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1 Origin and fate of long-chain polyunsaturated fatty acids in the Kerguelen

2 Islands region (Southern Ocean) in late summer

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11 Highlights:

10

FA profiles of SPOM from the Kerguelen region during the post-bloom period was
 dominantly composed of PUFA with high proportions of the two LC-PUFA 20:5n-3 and
 22:6n-3.

- Abundance of LC-PUFA in the mixed layer derived from the phytoplankton community
 composed of small species (prymnesiophytes and prasinophytes) and diatoms.
- 17 In the upper mesopelagic, LC-PUFA are maintained at high proportions according to
- 18 distinct pathways, export of diatoms for 20:5n-3 and zooplankton fecal material for 22:6n-
- 19
- SPOM revealed high nutritional quality in the upper water column (0-300m) both in the
 iron-fertilized area on the Plateau and outside in HNLC waters.

22 Abstract:

3.

Long-chain polyunsaturated fatty acids (LC-PUFA) are molecules produced at the basis of marine food webs and essential for ecosystem functioning. This study reports detailed fatty acid (FA) composition including the two LC-PUFA 20:5n-3 and 22:6n-3, in suspended organic matter (SPOM) from the upper 300 m collected in the Kerguelen Island region in the Southern Ocean during the post-bloom period (February–March 2018; project MOBYDICK). FA profiles were largely dominated by PUFA (53-69 % of Total Fatty Acid, TFA) regardless of stations and among PUFA, proportions of LC-PUFA were especially high, making up 27-44 % of TFA both in the ML 30 and upper mesopelagic. 20:5n-3 and 22:6n-3 co-occurred in the ML as a result of the post-bloom 31 phytoplankton community showing a mixed composition dominated by small size 32 phytoplankton (prymnesiophytes and prasinophytes) supplying 22:6n-3, and with diatoms in 33 lower proportions supplying 20:5n-3. Elevated levels of LC-PUFA were observed both inside the 34 iron-fertilized area on the Kerguelen Plateau and downstream, and outside in High Nutrient Low 35 Chlorophyll waters located upstream of the Plateau, and appeared unrelated to site. In the upper 36 mesopelagic, both LC-PUFA were maintained at high relative proportions suggesting an efficient 37 and possibly fast vertical transfer from the surface. Transfer with depth seems to proceed via 38 distinct pathways according to LC-PUFA. 20:5n-3 may be exported along with diatoms, 39 presumably in the form of large intact cells, aggregates as well as resting spores. For 22:6n-3, 40 transfer may involve a channeling through the heterotrophic food web resulting in its association with fecal material at depth. Channeling of 22:6n-3 could involve heterotrophic protists such as 41 42 dinoflagellates and ciliates grazing on small phytoplankton, as well as larger zooplankton such as 43 copepods and salps, possibly feeding on microzooplankton and producing fecal pellets rich in 44 22:6n-3. According to LC-PUFA content, SPOM present throughout the upper water column (0-45 300 m) appeared of high nutritional quality both on- and off-plateau, and represented a valuable source of food for secondary consumers and suspension feeders. 46

47 Keywords: Essential fatty acids, vertical distribution, fatty acid export, phytoplankton
48 diversity, diatoms, heterotrophic interactions, nutritional quality.

50 A. INTRODUCTION

51 The Southern Ocean (SO) is a vast and contrasted environment where the characteristics of 52 pelagic ecosystems are particularly diverse. The Antarctic Circumpolar Current (ACC) composed 53 of successive hydrographic fronts (Orsi et al., 1995) acts as a strong physical and biogeochemical 54 boundary for biological activity, and to the south and until the Antarctic sea ice, lies the largest 55 High Nutrient Low Chlorophyll (HNLC) zone of the world ocean. The productivity is low 56 throughout the year in HNLC open waters essentially due to a lack of dissolved iron (Blain et al., 57 2007; de Baar et al., 1995; Martin, 1990), and high concentrations of unused macronutrients 58 (phosphate, nitrate, silicic acid) persist in surface waters. In contrast, high productivity regimes 59 are only found close or downstream of Subantarctic islands (South-Georgia, Crozet, Kerguelen, 60 Heard) and continental shelves, which receive natural and persistent iron inputs (Blain et al., 2008; 61 Perissinotto and Duncombe Rae, 1990; Pollard et al., 2007; van der Merwe et al., 2015; Venables 62 and Moore, 2010). Iron fertilization allows phytoplankton, mainly initially composed of diatoms 63 and also *Phaeocystis*, to thrive during intense and long-lasting blooms in austral spring/summer 64 (Cavagna et al., 2015; Korb and Whitehouse, 2004; Mongin et al., 2008; Schallenberg et al., 2018; Schlosser et al., 2018; Seeyave et al., 2007). These blooms provide abundant food and energy to 65 66 heterotrophic organisms and higher trophic levels, and help to maintain highly productive food 67 webs extended from microbes to zooplankton, micronekton, up to emblematic top-predators of 68 the SO (marine mammals, sea-birds, penguins, whales) (El-Sayed, 1988; Evans and Brussaard, 69 2012; Pakhomov and McQuaid, 1996).

