higher than in <i>B. nutricula</i> (0.2 ± 0.1 pmol cell ⁻¹) while <i>G. radiolariae</i> exhibited intermediate average cell DMSPt content (0.1 ± 0.1 pmol cell ⁻¹). DMSPt concentration calculated from microscope-based estimates of cellular biovolume in free-living algae was one-to-three orders of magnitude higher than concentration in the radiolariae (462 ± 168 mmol L ⁻¹) showed
 322 323 324 325 326 327 328 329 330 	Cellular DMSPt content in cultures of free-living algae was markedly lower (range=0.01-1.24 pmol cell ⁻¹) according to their smaller size compared to their hosts (Table 1). Highest mean values were measured in <i>P. beii</i> (0.9 ± 0.2 pmol cell ⁻¹); these were 3-4-fold higher than in <i>B. nutricula</i> (0.2 ± 0.1 pmol cell ⁻¹) while <i>G. radiolariae</i> exhibited intermediate average cell DMSPt content (0.1 ± 0.1 pmol cell ⁻¹). DMSPt concentration calculated from microscope-based estimates of cellular biovolume in free-living algae was one-to-three orders of magnitude higher than concentration in the radiolariae (462 ± 168 mmol L ⁻¹) showed higher values than <i>Phaeocystis</i> RCC1383 (307 ± 47 mmol L ⁻¹) and <i>P. beii</i> (272 ± 49 mmol L ⁻

changes in the DMSPt cell content and concentration in relation to growth phase andphotoperiod at the time of harvesting the culture (Supporting information).

DMSP sulfur isotopic composition (δ^{34} S-DMSP) --- The mean δ^{34} S-DMSP in isolated 335 holobiont specimens of Acantharia-*Phaeocystis* sp. (δ^{34} S-DMSP = 19.8 ± 0.4‰) and 336 For a minifera-*P. beii* (δ^{34} S-DMSP = 20.1 ± 0.4‰) were very similar to each other, and also to 337 the natural microbial assemblage coexisting in the same surface water (δ^{34} S-DMSP = 19.7 ± 338 0.4‰, Fig. 3). These values were slightly lower than δ^{34} S of local seawater sulfate (δ^{34} S-SO₄²⁻ 339 =21.5 \pm 0.5%). In contrast, the Collodaria-*Brandtodinium* holobiont showed a very distinct 340 DMSP isotopic composition (δ^{34} S-DMSP = 23.5±0.8‰), significantly enriched not only 341 compared to other symbiotic groups (Acantharia and Foraminifera) and microbial assemblage, 342 but also compared to δ^{34} S-SO₄²⁻ in seawater (*p*<0.0001, F_{7.26}=46.9, one-way ANOVA, Fig. 343 3). 344

Sulfur isotopic composition for cultured microalgae showed differences among species, with *Brandtodinium nutricula* (18.2 ± 0.3‰) showing significantly lower δ^{34} S-DMSP values compared to *Phaeocystis* RCC1383 (20.2 ± 0.4‰, Tukey comparison test, *p*<0.001, Fig. 3). In the case of *B. nutricula* the δ^{34} S-DMSP values were significantly depleted relative to the Collodaria-*B. nutricula* holobiont (Tukey comparison test, *p*<0.0001), whereas the δ^{34} S-DMSP values of free-living *Phaeocystis* RCC1383 was not significantly different from that measured in the Acantharia-*Phaeocystis* holobionts (Fig. 3).

352 Sulfate assimilation into DMSP and δ^{34} S fractionation --- Free-living microalgae 353 growing in the 'Light' medium showed a progressive ³⁴S depletion of DMSP both with time 354 and relative to microalgae growing in the 'Heavy' medium, indicating active sulfate 355 assimilation from seawater for DMSP biosynthesis (Fig. 4a, b). Although relatively constant 356 during the 3-day incubation, the δ^{34} S-DMSP of the microalgae growing in the 'Heavy' media 35 was depleted relative to the seawater sulfate. The depletion was more pronounced for *B. nutricula* (-3‰) than for *Phaeocystis* RCC1383 (-1‰) (Fig. 4a, b). Contrary to free-living microalgae, δ^{34} S-DMSP in symbiotic Radiolaria incubated in 'Light' and 'Heavy' media remained similar and substantially ³⁴S-enriched relative to available sulfate sources during the incubation (Fig. 4c).

DMSP to **DMS** conversion and sulfur isotope fractionation --- δ^{34} S of DMSP and 362 derived DMS measured after daylight incubations showed differences between freshly-363 collected microbial assemblages and photosymbiotic radiolarians (Fig. 5). Sulfur isotopic 364 composition of DMS measured from *wsw*, which contained the bulk microbial assemblage 365 $(19.7 \pm 0.3 \%)$, was not significantly different from that measured in DMSP $(19.4 \pm 0.1 \%)$. 366 Fig. 5). Conversely, δ^{34} S-DMS measured from incubations containing photosymbiotic 367 Radiolaria (Acantharia-*Phaeocystis* holobiont, *fsw+rads*) was significantly lower (18.4 ± 0.4 368 %; p < 0.0001, F_{5.14}=12, one-way ANOVA). Specifically, DMS- δ^{34} S values measured in the 369 incubations with Radiolaria were significantly depleted relative to δ^{34} S-DMSPt values in both 370 A. elongata (1.2 ‰ mean difference) and 'Star' radiolarian morphotypes (1.5 ‰ mean 371 difference). Similarly, δ^{34} S-DMS in *fsw-only* treatment incubated in parallel (18.0 ± 0.5 ‰) 372 was depleted relative to δ^{34} S-DMSP in microbial assemblage and Radiolaria holobionts (Fig. 373 374 5).

375 DISCUSSION

376 DMSP cellular content in photosymbiotic plankton and partition among partners ---

The role of DMSP in coral symbiosis has received considerable attention in recent years
revealing high cell concentration and content of DMSP in the holobiont and different *Symbiodinium* sp. strains (Broadbent et al. 2002; Steinke et al. 2011; Deschaseaux et al. 2014)
and DMS production associated to coral reefs (Broadbent and Jones 2004; Raina et al. 2010; 16

Exton et al. 2015; Frade et al. 2016). One of the objectives of this study was to determine 381 whether photosymbiosis in planktonic organisms is systematically associated with high 382 concentrations of cellular DMSP. The high concentration and cellular content of DMSP 383 measured in symbiotic microalgal species cultured in their free-living stage but originally 384 isolated from Radiolaria and Foraminifera hosts (Table 1) is consistent with this hypothesis. 385 Highest DMSPt concentrations were observed among dinoflagellate strains (Table 1), which 386 are within the upper range of previously reported values for this phytoplankton class (Caruana 387 and Malin 2014). Although elevated DMSP concentrations are characteristic of Dinophyceae 388 and Prymnesiophyceae (Keller et al. 1989), the extremely high concentrations associated with 389 390 the symbiotic partners of Collodaria and planktonic Foraminifera, is suggestive of DMSP being an important attribute of the algal partner in rhizarian symbiotic associations. 391