70 In the current context of climate change, major modifications are predicted for the SO such as 71 sea surface warming, increasing stratification and reduction in nutrient supplies, southward 72 migration of oceanographic fronts, shrinking of sea ice, as well as ocean acidification, and it 73 already concerns some areas like the Antarctic peninsula (Constable et al., 2014; Moline et al., 74 2004). These climate-induced changes are likely to affect phytoplankton distribution and 75 composition (Deppeler and Davidson, 2017), and hence to have positive or negative cascade 76 effects on the whole food web. It is therefore important to determine what parameters control the 77 present-day stability, health and resilience of these pristine ecosystems.

78 Among the parameters that maintains ecosystem stability, the nutritional value of 79 phytoplankton is crucial, as it determines the amount of energy that can flow through the entire 80 food web and controls its overall functioning. Part of nutritional quality for marine organisms is 81 attributable to lipids, in particular to n-3 long-chain polyunsaturated fatty acids (n-3 LC-PUFA) 82 such as 20:5n-3 (eicosapentaenoic acid, EPA) and 22:6n-3 (docosahexaenoic acid, DHA). n-3 LC-83 PUFA are molecules essential to all organisms and are involved in a variety of physiological 84 processes such as growth, immunity, cell membrane function regulation and energy storage 85 (Guschina and Harwood, 2006; Parrish, 2013). They play important roles in trophic interactions, 86 starting from secondary consumers such as copepods and other crustacean zooplankton, for 87 which they determine growth potential, reproduction success and general fitness (Brett et al., 88 2009; Brett and Müller-Navarra, 1997; Müller-Navarra et al., 2000; Pond et al., 2005). LC-PUFA are 89 synthetized at the basis of the food web, essentially by phytoplankton and to some extent by small 90 heterotrophs, and the supplies to consumers occur necessarily through diet (Bec et al., 2006; Guo 91 et al., 2017; Klein-Breteler et al., 1999). The production of LC-PUFA by phytoplankton, and fatty 92 acids (FA) in general, is highly species-specific both at the phylum and class levels (Cañavate, 93 2019; Jónasdóttir, 2019; Parrish, 2013) and enables their use as valuable biomarkers (Dalsgaard et 94 al., 2003). Another consequence is that marine algae do not all have the same nutritional value, 95 the most valuable algae in terms of LC-PUFA proportions includes Prymnesiophyceae, 96 Pavlovaphyceae, Bacillariophyceae, Dinophyceae, Cryptophyceae, Eustigmatophyceae followed 97 by, Prasinophyceae and Mamiellophyceae (Jónasdóttir, 2019). The lowest contents are found in 98 Chlorophyceae, Trebouxiophyceae and especially in Cyanobacteria, which are devoid of LC-99 PUFA (Jónasdóttir, 2019).

In the SO, LC-PUFA are presumably supplied by two distinct groups of phytoplankton, diatoms and flagellates, characterized by specific FA profiles. Diatoms produce preferentially 20:5n-3 along with C16 monounsaturated fatty acids (MUFA) 16:1n-7, and C₁₆ PUFA 16:3n-4 and 16:4n-1 (Dalsgaard et al., 2003; Parrish, 2013; Volkman et al., 1998), with only small amounts of 22:6n-3 found in centric diatoms (Dunstan et al., 1993). The other essential n-3 LC-PUFA 22:6n-3 together with C18 PUFA 18:5n-3 are produced preferentially by dinoflagellates as well as some Prymnesiophyceae like the coccolithophore *Emiliana huxleyi* (Jónasdóttir, 2019; Okuyama et al.,

107 1992; Thomson et al., 2004) but this latter group is scarce in the SO, and thus it is not assumed to 108 be a potential producer. Another important Prymnesiophyceae species, *Phaeocystis sp.*, produces 109 low levels of LC-PUFA (Nichols et al., 1991; Skerratt et al., 1995), but it can represent an important 110 part of the autotrophic biomass especially in the early and late phase of the diatom bloom. This 111 general picture of LC-PUFA sources is generally adopted in diet studies and relies on the 112 identification of diagnostic FA in consumers (Atkinson et al., 2006; Ericson et al., 2018; Hellessey 113 et al., 2020). This generalized view is probably oversimplified as it mostly relies on FA signatures 114 obtained from culture and feeding experiments without accounting for field observations. 115 Moreover, it only takes into account a single autotrophic production pathway for LC-PUFA, 116 whereas alternative routes via heterotrophy and upgrading may exist. As an example, interactions 117 among grazers such as copepods and heterotrophic protozoans (dinoflagellates and ciliates) were 118 shown to substantially increase the nutritional quality of *Phaeocystis globosa*, most probably via the 119 production of LC-PUFA by heterotrophic upgrading (Tang et al., 2001).

120 For the SO, available data on the natural occurrence of LC-PUFA in phytoplankton, as well as 121 in protists in general, remain scarce, and it received a disproportionally low attention in 122 comparison to larger size organisms (Hellessey et al., 2020; Pond et al., 2005). The few available 123 data concern only surface samples obtained from highly productive regions of the marginal ice 124 zone and coastal Antarctic zone (Fahl and Kattner, 1993; Gillan et al., 1981; Hernando et al., 2018; 125 Nichols et al., 1993; Thomson et al., 2004) as well as of the frontal Subantarctic zone of the Indian 126 sector of the SO (Mayzaud et al., 2007, 2002). This clearly limits our understanding of LC-PUFA 127 cycling in both the surface euphotic zone where production takes place, and in the water column 128 where organic matter (OM) and associated LC-PUFA are considered as labile fraction, and could 129 be degraded, transformed, and even transferred to benthic communities (Budge and Parrish, 1998; 130 Rembauville et al., 2018; Wilson et al., 2010). Recent results obtained from five long-term sediment 131 traps deployed in naturally iron-fertilized settings and HNLC waters of the Polar Front (PF) 132 region (South Georgia, Crozet, Kerguelen) indicate large regional and seasonal fluctuations in FA 133 and PUFA composition of sinking OM (Rembauville et al., 2018). Variations are tightly linked to 134 the ecological vectors responsible for the export of OM (e.g., diatom resting spores, phytoplankton 135 aggregates, zooplankton fecal pellets) (Rembauville et al., 2016, 2015a, 2015b). On the Kerguelen Plateau, LC-PUFA such as 20:5n-3, which is usually associated to diatoms resting spores (mainly *Chaetoceros*), could be transferred at depth towards the sediments (Rembauville et al., 2018).

138 Among naturally Fe-fertilized regions, the Kerguelen Island area has received considerable 139 interest over recent decades as it is well suited for studying the contrasted properties of pelagic 140 ecosystems found in the SO. The Kerguelen Plateau, which extends to the southeast of the islands, 141 is a large-scale topographic feature with shallow bathymetry (< 700 m) that benefits from 142 enhanced Fe inputs via diapycnal mixing (Blain et al., 2008; Park et al., 2008) and is subjected to a 143 large-scale (45,000 km²) and long-lasting (October to February) diatom bloom (Mongin et al., 144 2008). Bloom phenology consists of two successive phases (Pellichero et al., 2020) with a first phase 145 occurring in spring and dominated by small size diatoms forming long-chains (Chaetoceros, Pseudo-nitzschia) (Lasbleiz et al., 2016) and a second phase occurring in summer and dominated 146 147 by larger size diatoms (Eucampia, Corethron) (Armand et al., 2008; Blain et al., 2021; Liu et al., 2020). 148 Productivity is also stimulated downstream of the Plateau (according to the West to East ACC 149 circulation) due to lateral advection of Fe-rich waters (Quéroué et al., 2015; Trull et al., 2015; van der Merwe et al., 2015), which promotes a diatom bloom from October to January. This latter 150 151 bloom is shorter in duration and lower in intensity, and composed of different phytoplankton 152 community composition in comparison to the central Plateau (Armand et al., 2008; Lasbleiz et al., 153 2016). Upstream of the Plateau, south-west of the Kerguelen Islands, open waters are away from 154 the influence of the Fe fertilized area, productivity is drastically reduced and considered representative of HNLC conditions found in the SO (Rembauville et al., 2017). Phytoplankton 155 156 biomass stays low throughout the year (climatological Chlorophyll a (Chl-a) < 0.3 µg.L⁻¹) except 157 during a short time period (~1 month) in December/January (Fiala et al., 1998; Kopczyńska et al., 158 1998). Under HNLC regime, pico- and nano- communities composed mostly by Phaeocystis sp. 159 predominate most of the time, except during summer when microphytoplankton, such as small 160 diatoms (Fragilariopsis sp.) and autotrophic dinoflagellates, are more abundant (Armand et al., 161 2008; Fiala et al., 1998; Kopczyńska et al., 1998; Lasbleiz et al., 2016; Rembauville et al., 2017).

162 The MOBYDICK project (Marine Ecosystem Biodiversity and Dynamics of Carbon around 163 Kerguelen: an integrated view) was designed to complement the available description of the

164 Kerguelen region ecosystems and the main objectives were 1) to track carbon from its initial 165 fixation at the surface to its channeling through the food web and its transfer at depth via export, 166 and 2) to perform a detailed description of the diversity at each trophic level. The oceanographic 167 survey was carried out in late summer/early autumn (February-March 2018) corresponding to the 168 demise of the diatom bloom, a period that was not previously investigated by other surveys 169 carried out during the onset (KEOPS2, October-November 2011) and decline (KEOPS1, January-170 February 2005) of the bloom. In this study, we investigated the distribution of FA, including the 171 essential LC-PUFA 20:5n-3 and 22:6n-3, in suspended organic matter (SPOM) collected in the 172 upper water column. Our aims were to explore the origin and fate of essential LC-PUFA from the 173 surface to the upper mesopelagic zone at sites with contrasted seasonal productivity regimes 174 found on and off the Kerguelen Plateau. Statistical analyses were used to identify the main drivers 175 of FA profile variability. A selection of FA specific to phytoplankton classes and zooplankton 176 activity was used to identify the main producers of LC-PUFA and to explore the impact of 177 heterotrophic interactions in the upper water column. Finally, PUFA contents were used to 178 discuss the nutritional quality of suspended OM and a comparison between iron-fertilized and 179 HNLC conditions was addressed.

180 B. MATERIAL & METHODS

181 Studied area

182 The MOBYDICK survey was conducted in the Kerguelen Islands area in the Indian sector of 183 the SO on board of R/V Marion Dufresne II during late-austral summer 2018 from February 18th to 184 March 28th coinciding with the post bloom phase. Four stations (M1, M2, M3, and M4, Figure 1), 185 were selected to represent the contrasted production regimes encountered in the area and defined 186 on a seasonal basis. Station M2 (bottom depth 520 m) is located in the Fe-enriched area on the 187 central Kerguelen Plateau. This station corresponds to the station A3 investigated during KEOPS1 188 and KEOPS2 programs. Station M1 (bottom depth 2723 m), located downstream of the Plateau, 189 benefits also from enhanced Fe supplies and exhibit a seasonal moderate production regime. 190 Station M3 (bottom depth 1730 m) and M4 (bottom depth 4731 m) are both located upstream of 191 the Plateau and are representative of HNLC conditions. Station M3, previously investigated as 192 part of the KERFIX time series program (1990-1995) (Jeandel et al., 1998), was at the time of 193 sampling north of the PF in the Polar Frontal Zone (PFZ) according to Pauthenet et al. (2018), 194 while the three other stations were permanently south of the PF. During the survey, repeated 195 visits were performed at most stations at around 10-day interval except at station M1, which was 196 visited only once. As shown in Table 1, three visits were performed at station M2 (M2-1, M2-2, 197 and M2-3) and two at stations M3 and M4 (M3-1, M3-3, M4-1, M4-2).