The endosymbiotic nature of the Radiolarian-microalgae associations precludes direct 392 measurements of the DMSP content in symbiotic microalgae cells within the radiolarian host 393 (i.e. in hospite). However, the DMSP concentrations estimated for microalgae, based on the 394 DMSP content in the holobiont and the number of microalgae cells hosted, yielded extremely 395 high concentrations (Table 1) that matched the highest values estimated for photosymbiotic 396 anemone and coral species (Broadbent et al. 2002; Van Alstyne et al. 2006). In the case of the 397 colonial and solitary Radiolaria-microalgae holobiont, DMSP content per cell estimated for B. 398 nutricula in hospite was 3-, 8-fold and 20-fold higher than for free-living cells, respectively. 399 This difference was even larger for *Phaeocystis* sp. with 64-100-fold higher cellular content 400 estimated for symbiotic algae in 'A. elongata' and 'Translucid' morphotypes of Acantharia, 401 respectively, compared to the free-living algae, which would result in unrealistic (40-60 mM 402 DMSP) cellular concentrations. In other words, if the 15-38 pmol DMSP cell⁻¹ measured in 403 the Acantharia-Phaeocystis sp. partnership (Table 1) was allocated entirely to symbiotic algae 404

with similar concentrations to those measured in free-living cultures, the host should harbor 405 1250-3000 algal cells, which is a 100-fold more than the 10-20 cells commonly found in its 406 cytoplasm. Acknowledging the limits of our data, these calculations suggest that DMSP 407 could be present in the host as well. Whether the DMSP is translocated from the algae to the 408 host (Van Alstyne et al. 2009) and/or produced by the latter (Raina et al. 2013) cannot be 409 concluded from concentration measurements alone. Regardless of its origin, the moderate-to-410 high cellular content of DMSP estimated for the host and the algae are consistent with recent 411 observations in Radiolaria (Decelle et al. 2012) and other photosymbiotic organisms (Hill et 412 al. 2000; Stefels 2000; Broadbent et al. 2002; Van Alstyne et al. 2006), and suggest that both 413 partners could mutually benefit from one or more of the multiple eco-physiological roles 414 attributed to dimethyl sulfur compounds, e.g., osmoregulation, oxidant scavenger or info-415 chemical (Stefels 2000; Seymour et al. 2010; Raina et al. 2013). 416

417 Sulfate assimilation and DMSP production in photosymbiotic associations – Highly

enriched δ^{34} S-DMSP observed in Collodaria-*Brandtodinium* sp. holobiont specimens (23.5 ± 418 0.8 ‰; Fig. 3) was particularly unexpected. This association showed significantly enriched 419 δ^{34} S-DMSP not only relative to seawater SO₄²⁻ but also compared to other symbiotic 420 radiolarians and all measurements previously reported for both macro- and microalgae, which 421 showed a slight ³⁴S-depletion in DMSP compared to seawater SO_4^{2-} (Oduro et al. 2012; 422 Amrani et al. 2013; Said-Ahmad and Amrani 2013)(Fig. 3). This distinctive sulfur isotopic 423 composition could result from differential fractionation associated with the multi-step 424 assimilation of sulfate into methionine (intracellular precursor to DMSP) and/or may reflect 425 differences in the DMSP biosynthetic pathway. The 1-3‰ ³⁴S depletion in DMSP relative to 426 sulfate during assimilation of sulfate is in agreement with previous reports (e.g. Kaplan & 427 Rittenberg 1964). All three biosynthetic pathways for DMSP described to date for higher 428

plants and algae rely on assimilatory sulfate reduction (Gage et al. 1997; Summers et al. 429 1998). However, our incubation experiments with isotopically distinct sulfate solutions 430 showed that while free-living microalgae did actively take up and assimilate sulfate into 431 DMSP in time scales relevant for phytoplankton growth (Fig. 4a, b), collodarians with 432 elevated cellular content of DMSP showed no isotopic evidence of assimilatory sulfate 433 reduction into DMSP (Fig. 4c). This could be due to slower growth rates of these giant 434 protists compared to microalgae, although the high photosynthetic rates of solitary Radiolaria 435 (Caron et al. 1995) and abundance of endosymbiotic microalgae in incubated specimens 436 (Table 1, Fig.1) could suggest additional explanations to the lack of sulfate uptake. The 437 holobiont may for instance rely on inorganic or reduced sulfur species (e.g. methionine, 438 cysteine) stored by the algae for DMSP synthesis. Radiolarians are active grazers (Anderson 439 1978; Swanberg and Anderson 1985; Suzuki and Not 2015), and captured prey and digestion 440 441 products could provide such sulfur compounds to the symbiotic microalgae, reducing its dependence on sulfate uptake. It is worth noting however, that the assimilatory sulfate 442 443 reduction only represents a small proportion of the total energetic cost (NADPH and ATP 444 molecules) associated with *de novo* synthesis of DMSP (Lavoie et al. 2016).

In contrast to the collodarians, the δ^{34} S-DMSP measured in individual specimens of 445 Acantharia-*Phaeocystis* (19.8 \pm 0.4‰) and *Globigerinella* sp.-*P. beii* (20.1 \pm 0.4‰), was very 446 similar to the isotopic composition obtained from simultaneous $(19.7 \pm 0.4\%)$ and previously 447 reported bulk δ^{34} S-DMSP measurements of surface microbial assemblages (19.7 ± 0.5%), 448 Amrani et al. 2013) (Fig. 3). In this regard, the δ^{34} S-DMSP of cultured *Phaeocystis* RCC1383 449 was similar to that measured in the Acantharia-Phaeocystis sp. holobiont (Fig. 3). The 450 reasons for the different sulfur isotopic behavior between Acantharia-Phaeocystis sp. and 451 Collodaria-B. nutricula associations is unclear to us, but given the phylogenetically distant 452

microalgal partners involved in these symbiotic association (*i.e.* Dinophyceae and 453 Prymnesiophyceae), this isotopic variability could reflect differences in the metabolic 454 capacities of the symbiotic algae and/or the holobiont. Differences in δ^{34} S-DMSP observed 455 between B. nutricula and Phaeocystis RCC1383 (Fig. 3) were of similar magnitude to those 456 reported between macro and microalgae species (Oduro et al. 2012). The observed differences 457 between the two algal strains and the remarkably distinct δ^{34} S-DMSP observed in freshly 458 collected Collodaria-B. nutricula associations (Fig. 3) contrast with the homogeneity in δ^{34} S-459 DMSP composition observed across diverse marine microbial communities (Amrani et al. 460 2013) and illustrate the isotopic variability potentially hidden in natural microbial 461 communities. Little is known about the isotopic variability associated with taxonomic and 462 functional diversity of major DMSP producers. Nonetheless, if adequately characterized, this 463 variability offers an opportunity not only to investigate physiological differences between 464 465 species in the lab, but also to refine our current ability to track specific sources of DMSP, and derived DMS, in highly intermixed and diverse natural microbial communities (Kaye 1987; 466 Calhoun et al. 1991; Said-Ahmad and Amrani 2013). 467