198

Hydrological and biogeochemical data



Figure 1 Map of MOBYDICK station locations and monthly mean (March 2018) surface Chlorophyll-a concentrations (µg.L⁻¹) obtained from 4 km resolution Global Ocean Satellite Observations (Copernicus-Globcolour, Copernicus Marine Service, http://marine.copernicus.eu/). The black line refers to 1000 m bathymetry and the blue refers to the approximate position of the PF for the Feb.-Mar. period and drawn according to Pauthenet et al. (2018).

Vertical profiles of temperature, dissolved oxygen, and salinity were obtained at all stations using a SeaBird 911-plus CTD (Conductivity, Temperature, and Density) unit mounted on the rosette. Chl-a concentrations and PAR (photosynthetically available radiation) levels were measured using a fluorometer and LI-COR sensor, respectively. The mean depth of the mixed layer (MLD) was estimated based on a difference in potential density of 0.03 to the surface value (10 m) and using all CTD casts performed during the occupation of stations (Lafond et al., 2020).

- 205 The depth of the euphotic zone (Z_e) corresponded to the depth where light intensity was at least
- 206 1% of incident light at the surface (Table 1). Averaged Chl-a, silicic acid (Si(OH)₄) and ammonium
- 207 (NH4⁺) concentrations for the mixed layer (ML) are also shown in Table 1, details on the analytical
- 208 methods can be found elsewhere (Irion et al., 2020; Lafond et al., 2020).

Table 1. MOBYDICK station details, averaged depth of the mixed layer (MLD), depth of the euphotic zone (Ze), and averaged concentrations of Chl-a, silicic acid, ammonium in the mixed layer.

Station	Lat	Lon	Bottom depth	date	Visit	MLD ^a	Ze ^b	Chl-a		Si(OH)4		NH4	
	°S	°E	m			m	m	µg.L-1		µmol.L-1		µmol.L-1	
								Mean	S.D	Mean	S.D	Mean	S.D
M1	49.9	74.9	2723	10/03/2018	M1	63	89	0.34	0.04	6.7	0.3	0.57	0.21
M2	50.6	72.0	520	27/02/2018	M2-1	79	64	0.27	0.02	1.4	0.4	0.75	0.08
				08/03/2018	M2-2	73	61	0.31	0.05	1.7	0.8	1.12	0.03
				17/03/2018	M2-3	80	58	0.59	0.03	2.8	0.3	0.95	0.06
M3	50.7	68.1	1730	05/03/2018	M3-1	74	93	0.21	0.03	2.9	1.0	0.63	0.22
				19/03/2018	M3-3	96	105	0.14	0.00	2.3	0.2	0.73	0.01
M4	52.6	67.2	4731	03/03/2018	M4-1	69	95	0.19	0.01	4.4	0.4	0.37	0.03
				14/03/2018	M4-2	96	101	0.21	0.01	5.3	1.0	0.54	0.11

^a MLD: Average mixed layer depth estimated from multiple CTD casts at each station and according to potential density fluctuations < 0.03 kg.m⁻³ relative to 10 m depth.

^bZe: Depth of the euphotic zone defined as 1% of surface photosynthetic active radiation.

209

210 Sampling of suspended particles

211 Suspended particulate organic matter (SPOM) was sampled for FA determination, at all 212 visited stations (8 stations in total) using five in situ pumping systems (ISP) allowing large volume 213 of seawater to be filtered (200-1500 L) and sufficient amount of material to be collected. The five 214 ISP were deployed from the surface euphotic zone to the upper mesopelagic, down to 200 m (M1, 215 M2-2, M2-3, M4-2), 300 m (M2-1, M3-1, M3-3), and 600 m depth (M4-1). For three stations (M1, 216 M2-1, M4-1), only four depths could be sampled due to ISP failures. ISP were equipped for 217 sequential filtration, and we considered here only the 1-50 µm size fraction, which corresponds to 218 the largely dominant fraction in terms of Particulate Organic Carbon (POC, averaged weight 219 proportion 93% of total POC, n=37). For the first six stations (M1, M2-1, M2-2, M4-1, M4-2, M3-1),

the 1-50 μ m size fraction was obtained from a single filter made of high-purity quartz microfiber (QMA, Sartorius, France) of 1 μ m nominal pore size, and placed after two successive nylon (NITEX) screen pre-filters of 300 and 50 μ m mesh size. For the last two stations (M2-3 and M3-3), due to a change in the mesh size of screens (50 and 20 μ m), the 1-50 μ m fraction was obtained from the QMA filter (1-20 μ m) and from the intermediate screen of 20 μ m mesh size (20-50 μ m).

225 Sampl

Samples processing on board

226 After ISP recovery, the QMA filters were processed on-board as follows: subsampling was 227 conducted directly on the filter using a 25 mm plexiglass punch previously cleaned with ethanol. For FA analyses, four punches were taken and then extracted in 6 mL of 2:1 (v:v) 228 229 chloroform:methanol solvent and preserved at -20°C. For POC measurements, four other punches 230 were taken, dried at 55°C for 24 hours, and stored at room temperature. For the NITEX screens, 231 the 142 mm filters were cut into quarters using a scalpel (ethanol cleaned), and one quarter was 232 dedicated to POC analyses and another to FA analyses. The two remaining quarters were kept as 233 spare at -20°C. Particles from the NITEX were resuspended using filtered seawater (0.4 µm) and 234 recollected on a pre-combusted 0.7 µm nominal pore-size glass fiber filters (Whatman GF/F, 235 Maidstone, UK). The GF/F filters were processed as described for QMA filters for FA (dived in 6 236 mL of chloroform:methanol solvent and stored at -20°C) and POC analyses. Lipid extracts 237 dedicated to FA analyses were stored at -20°C during the cruise, shipped to home laboratory 238 (France) with dry ice and stored at -20°C until FA analyses.

239 Particulate organic carbon analyses

POC samples were fumed with hydrochloric acid (10M) for 4 hours to remove particulate inorganic carbon, dried at ambient temperature for 2 hours, subsampled with a 13 mm punch and encapsulated into tin caps following protocols by Brook et al. (2003) and Trull et al. (2015). POC content was determined using a Flash Elemental Analyzer 2000 coupled to a Thermo Fisher Delta V Plus stable light isotope ratio mass spectrometer system (IRMS, Bremen, Germany). Data were corrected for filter blank contribution, and acetanilide (C₈H₉NO) was used as a standard for POC content. Final concentrations of POC in seawater are reported in µmol.L⁻¹. 247

Fatty acids analysis

248

a. Lipid extraction and purification

249 Samples for FA analyses were re-extracted to remove any residual seawater. The initial lipid 250 extract was transferred in a new 22 mL vial, and then re-extracted with 3 mL of chloroform. The 251 sample was vigorously shaken and then centrifuged to insure a good phase separation. For each 252 sample, three re-extractions were performed. Lipid extracts were then evaporated with N2 gas, 253 resuspended in 6 mL of chloroform:methanol (2:1 v:v) and stored at -20°C until analysis. Lipid 254 extracts were separated into neutral and polar lipids following the method of Remize et al.(2020). 255 Briefly, 3 mL of total lipid extract was evaporated with nitrogen, recovered with three washes 256 using chloroform:methanol (final volume 1.5 mL, 98:2 v:v) and spotted at the top of a silica gel 257 column (40 mm ×4 mm, silica gel 60A 63–200 µm rehydrated with 6% H2O, 70–230 mesh, Sigma-258 Aldrich, Darmstadt, Germany). The neutral lipid fraction (NL) was eluted using 259 chloroform:methanol (10 mL 98:2 v:v) and the polar lipid fraction (PL) with methanol (20 mL). 260 Both fractions were then collected in glass vials, and an internal standard (C23:0, 2.3 µg) was 261 added.

262

b. Transesterification of FAME

Fatty acids methyl esters (FAME) transesterification was conducted according to the protocol described by Mathieu-Resuge et al. (2019). In brief, after evaporation to dryness of the NL and LP fractions, transesterification was performed by adding 0.8 mL of H₂SO₄/methanol mixture (3.4% v:v) to the lipid extract and heated at 100 °C for 10 min. Hexane (0.8 mL) and distilled water saturated with hexane (1.5 mL) were added. The lower MeOH–water phase was discarded after homogenization and centrifugation. Hexane fraction containing FAME was washed two more times with another 1.5 mL of distilled water.

270

c. Fatty acid analysis by gas chromatography

Analyses of FAME were performed on a Varian CP8400 gas chromatograph (Agilent, Santa Clara
CA, USA) using simultaneously two separations on a polar column (ZBWAX: 30 mm × 0.25 mm
ID × 0.2 μm, Phenomenex, Torrance, CA, USA) and an apolar column (ZB5HT: 30 m × 0.25 mm

- 274 ID \times 0.2 μm Phenomenex, Torrance, CA, USA). The temperature program used by the gas
- chromatograph was the following: first, initial heating to 0 from 150 °C at 50 °C.min⁻¹, then to 170
- 276 °C at 3.5 °C.min⁻¹, to 185 °C at 1.5 °C.min⁻¹, to 225 at 2.4 °C.min⁻¹ and finally to 250 °C at 5.5 °C.min⁻¹
- and maintained for 15 min. The FAME were identified by comparison of their retention time with
- 278 commercial and in-house standards mixtures as described in Remize et al. (2020).

Table 2. Complete list of FA determined in this study and grouped by compound family with saturated fatty acids (SAFA), monounsaturated fatty acids (MUFA), polyunsaturated fatty acids (PUFA), and branched fatty acids (Branched). Major FA (>1% of TFA of the mean of all samples) are highlighted in bold.