Sulfur isotope fractionation associated with DMSP cleavage into DMS --- In addition 468 to the sulfur isotopic composition of DMSP, it is important to constrain the isotopic 469 fractionation associated with the cleavage of DMSP into DMS. The two previous studies 470 assessing this fractionation reported contradictory patterns; Oduro et al. (2012) reported ³⁴S 471 depletion for the DMS produced by macroalgae, while Amrani et al. (2013) reported small to 472 negligible ³⁴S enrichment from parallel measurements of DMSP and DMS *in situ*. Our results 473 from the experimental incubations conducted with natural microbial assemblages and 474 symbiotic radiolarians (Fig. 5), although limited in data and scope, offer new insights that 475 may contribute to reconcile contrasting previous observations. It is important to note that we 476

are not calculating fractionation factors (α), but rather apparent fractionation (Δ^{34} S) from the 477 difference between the product and the reactant. Consistent with Amrani et al. (2013), δ^{34} S-478 DMSP and δ^{34} S-DMS values were similar in whole seawater incubated from the Red Sea. 479 suggesting negligible fractionation is associated with microbial cleavage of DMSP. However, 480 the δ^{34} S value of DMS produced by symbiotic acantharians was lower (-1.2, -1.5‰) than that 481 of DMSP (Fig. 5), resulting in a DMS fractionation similar in sign and magnitude to that 482 reported for macroalgae by Oduro et al. (2012). These differences in the DMSP to DMS 483 484 fractionation between the microbial assemblage, symbiotic radiolarians and macroalgae may reflect their taxonomic and functional diversity. However, similar ³⁴S depletion of DMS was 485 observed between *fsw-only* and *fsw+rads* incubations (Fig. 5). Evidences exist that the 486 filtration process can break phytoplankton cells and enrich the filtered seawater with 487 dissolved DMSP and extracellular activity of DMSPlyase (Kiene and Slezak 2006) to the 488 489 point that it is common to measure substantial DMS production in fsw (Galí et al. 2011). During the 4-hour experiments, DMS was produced in both *fsw-only* and *fsw+rads* 490 incubations, although, the latter, containing approximately 100 specimens of Acantharia (see 491 methods), produced larger amounts of DMS (Fig. 5). Overall, δ^{34} S values of DMS produced 492 in natural microbial communities were similar or slightly higher than δ^{34} S values of DMSP 493 (Amrani et al., 2013; this study), while cultures or incubations with isolated organisms in 494 filtered seawater produced ³⁴S-depleted DMS (Oduro et al., 2012; this study). This may 495 reflect the interplay among DMS sources and sinks: DMSPlyase selection for DMSP with 496 lower δ^{34} S would produce ³⁴S depleted DMS, but preference of DMS-consumption processes 497 (e.g. bacterial uptake, photochemistry, ventilation, and bacterial uptake) for ³⁴S-depleted DMS 498 would buffer the overall result. This possibility was already discussed by Amrani et al. 499 (2013), who invoked these DMS removal processes as potential mechanisms contributing to 500

're-enrich' the DMS pool. To further constrain the contribution of each physical and 501 biological process to this putative isotopic compensatory effect, we can compare their 502 prevalence in the different incubations and natural measurements available from this and 503 previous studies. The effect of ventilation can be ruled out because our incubation bottles did 504 not allow for gas-exchange; also, the fact that the fractionation buffering effect had been 505 observed at depths were ventilation is low further underscores the minor influence of 506 ventilation (Amrani et al. 2013). The similarity in the light conditions for the three incubation 507 508 types (*wsw*, *fsw-only*, and *fsw+rads*), seem to argue against the effect of photochemistry upon DMS fractionation as well. Finally, although not directly measured, bacterial abundance, and 509 likely bacterial DMS consumption, should have been largely suppressed in 0.2 um-filtered 510 seawater during the short duration (4-hours) incubations. Filtered seawater showed ³⁴S-511 depletion of DMS relative to DMSP (Oduro et al. 2012; *fsw-only* incubation this study) while 512 whole seawater, containing natural abundances of bacteria, showed similar δ^{34} S values for 513 DMS and DMSP (Amrani et al. 2013; wsw incubation this study, Fig. 2b). These observations 514 515 are not sufficient to confirm, but are consistent with, a hypothetical scenario where bacterial uptake contributes the most to the homogenization of δ^{34} S DMS(P) in natural settings. Future 516 experiments assessing 1) the fractionation towards ³⁴S depleted DMS preference of DMSP 517 lyases, and 2) the magnitude and sign of the sulfur isotopic fractionation associated with 518 bacterial uptake are needed to test this hypothesis. 519

Photosymbiosis contribution to the marine sulfur cycle --- Current understanding of
DMS(P) cycling in pelagic ecosystems recognizes phytoplankton, particularly the nano-size
fraction (2-20 μm) with abundant representatives of high DMSP producer classes such as
Prymnesiophyceae and Dinophyceae, and also diatoms (Bucciarelli et al. 2013), as the
primary DMSP producers (Keller et al. 1989; Stefels et al. 2007). However, the concentration

of DMSP observed in symbiotic Radiolaria and Foraminifera, suggests that larger 525 526 photosymbiotic plankton (50-2000 µm) that harbor photosynthetic endosymbionts from these same phytoplankton groups (Siano et al. 2010; Decelle et al. 2012; Probert et al. 2014; Yuasa 527 et al. 2016), can represent a significant source of marine DMS(P) particularly in (sub)tropical 528 latitudes where high abundances have been reported (Caron et al. 1995; Dennett et al. 2002; 529 Biard et al. 2016). To evaluate this hypothesis we calculated the Radiolaria-associated 530 contribution to total water column DMSP in the euphotic zone of a number of stations from 531 the (sub)tropical-oriented *Tara* Oceans expedition track, using the mean group-specific 532 DMSP concentration for Radiolaria (Table 1) and in situ abundance and biovolume obtained 533 534 with the Underwater Video Profiler (UVP)(Biard et al. 2016), and micro- and nano-535 phytoplankton-associated DMSP estimated from chlorophyll-based empirical relationships (Belviso et al. 2004). The potential contribution of large symbiotic Radiolaria (UVP lower 536 detection threshold = 600 μ m) to water column DMSP was low on average (1.1 ± 2.2%, mean 537 \pm SD, n=19) although peaks of moderate contribution (8%) were inferred associated with 538 539 Acantharia. Despite being inefficiently captured by the UVP – Acantharia are numerically dominated by <150 µm forms (Michaels 1988; Caron and Swanberg 1990) – these group of 540 Radiolaria potentially contribute the most DMSP in virtue of their high DMSP intracellular 541 concentration. Moreover, the Radiolaria-associated DMSPt may represent an important 542 543 source of sulfur for higher trophic levels that cannot effectively prev on smaller nanophytoplankton. Considering all these caveats, this conservative approximation indicates 544 that photosymbiotic plankton can represent a significant source of DMS(P), particularly in 545 tropical and subtropical oceans, where the high DMSP producers that are typical of temperate 546 and sub-polar latitudes (e.g. E. huxleyi or Phaeocystis sp.), are less abundant and Radiolaria 547 and Foraminifera represent an important component of plankton communities (Caron et al. 548

1995; Decelle et al. 2015; Biard et al. 2016), and can account for up to 20% of primary 549 production (Michaels 1988). In this regard, the distinct sulfur isotopic composition of DMSP 550 in Collodaria-Brandtodinium holobiont, one of the most abundant and ecologically relevant 551 group of photosymbiotic plankton (Dennett et al. 2002; de Vargas et al. 2015; Guidi et al. 552 2016), presents the possibility of assessing the contribution of photosymbiotic plankton to 553 community DMSP production in the future. Beyond biogeochemical considerations, the 554 highly ³⁴S enriched DMSP and the lack of apparent sulfate assimilation into DMSP observed 555 556 in Collodaria-Brandtodinium associations suggest that DMSP biosynthetic pathways could be linked to the recycling of organic sulfur between symbiotic partners. If use of organic sulfur 557 for DMSP synthesis also occurs among free-living phytoplankton, we may be able to 558 discriminate between 'new' (from sulfate) vs 'recycled' (from organic S) DMSP production 559 by their differential S-isotopic signature, in a similar way to approaches using nitrogen 560 561 isotopes to apportion new and recycled nitrogen sources for primary production.

563 **References**

- Amrani, A., W. Said-Ahmad, Y. Shaked, and R. P. Kiene. 2013. Sulfur isotope homogeneity
 of oceanic DMSP and DMS. Proc. Natl. Acad. Sci. U S A 110: 18413-18418.
- Amrani, A., A. L. Sessions, and J. F. Adkins. 2009. Compound-specific delta s-34 analysis of
- volatile organics by coupled GC/multicollector-ICPMS. Anal. Chem. **81:** 9027-9034.
- 568 Anderson, O. R. 1978. Light and electron-microscopic observations of feeding-behavior,
- nutrition, and reproduction in laboratory cultures of thalassicolla-nucleata. Tissue &
 Cell 10: 401-412.
- 571 ---. 1983. Radiolaria. Springer.
- Andreae, M. O. 1997. Atmospheric aerosols: Biogeochemical sources and role in atmospheric
 chemistry. Science 276: 1052-1058.
- Bates, T. S., B. K. Lamb, A. Guenther, J. Dignon, and R. E. Stoiber. 1992. Sulfur emissions to
 the atmosphere from natural sources. J. Atmos. Chem. 14: 315-337.
- 576 Belviso, S., C. Moulin, L. Bopp, and J. Stefels. 2004. Assessment of a global climatology of
- 577 oceanic dimethylsulfide (DMS) concentrations based on seawifs imagery (1998-2001).
 578 Can. J. Fish. Aquat. Sci. 61: 804-816.
- 579 Berges, J. A., D. J. Franklin, and P. J. Harrison. 2001. Evolution of an artificial seawater
- 580 medium: Improvements in enriched seawater, artificial water over the last two
- 581 decades. J. Phycol. **37:** 1138-1145.
- Biard, T., and others. Global in situ imaging observations reveal the biomass of rhizaria in the
 oceans. DOI 10.1038/nature17652
- Blunden, G., B. E. Smith, M. W. Irons, M. H. Yang, O. G. Roch, and A. V. Patel. 1992.

585 Betaines and tertiary sulfonium compounds from 62 species of marine-algae.

586 Biochem. Syst. Ecol. **20:** 373-388.

- Broadbent, A. D., and G. B. Jones. 2004. DMS and DMSP in mucus ropes, coral mucus,
 surface films and sediment pore waters from coral reefs in the great barrier reef. Mar.
 Freshwater Res. 55: 849-855.
- Broadbent, A. D., G. B. Jones, and R. J. Jones. 2002. DMSP in corals and benthic algae from
 the great barrier reef. Estuar. Coast. Shelf Sci. 55: 547-555.
- 592 Bucciarelli, E., C. Ridame, W. G. Sunda, C. Dimier-Hugueney, M. Cheize, and S. Belviso.
- 593 2013. Increased intracellular concentrations of DMSP and DMSO in iron-limited
 594 oceanic phytoplankton thalassiosira oceanica and trichodesmium erythraeum. Limnol.
 595 Oceanogr. 58: 1667-1679.
- Calhoun, J. A., T. S. Bates, and R. J. Charlson. 1991. Sulfur isotope measurements of
 submicrometer sulfate aerosol-particles over the pacific-ocean. Geophys. Res. Lett.
 18: 1877-1880.
- 599 Caron, D. A., P. D. Countway, A. C. Jones, D. Y. Kim, and A. Schnetzer. 2012. Marine
- protistan diversity, p. 467-493. *In* C. A. Carlson and S. J. Giovannoni [eds.], Annual
 review of marine science, vol 4. Ann. Rev. Mar. Sci.
- Caron, D. A., A. F. Michaels, N. R. Swanberg, and F. A. Howse. 1995. Primary productivity
 by symbiont-bearing planktonic sarcodines (acantharia, radiolaria, foraminifera) in
 surface waters near bermuda. J. Plankton Res. 17: 103-129.
- Caron, D. A., and N. R. Swanberg. 1990. The ecology of planktonic sarcodines. Rev. Aquat.
 Sci. 3: 147-180.
- Caruana, A. M. N., and G. Malin. 2014. The variability in DMSP content and DMSP lyase
 activity in marine dinoflagellates. Prog. Oceanogr. 120: 410-424.
- 609 Charlson, R. J., J. E. Lovelock, M. O. Andreae, and S. G. Warren. 1987. Oceanic
- 610 phytoplankton, atmospheric sulfur, cloud albedo and climate. Nature **326**: 655-661.

- de Vargas, C., and others 2015. Eukaryotic plankton diversity in the sunlit ocean. Science
 348.
- 613 Decelle, J., S. Colin, and R. A. Foster. 2015. Photosymbiosis in marine planktonic protists. In
- S. Ohtsuka and T. Suzaki, Horiguchi, T., Suzuki, N., Not, F. [eds.], Marine protists
 diversity and dynamics. Springer.
- Decelle, J., and others 2012. An original mode of symbiosis in open ocean plankton. Proc.
 Natl. Acad. Sci. U S A 109: 18000-18005.
- 618 Dennett, M. R., D. A. Caron, A. F. Michaels, S. M. Gallager, and C. S. Davis. 2002. Video
- plankton recorder reveals high abundances of colonial radiolaria in surface waters of
 the central north pacific. J. Plankton Res. 24: 797-805.
- Deschaseaux, E. S. M., and others 2014. Effects of environmental factors on dimethylated
 sulfur compounds and their potential role in the antioxidant system of the coral
 holobiont. Limnol. Oceanogr. 59: 758-768.
- Exton, D. A., T. J. McGenity, M. Steinke, D. J. Smith, and D. J. Suggett. 2015. Uncovering
 the volatile nature of tropical coastal marine ecosystems in a changing world. Glob.
 Chang. Biol. 21: 1383-1394.
- 627 Frade, P., V. Schwaninger, B. Glasl, R. W. Hill, R. Simó, and G. J. Herndl. 2016.
- Dimethylsulfoniopropionate in corals and its interrelations with bacterial assemblagesin coral surface mucus. Environ. Chem. 13: 252-265.
- Gage, D. A., D. Rhodes, K. D. Nolte, W. A. Hicks, T. Leustek, A. J. L. Cooper, and A. D.
- Hanson. 1997. A new route for synthesis of dimethylsulphoniopropionate in marine
 algae. Nature **387**: 891-894.
- Gali, M., V. Salo, R. Almeda, A. Calbet, and R. Simó. 2011. Stimulation of gross
 dimethylsulfide (DMS) production by solar radiation. Geophys. Res. Lett. 38.