Group	Fatty acids
SAFA	14:0 , 15:0, 16:0 , 17:0, 18:0 , 20:0, 21:0, 22:0, 24:0
MUFA	14:1n-5, 15:1n-5, 16:1n-9, 16:1n-7 , 16:1n-5, 17:1n-7, 18:1n-9 , 18:1n-7 , 18:1n-5, 20:1n-9, 20:1n-7, 22:1n-9, 22:1n-7, 24:1n-9
	C16: 16:2n-7, 16:2n-6, 16:2n-4, 16:3n-4, 16:3n-6, 16:3n-3, 16:4n-3, 16:4n-1
	C18: 18:2n-6, 18:2n-4, 18:3n-6, 18:3n-4, 18:3n-3, 18:4n-1, 18:4n-3, 18:5n-3
PUFA	C ₂₀ : 20:2n-6, 20:3n-6, 20:3n-3, 20:4n-6, 20:4n-3, 20:5n-3
	C21: 21:5n-3
	C ₂₂ : 22:2n-6, 22:4n-6, 22:5n-6, 22:5n-3, 22:6n-3
Branched	l iso15:0, ante15:0, iso16:0, iso17:0

FA concentrations reported here correspond to the total lipid fraction (sum of PL and NL fractions) measured in 1-50 μ m SPOM. Concentrations are reported in μ g.L⁻¹ (of seawater) as well as in percent mass proportion relative to total fatty acids (%TFA). The complete list of FA considered and grouped by compound family is shown in Table 2. FA are reported using a shorthand notation of A:Bn-x, where A indicates the number of carbon atoms, B is the number of double bonds and x indicates the position of the first double bond relative to the terminal methyl group (Budge et al., 2006).

286 Fatty acid profiles in phytoplankton classes

In Table S1, we summarized available information on FA composition of phytoplankton classes provided by the meta-analysis of Jónasdóttir (2019) and Cañavate (2019) as well as the screening study of Mitani et al. (2017). This allows to highlight the relationships between some specific FA abundances and phytoplankton classes.

291 As seen in Table S1, the percentage of 14:0 is usually above 10% in Mamiellophycea (MAM), 292 Pavlovophycea (PAV), Pelagophycea (PEL), Cosciondiscophycea (COS) and Mediophycea (MED) 293 and above 15% in Coccolithophycea (COC), Prymnesiophycea (PRY), Pinguiophycea (PIN) and 294 Fragilariophycea (FRA). The percentage of 16:1n-7 is above 15% in PAV, Bacillariophyceae (BAC), 295 COS, FRA, and MED. The sum of 16:2n-7, 16:2n-4, 16:3n-4 and 16:4n-1 account for 12% in BAC, 296 21% in COS, 12% in FRA and 19% in MED while they are only present in trace amounts in other 297 phytoplankton classes. The percentage of 16:4n-3 is above 10% in Chlorophyceae (CHL), 298 Chlorodendrophyceae (CHD), Pyramimonadophyceae (PYR), Prasinophyceae (PRA) and MAM. 299 Percentage of 18:2n-6 is present at 11% in Cyanophyceae (CYA) and 14.5% in Trebouxiophyceae 300 (TRE) while 18:3n-6 is only above 5% in CYA. 18:3n-3 and 18:4n-3 are present in most 301 phytoplankton classes; 18:3n-3 is above 15% in CYA, CHL, TRE, CHD, and Cryptophycea (CRY) 302 while 18:4n-3 is above 12% in PYR, PRA, MAM, CRY PRY, Raphydophyceae (RAP), and PEL. The 303 percentage of 18:5n-3 is especially high in Dinophyceae (DIN) (16%) and present above 5% in PYR, 304 PRA, COC, and PRY. The percentage of 16:4n-3 is above 12% in CHL, CHD, PYR, PRA and MAM, 305 all belonging to the chlorophyta phylum. 20:5n-3 (EPA) ranged between 15% and 20% in PAV, 306 RAP, BAC, COS, FRA, and MED and above 20% in Porphyridophyceae and PIN. 22:6n-3 (DHA) 307 is above 10% in COC, and PRY and above 15% in DIN.

308 Fatty acids associated to heterotrophic organisms

Branched FA (iso15:0, anteiso15:0, iso16:0 and iso17:0) are associated to bacteria (Parrish, 2013;
Volkman et al., 1998; Wilson et al., 2010). The MUFA 20:1n-9, 22:1n-9, 22:1n-7 and 22:1n-11 are
specifically produced by zooplankton such as Calanoids (Brett et al., 2009; Dalsgaard et al., 2003;
Kattner and Hagen, 1995; Mayzaud et al., 2007; Parrish, 2013; Wilson et al., 2010) and are not found
in phytoplankton.

314 **FA-based nutritional quality index**

FA data were used to estimate the nutritional quality of the 1-50 μm SPOM following the FAbased nutritional quality index (NQI) developed by Cañavate (2019). The NQI was calculated as
follow:

NQI =
$$[(15 * DHA + 10 * EPA + 2 * ARA) * 0.8 + (1.8 * \Sigma C18PUFA)] * \log(\frac{n-3}{n-6})$$

DHA, EPA and 20:4n-6 (arachidonic acid, ARA) are expressed as % of TFA. ΣC18PUFA is the
sum of 18:2n-6, 18:3n-3, 18:4n-3 and 18:5n-3 percentages and n-3/n-6 is the ratio of total n-3 PUFA
to total n-6 PUFA.