- Garces, E., E. Alacid, A. Rene, K. Petrou, and R. Simó. 2013. Host-released dimethylsulphide
 activates the dinoflagellate parasitoid parvilucifera sinerae. ISME J. 7: 1065-1068.
- 637 Giesemann, A., H. J. Jager, A. L. Norman, H. P. Krouse, and W. A. Brand. 1994. Online
- sulfur-isotope determination using an elemental analyzer coupled to a massspectrometer. Anal. Chem. 66: 2816-2819.
- Guidi, L., S. Chaffron, L. Bittner and others. 2016. Plankton networks driving carbon export
 in the oligotrophic ocean. Doi:10.108/nature16942
- Hill, R. W., J. W. H. Dacey, and A. Edward. 2000. Dimethylsulfoniopropionate in giant clams
 (tridacnidae). Biol. Bull. 199: 108-115.
- Kaplan, I. R., and S. C. Rittenberg. 1964. Microbiological fractionation of sulphur isotopes. J.
 Gen. Microbiol. 34: 195-212.
- Karsten, U., S. Koch, J. A. West, and G. O. Kirst. 1996. Physiological responses of the
 eulittoral macroalga stictosiphonia hookeri (rhodomelaceae, rhodophyta) from
 argentina and chile: Salinity, light and temperature acclimation. Eur. J. Phycol. 31:
 361-368.
- Kaye, J. A. 1987. Mechanisms and observations for isotope fractionation of molecular-species
 in planetary-atmospheres. Rev Geophys 25: 1609-1658.
- Keller, M. D., Selvin, R. C., Claus, W. & Guillard, R. R. L. 1987. Media for the culture of
 oceanic ultraphytoplankton. J. Phycol. 23:633–8.
- Keller, M. D., W. K. Bellows, and R. R. L. Guillard. 1989. Dimethyl sulfide production in
 marine phytoplankton Acs Symposium Series 393: 167-182.
- Kiene, R., and D. Slezak. 2006. Low dissolved DMSP concentrations in seawater revealed by
- 657 small-volume gravity filtration and dialysis sampling. Limnol. Oceanogr. Methods 4:658 80-95.

Kirst, G. O. 1990. Salinity tolerance of eukaryotic marine-algae. Annu. Rev. Plant. Biol. 41:
21-53.

Kirst, G. O., C. Thiel, H. Wolff, J. Nothnagel, M. Wanzek, and R. Ulmke. 1991.

- Dimethylsulfoniopropionate (DMSP) in ice-algae and its possible biological role. Mar.
 Chem. 35: 381-388.
- Lana, A., R. Simó, S. M. Vallina, and J. Dachs. 2012. Re-examination of global emerging
 patterns of ocean DMS concentration. Biogeochemistry 110: 173-182.

666 Lavoie, M., Levasseur, M., and M. Babin. 2015. Testing the potential ballast role for

- dimethylsulfoniopropionate in marine phytoplankton: A modeling study. J PlanktonRes. 0:1-13.
- Lavoie, M., Raven, J. A., and M. Levasseur. 2016. Energy cost and putative benefits of
 cellular mechanisms modulating buoyancy in a flagellate marine phytoplankton. J.
 Phycol. 52:239-51.
- Lyon, B. R., J. M. Bennett, P. A. Lee, M. G. Janech, and G. R. DiTullio. 2016 Role of
- dimethylsulfoniopropionate as an osmoprotectant following gradual salinity shifts in
 the sea-ice diatom fragilariopsis cylindrus. Environ. Chem. 13: 181-194.
- Malin, G., and M. Steinke. 2004. Dimethyl sulfide production: What is the contribution of the
 coccolithophores? In: Thierstein HR, Young JR (eds) Coccolithophores: from
 molecular processes to global impact. Springer, Berlin, p 127–164
- Marie, D., F. Partensky, S. Jacquet, and D. Vaulot. 1997. Enumeration and cell cycle analysis
 of natural populations of marine picoplankton by flow cytometry using the nucleic
- acid stain SYBR green I. Applied and environmental microbiology 63: 186-193.
- 681 Michaels, A. F. 1988. Vertical-distribution and abundance of acantharia and their symbionts.

682 Mar. Biol. **97:** 559-569.

- Norman, A. L., L. A. Barrie, D. Toom-Sauntry, A. Sirois, H. R. Krouse, S. M. Li, and S. 683
- Sharma. 1999. Sources of aerosol sulphate at alert: Apportionment using stable 684 isotopes. J. Geophys. Res.: Atmospheres 104: 11619-11631. 685
- Not, F., J. del Campo, V. Balague, C. de Vargas, and R. Massana. 2009. New insights into the 686 diversity of marine picoeukaryotes. Plos One 4: e7143 687
- Nowack, E. C., and M. Melkonian. 2010. Endosymbiotic associations within protists. Phil. 688 Trans. R. Soc. B, Biol. Sci. 365: 699-712. 689
- 690 Oduro, H., K. L. Van Alstyne, and J. Farquhar. 2012. Sulfur isotope variability of oceanic
- DMSP generation and its contributions to marine biogenic sulfur emissions. Proc. 691 Natl. Acad. Sci. US A 109: 9012-9016. 692
- Patris, N., R. Delmas, M. Legrand, M. De Angelis, F. A. Ferron, M. Stiévenard, and J. Jouzel. 693 2002. First sulfur isotope measurements in central greenland ice cores along the 694 695 preindustrial and industrial periods. J. Geophys. Res: Atmos 107: ACH 6-1-ACH 6-11.
- 696
- 697 Probert, I., R. Siano, C. Poirier and others 2014. Brandtodinium gen. Nov. And b. Nutricula comb. Nov. (dinophyceae), a dinoflagellate commonly found in symbiosis with 698 polycystine radiolarians. J. Phycol. 50: 388-399. 699
- Quinn, P. K., and T. S. Bates. 2011. The case against climate regulation via oceanic 700 701 phytoplankton sulphur emissions. Nature 480: 51-56.
- Raina, J. B., E. A. Dinsdale, B. L. Willis, and D. G. Bourne. 2010. Do the organic sulfur 702 compounds DMSP and DMS drive coral microbial associations? Trends Microbiol. 703 **18:** 101-108. 704
- Raina, J. B., D. M. Tapiolas, S. Foret and others 2013. DMSP biosynthesis by an animal and 705 706 its role in coral thermal stress response. Nature 502: 677-680.
 - 30

707	Said-Ahmad, W., and A. Amrani. 2013. A sensitive method for the sulfur isotope analysis of
708	dimethyl sulfide and dimethylsulfoniopropionate in seawater. Rapid Commun. Mass
709	Sp.: RCM 27: 2789-2796.
710	Savoca, M. S., and G. A. Nevitt. 2014. Evidence that dimethyl sulfide facilitates a tritrophic
711	mutualism between marine primary producers and top predators. Proc. Natl. Acad.
712	Sci. U S A 111: 4157-4161.
713	Schoemann, V., S. Becquevort, J. Stefels, V. Rousseau, and C. Lancelot. 2005. Phaeocystis
714	blooms in the global ocean and their controlling mechanisms: A review. J. Sea. Res.
715	53: 43-66.
716	Seymour, J. R., R. Simó, T. Ahmed, and R. Stocker. 2010. Chemoattraction to
717	dimethylsulfoniopropionate throughout the marine microbial food web. Science 329:
718	342-345.
719	Siano, R., M. Montresor, I. Probert, F. Not, and C. de Vargas. 2010. Pelagodinium gen. Nov.
720	And p. Beii comb. Nov., a dinoflagellate symbiont of planktonic foraminifera. Protist
721	161: 385-399.
722	Simó, R. 2001. Production of atmospheric sulfur by oceanic plankton: Biogeochemical,
723	ecological and evolutionary links. Trends Ecol. Evol. 16: 287-294.
724	Stefels, J. 2000. Physiological aspects of the production and conversion of DMSP in marine
725	algae and higher plants. J. Sea. Res. 43: 183-197.
726	Stefels, J., M. Steinke, S. Turner, G. Malin, and S. Belviso. 2007. Environmental constraints
727	on the production and removal of the climatically active gas dimethylsulphide (DMS)
728	and implications for ecosystem modelling. Biogeochemistry 83: 245-275.

729	Steinke, M., P. Brading, P. Kerrison, M. E. Warner, and D. J. Suggett. 2011. Concentrations
730	of dimethylsulfoniopropionate and dimethyl sulfide are strain-specific in symbiotic
731	dinoflagellates (symbiodinium sp., dinophyceae). J. Phycol. 47: 775-783.

- 732Stemmann, L., M. Youngbluth, K. Robert and others 2008. Global zoogeography of fragile
- macrozooplankton in the upper 100-1000 m inferred from the underwater video
- 734 profiler. ICES J. Mar. Sci. **65:** 433-442.
- Stoecker, D. K., M. D. Johnson, C. deVargas, and F. Not. 2009. Acquired phototrophy in
 aquatic protists. Aquat. Microb. Ecol. 57: 279-310.
- 737 Summers, P., K. Nolte, A. Cooper, T. Leustek, D. Rhodes, and A. Hanson. 1998.
- 738 Identification and stereospecificity of the first three enzymes of 3-
- dimethylsulfoniopropionate biosynthesis in a chlorophyte alga. Plant. Physiol. 116:369-378.
- Sunda, W., D. J. Kieber, R. P. Kiene, and S. Huntsman. 2002. An antioxidant function for
 DMSP and DMS in marine algae. Nature 418: 317-320.
- 743 Suzuki, N., and F. Not. 2015. Biology and ecology of radiolaria. 179-222.
- Swanberg, N. R., and O. R. Anderson. 1985. The nutrition of radiolarians: Trophic activity of
 some solitary spurnellaria. Limnol. Oceanogr. 30: 646-652.
- 746 Vairavamurthy, A., M. O. Andreae, and R. L. Iverson. 1985. Biosynthesis of dimethylsulfide
- and dimethylpropiothetin by hymenomonas-carterae in relation to sulfur source andsalinity variations. Limnol. Oceanogr. 30: 59-70.
- 749 Van Alstyne, K. L., V. J. Dominique, III, and G. Muller-Parker. 2009. Is
- dimethylsulfoniopropionate (DMSP) produced by the symbionts or the host in an
- anemone-zooxanthella symbiosis? Coral Reefs **28**: 167-176.

- Van Alstyne, K. L., P. Schupp, and M. Slattery. 2006. The distribution of
- dimethylsulfoniopropionate in tropical pacific coral reef invertebrates. Coral Reefs 25:
 321-327.
- Yuasa, T., T. Horiguchi, S. Mayama, and O. Takahashi. 2016. Gymnoxanthella radiolariae
- gen. Et sp. Nov. (dinophyceae), a dinoflagellate symbiont from solitary polycystine
- radiolarians. J. Phycol. **52:** 89-104.

758 ACKNOWLEDGMENTS

We thank the ASSEMBLE (Association of European Marine biological Laboratories) 759 European program for supporting the field campaigns conducted from the Interuniversity 760 761 Institute for Marine Science (IUI, Israel) and the Observatory Oceanologique de Villefranche sur Mer (obs-vlfr, France). Particularly to Simon Berkowicz, Keren Zandbak and Yeala 762 763 Shaked for organizing our visit and help with the sampling and experimental logistics at the IUI; John Dolan for sharing his laboratory facilities in obs-vlfr, Fabien Lombard and Sophie 764 Marro for their time and assistance with the sampling and logistic support. Many thanks to 765 766 Ian Probert and Estelle Bigeard for their invaluable contribution during the sampling and 767 experiments set up. We are in debt with Eva Bucciarelli for generously sharing her time and analytical facilities at LEMAR (Brest, France) for DMSP analysis. We thank Cliff Law for his 768 comments and help copy editing the document. A.G.R. was supported by a Region Bretagne 769 postdoctoral fellowship SYMBIOX and by the National Institution for Water and 770 Atmospheric research (NIWA, New Zealand). T.B. was supported by the UMPC Emergence 771 program and L.P. by the SNF Seastar project. LP was supported by the Swiss National 772 Science Foundation postdoctoral grant P2GEP3 148800. A.A. thanks the Binational Science 773 Foundation Grant 2010407 for partial funding of this work. R.S. acknowledges support from 774 the Spanish Ministry of Science and Innovation through the SUMMER project (CTM2008-775 03309). 776