322 Statistical analysis

323 Principal Component Analysis (PCA) was used to characterize relationships among FA and to discriminate FA profiles according to depth and station. The PCA was performed with 36 FA 324 325 out of 55 analyzed whose average abundance was >0.1% of TFA. Resulting components 1, 2 and 326 3 were analyzed using one-way analysis of variance (ANOVA) to identify differences of FA 327 profiles according to depth intervals (ML, MLD-150 m, 150-300 m) and stations. Categories of 328 branched FA, SAFA, MUFA, short-chain PUFA (SC-PUFA), and LC-PUFA were analyzed by non-329 parametric one-way ANOVA (Krustal-Wallis) with post-hoc analysis to determine significant 330 differences between depth zones (ML, MLD-150 m, 150-300 m). Furthermore, spearman tests were 331 conducted to explore the relationship between some FA and depth. In the surface mixed layer 332 (ML), FA profiles of the 1-50 µm fraction were analyzed by non-parametric one-way ANOVA 333 (Krustal-Wallis) with post-hoc analysis to determine significant differences among stations. Spearman tests were also conducted to explore the relationship between FA profiles and physico-334 335 chemical parameters (Temperature, Salinity, Oxygen, NH4⁺, Si(OH)4, NO3⁻) in the ML. Differences 336 were considered statistically significant if p<0.05. All statistical analyses were performed using 337 Statgraphics Plus statistical software (Manugistics, Rockville, MD, USA).

338 C. RESULTS

339 Hydrological and biogeochemical context

340 During the first half of the MOBYDICK survey, from late February to early March, the MLD 341 was relatively shallow at all stations (Table 1), varying from 63 m at station M1 to 79 m at station 342 M2-1, and it was consistent with late summer conditions (stable upper water column and low 343 wind stress). After a storm event that occurred on the 10th of March, the MLD deepened 344 consistently at stations M4-2 and M3-3 reaching 96 m, and to a lesser extent on the Plateau at 345 station M2-3 (80 m). The depth of Ze was shallow on the plateau (58-64 m) and above the MLD. 346 Ze was deeper at the other stations (89 m at M1 down to 105 m at M3-3) and it was below the 347 MLD indicating no light limitation in the ML.

In the ML, Chl-a exhibited low values at all stations, with 0.27-0.59 μ g.L⁻¹ and 0.14-0.34 μ g.L⁻¹ on- and off-plateau, respectively. At station M2, Chl-a almost doubled between the second (M2-2) and the third visit (M2-3). Silicic acid had variable concentrations in the ML (Table 1). These were the lowest on the plateau at M2-1 and M2-2 (1.4-1.7 μ M), intermediate at M3 (2.3-2.9 μ M) and the highest at M4 (4.8 μ M) and M1 (6.7 μ M). As a comparison, ammonium concentrations, an indicator of heterotrophic excretion, exhibited a reverse trend with highest values on the plateau (0.7-1.1 μ M) decreasing to 0.6-0.7 μ M at M3 and M1, and were the lowest at M4 (0.3-0.5 μ M).

355 Vertical distribution of TFA concentrations

356 Depth weighted average (DWA) of TFA concentrations obtained for three depth intervals 357 (ML, MLD-150 m, 150-300 m) are shown in Figure 2. Considering all stations and visits, TFA concentrations were significantly (p<0.001) higher in the ML (range: 1.4-3.6 µg.L⁻¹) as compared 358 359 to the deeper zones (MLD-150 m and 150-300 m). Although TFA concentrations tended to be lower 360 in the 150-300 m depth interval (range: 0.2-0.7 µg.L⁻¹) compared to the MLD-150 m zone (range: 361 0.6-1.6 µg.L⁻¹), the difference was not significant. TFA concentrations were significantly related to 362 POC (R²=0.91 and p<0.001, n=24) and therefore to biomass variability. Based on the slope of the 363 regression, the average concentration of TFA in biomass was estimated to be 79±5 µg.mgC⁻¹.

364 In the ML, TFA concentrations were not significantly (p>0.05) different between on- and off-Plateau stations despite different temporal trends. On the Plateau, TFA concentrations increased 365 from 1.7 to 2.9 µg.L⁻¹ between stations M2-1 and M2-3. Off the Plateau, TFA concentrations 366 decreased during the course of the survey, from 2.3 to 1.5 µg.L⁻¹ at station M3 and from 2.4 to 1.4 367 µg.L⁻¹ at station M4. The highest TFA concentration in the ML (3.6 µg.L⁻¹) was found at station M1 368 369 downstream of the Plateau. Below the ML, no significant differences (p>0.05) were observed 370 according to station locations. On the Plateau, TFA concentration was higher during the first visit 371 of M2-1 (1.6 µg.L⁻¹) decreasing to 0.6 µg.L⁻¹ at M2-2, ten days later.



Figure 2: Depth weighted average TFA concentrations (μ g.L-1) obtained in three depth intervals (ML, MLD-150m, 150-300m) according the station-visit sampled during the cruise. Also shown, the MLD (m) which was estimated for all stations.

372 FA partitioning according to unsaturation level

A preliminary partitioning of FA was performed according to the unsaturation level as described in Table 2 and considering branched FA, SAFA, MUFA and PUFA. Within PUFA, we further distinguished SC-PUFA with 16 and 18 atoms of carbon from long-chain PUFA (LC-PUFA) containing 20 and 22 atoms of carbon, which included the essential 20:5n-3 and 22:6n-3. 377 Depth weighted average percentages (relative to TFA) of these five groups are shown in Figure 3.





Figure 3: Depth-weighted averaged percentages (relative to TFA) of branched FA (BRCH), saturated FA (SAFA), monounsaturated FA (MUFA), short-chain (C16-C18) and long-chain (C20-C22) polyunsaturated FA (SC-PUFA and LC-PUFA) in three depth intervals (ML, MLD-150, 150-300m).