778 Figure Captions

- 779 Figure 1. Symbiotic Radiolaria and Foraminifera specimens isolated from the NW Mediterranean and Red sea surface waters imaged under the binocular (a, b, d, e, f, g) and light microscopy (c, 780 h. i). Single-celled acantharians Amphilonche elongata (a), 'Star' (b), 'Translucid' (c) 781 782 morphotypes with *Phaeocystis* sp. endosymbiotic algae. Single-celled Foraminifera 783 Globigerinella sp. with Pelagodinium beii endosymbionts (d), Solitary Thalassicolla sp. (e), and colonial Sphaerozoum sp. (f, h) and Collozoum sp. (g, i) collodarian specimens with 784 Brandtodininium nutricula endosymbionts. 785 Figure 2. Schematic representation of the experimental design and sequence of operations followed for 786 (a) the sulfate assimilation and (b) the DMS production experiments described in the 787 788 methods. Figure 3. Sulfur isotopic composition of intracellular DMSP (δ^{34} S-DMSP) of freshly-collected 789 790 symbiotic Radiolaria (Acantharia, solid-brown; Collodaria, green stars) and Foraminifera 791 (red square), of cultured free-living symbionts isolated from the same radiolarian groups B. 792 nutricula (green triangles) and Phaeocystis (empty-brown), and of natural microbial assemblages collected from surface waters. Seawater sulfate isotopic composition (δ^{34} S-793 SO_4^{2-}) (grev diamond). The range of δ^{34} S-DMSP from previous measurements in a bloom of 794 Prorocentrum minimum and isolated macroalgae (Oduro et al. 2012) and surface seawater 795 796 (Amrani et al. 2013, area between dash lines) are shown as reference. Figure 4. Sulfur isotopic composition of sulfate (δ^{34} S-SO₄²⁻) and intracellular DMSP (δ^{34} S-DMSP) in 797 cultured free-living Phaeocystis RCC1383 (A), Brandtodinium nutricula (B), and freshly-798
- 800 culture media containing sulfate with standard ('Standard') and light ('Light') δ^{34} S-SO₄²⁻, 801 respectively. Cell abundance for free-living cultured microalgae incubations is shown.

collected Collodaria-Brandtodinium holobiont (C) incubated for 3-day in two different

Figure 5. Sulfur isotopic composition of particulate DMSP (δ^{34} S-DMSP, black bars) and dissolved 802 DMS (δ^{34} S-DMS, white bars) in freshly-collected symbiotic Radiolaria (Acantharia-803 804 *Phaeocystis*) and microbial assemblage sampled from the same waters in the Red Sea where the Radiolaria were collected, measured after 4 hours of incubation under natural sunlight 805 806 conditions in surface waters. Squares with the error bars represent the concentration of 807 dissolved DMSP (black) and DMS (white) at the end of the incubation. Error bars represent 808 the standard error of the mean. Microbial assemblage (wsw), Radiolaria assemblage 809 (fsw+rads), Filtered seawater (fsw-only).

35

810 TABLE 1. Cell and colony size, DMSPt cell content, and intracellular concentration in uncultured

symbiotic Radiolaria and Foraminifera (host+microalgae), and cultured free-living symbiotic

812 microalgae isolated from the same species of Radiolaria and Foraminifera. The abundance of

radiolarian cells per colony and the abundance of symbiont cells per holobiont were assessed from

814 image analysis of freshly-collected organisms. DMSPt cellular content per symbiont *in hospite*

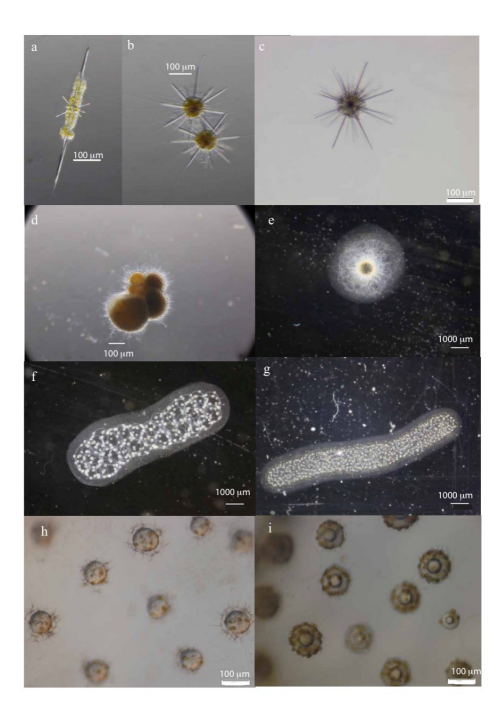
estimated assuming all DMSPt measured in the holobiont is allocated in the symbiotic microalgae. Cc

refers to central capsule, n/a and n/d refer to not applicable and not determined, respectively. The

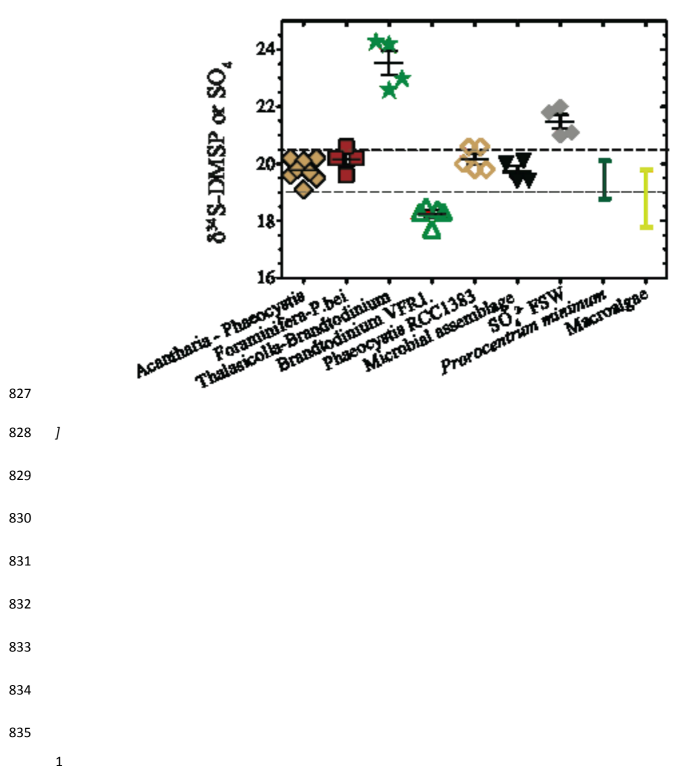
817 number of measurements are shown in brackets. The errors refer to the standard deviation of the mean.

TABLE 1 (All growth phases)			Size (um) Major axis (µm)	Size (um) Minor axis (µm)	DMSPt Content pmol cell ⁻¹	DMSPt Concentration mmol L ⁻¹	Number central capsules colony ⁻¹
Collodaria-Brandtodin	nium sp.						
(Radiolaria)							
Collozoum sp.	Colonial	Symbiotic	7267 ±1793(3)	3754 ±243(3)	3652 ±2008(3)	$0.1 \pm 0.1(3)$	509 ±72(3)
Sphaerozoum sp.	Colonial	Symbiotic	8242 ±2041(4)	3463 ±1327(4)	3135 ±236(3)	0.1 ±0.01(3)	166 ±65(17)
<i>Thalassicolla</i> sp.	Solitary	Symbiotic	3972 ±1148(5)	3517 ±939(5)	2757 ±750(5)	0.2 ±0.1 (5)	n/a
Acantharia-Phaeocyst (Radiolaria)	is sp.						
Amphilonche elongata	Single- celled	Symbiotic	391 ±117(31)	64 ±30(31)	15.2 ±5.4(4)	$17.1 \pm 6.1(4)$	n/a
Translucid' morphotype	Single- celled	Symbiotic	483 ±188(12)	437 ±182(12)	20.9 ±3.6(3)	0.3 ±0.1(3)	n/a
Star shape' morphotype	Single- celled	Symbiotic	307 ±114(42)	242 ±119(42)	36.4 ±10.4(5)	2.4 ±0.7 (5)	n/a
Foraminifera-P. beii							
<i>Globigerinella sp.</i> (Planktonic)	Single- celled	Symbiotic	235 ±87(31)	209 ±58(31)	37.8 ±15.5(5)	7.5 ±3.1(5)	n/a
Free-living symbiotic							
Brandtodinium nutricula (RCC3468) (Dinophyceae)	Single- celled	Symbiotic	12 ±0.9(15)	8.6 ±0.7(15)	0.2 ±0.1(12)	490 ±107(12)	n/a
Pelagodinium beii (RCC1491) (Dinophyceae)	Single- celled	Symbiotic	20.5 ±1.8(18)	17.7 ±2.3(18)	0.9 ±0.2 (12)	272 ±49(12)	n/a
<i>Gymnoxanthella</i>	Single-	Symbiotic	9.1	7.9	0.1	462	n/a

radiolariae	celled		±1.0(15)	±0.6(15)	$\pm 0.1(12)$	±168(12)	
(RCC3507)							
(Dinophyceae)							
Phaeocystis	Single-	Symbiotic	4.0	4.0	0.01	307	n/a
(RCC1383)	celled		nominal	nominal	$\pm 0.001(3)$	$\pm 47(3)$	
						• (-)	
(Prymnesiophyceae)						. (-)	

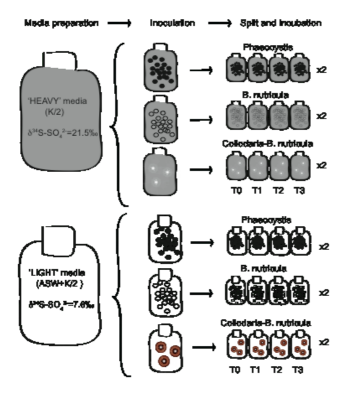


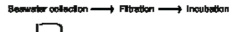
826 Figure 2

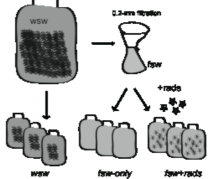


a - Sulfate assimilation experiment

b - DMS production experiment

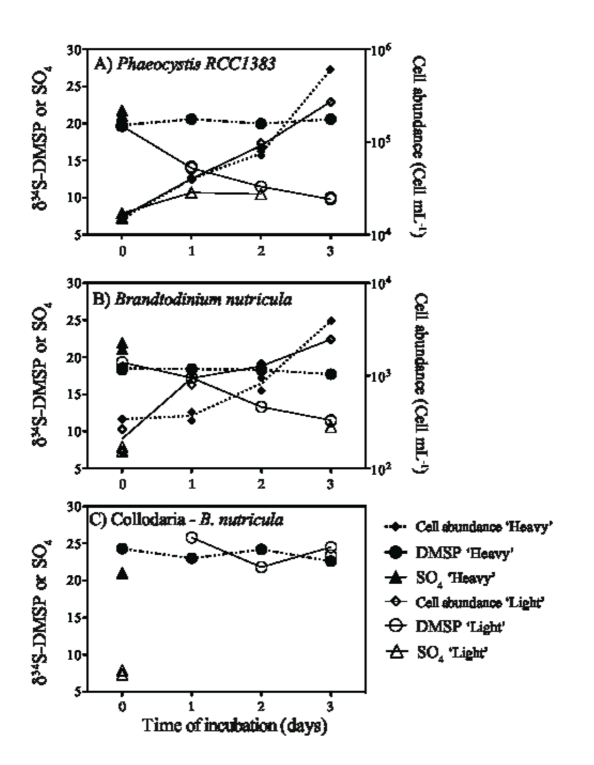


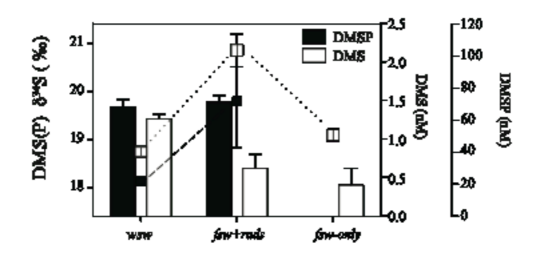




838

839 Figure 4

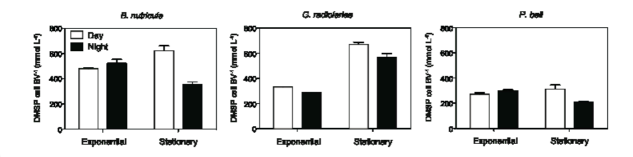




845

846 <u>SUPPORTING INFORMATION</u>

847 Intracellular DMSPt concentration of symbiotic microalgae as a function of growth phase and diel cycle. The large error associated to the mean DMSPt cell content and concentration values of 848 849 cultured symbiotic microalgae (Table 1) reflected the variability associated to growth phase and 850 photoperiod at the time of harvesting the culture. The photoperiod (day vs. night) had a significant 851 effect on DMSP cellular concentration for *B. nutricula* and *G. radiolariae*, while this was only marginally significant for *P. beii* (two-way ANOVA, p=0.066). The effect of growth phase 852 853 (exponential vs. stationary) on DMSPt content was only significant for G. radiolariae (two-way 854 ANOVA, p<0.0001), while the interaction between both photoperiod and growth phase was 855 significant for B. nutricula and P. beii (p<0.001). During exponential phase, DMSP cell content was 856 slightly lower during the day than night for B. nutricula and P. beii, while the cellular DMSP 857 concentrations decreased substantially at night for these strains. In G. radiolariae DMSP content decreased at night during both exponential and stationary growth phases (p=0.003, Supplementary 858 859 Figure 1).





861 Supplementary Figure 1. Mean intracellular DMSP concentration in free-living cultures of

- 862 Brandtodinium nutricula (A), Gymnoxanthella radiolariae (B), and Pelagodinium bei (C) harvested at
- 863 different growth phases (exponential vs. stationary) and times of the daily photoperiod (day vs night).
- 864 Error bars represent the standard error of the mean. A) Growth $F_{1,8}=0.17$, p=0.69, photoperiod
- 865 $F_{1,8}=16.6 \text{ p}=0.0036$, interaction $F_{1,8}=32.1$, p=0.0005; B) Growth $F_{1,8}=304$, p<0.0001, photoperiod
- 866 $F_{1,8}=17.7$, p=0.0030, interaction $F_{1,8}=2.7$, p=0.14; C) Growth $F_{1,8}=1.4$, p=0.27, photoperiod $F_{1,8}=4.5$,
- 867 p=0.066, interaction $F_{1,8}$ =12.9, p=0.0071. Two-way ANOVA not repeated measures.