379 followed by SAFA (21% on average), MUFA (15% on average), and Branched FA (2% on average). 380 Within PUFA, LC-PUFA were dominant (range: 34-44%) and showed increasing and significantly 381 different (p<0.001) proportions according to depth interval, 34% in the ML, 37% in the MLD-150 382 m, and 45% in the 150-300 m. On the opposite, SC-PUFA decreased substantially with depth, from 383 30% in the ML down to 12% in the 150-300 m depth interval, and differences between all depth 384 intervals were significant (p<0.001). For SAFA and MUFA, proportions increased with depth. 385 Differences were significant (p<0.01) only in the 150-300 m compared to the ML and MLD-150 m 386 for SAFA, and in the ML compared to the MLD-150 m and 150-300 m for MUFA.

In the ML, no significant differences were observed for SAFA, MUFA, and Branched between
 on- and off-plateau. LC-PUFA tended to be higher off-plateau compared to on-plateau while SC PUFA tended to be higher on the plateau vs off-plateau. Below the ML, no significant differences

were observed among stations. However, the decrease of SC-PUFA with depth was faster on the
plateau (from 34 to 20%) as compared to off-plateau (from 28 to 24%).

LC-PUFA were dominated by 20:5n-3 and 22:6n-3, both increasing significantly with depth from 13% to 19% for 20:5n-3 and from 18% to 20% for 22:6n-3. SC-PUFA decreased significantly with depth due to the decreases of 18:3n-3, 18:4n-3 and 18:5n-3. In contrast, C16 PUFA associated to diatoms were significantly higher in deep zones (MLD-150m and 150m-300m divisions) than in the ML.

397 **Overall pattern in individual FA abundance**

398 Principal component analysis was applied to 36 FA relative abundances (expressed as % of 399 TFA) by examining 37 cases including FA compositions from 8 stations at various depths and 400 visits. As shown in Figure 4, axis 1 and 2 explained 34.1% and 17.1% of the variability, respectively, 401 and axis 3 12.3% (not shown in Fig. 4). These three axis accounted for >63% of total variability. 402 The positive side of axis 1 was driven by 15:0, 16:0, 18:1n-7, 20:1n-9, 22:1n-9, and 22:1n-7 while 403 17:1n-7, 18:2n-6, 18:3n-6, 18:3n-3, 18:4n-3 and 18:5n-3 were correlated with the negative side of 404 axis 1. The second axis of the PCA was positively associated to diatom markers including 16:2n-405 7, 16:2n-4, 16:3n-4, 16:4n-1 and 20:5n-3 and negatively associated to 18:1n-9. The third axis was 406 positively correlated to 16:4n-3. All correlations previously listed (positive or negative) between 407 FA variables and principal components 1, 2, and 3 were highly significant and had correlation 408 coefficients greater than 0.7. As PC1, PC2, and PC3 revealed good correlations (positive and 409 negative) between FA variables, non-parametric analyses of variance with PC1, PC2, and PC3 as 410 the dependent variables and depth intervals and stations as the independent variables were 411 performed. The analysis showed that differences in FA profiles attributable to depth were highly 412 significant (Figure 5) for both PC1 and 2 with p values <0.001 in both cases. While PC1 value 413 increased steadily with depth zones, PC2 value of the MLD-150 m was higher than in the ML and 414 150-300m zones. While PC1 and PC2 were not significantly different according to stations, PC3 415 allowed significant differentiation between on-plateau station M2 and off-plateau stations M3 and 416 M4 (p<0.01, Figure 5).



Component 1 34.1 %

Figure 4: Correlation circle of 36 fatty acid variables according to the two principal components 1 and 2 obtained from the Principal Component Analysis, which used 37 cases including FA compositions of 1-50 μ m suspended particulate organic matter from 8 stations at various depths. The outer circle represents the unit circle in terms of correlation coefficient (positive or negative along each axis). The closer the FA variables are to the outer circle, the higher the correlation coefficient.



Figure 5: (Left) Non-parametric analysis of variance with PC1 and PC2 as the dependent variables (all stations pooled) and depth intervals as the independent variable; (Right) Non-parametric analysis of variance with PC2 and PC3 as the dependent variables (all depth intervals pooled) and stations as the independent variable. Letters a, b and c indicate statistical differences among conditions.



421 FA variability according to depth and stations

Figure 6: Percentages (relative to TFA) of C18 PUFA (sum of 18:3n-3, 18:4n-3 and 18:5n-3), C16 PUFA (sum of 16:2n-7, 16:2n-4, 16:3n-4 and 16:4n-1), 22:6n-3 and 20:5n-3 according to depth intervals. Letters a, b and c indicate statistical differences among conditions.

422 Furthermore, we explored the relationships between proportions of individual FA and depth 423 (considering all stations together). Depth explained significantly the variability of 17:1n-7 (R²=61%, p<0.0001), 18:2n-6 (R²=30%, p<0.001), 18:3n-6 (R²=37%, p<0.001), 18:3n-3 (R²=44%, 424 p<0.0001), 18:4n-3 (R²=49%, p<0.0001), 18:5n-3 (R²=44.7%, p<0.0001) through negative linear 425 426 regressions. R² values were improved using Log regression for 18:2n-6 (R²=50%, p<0.0001), 18:3n-427 3 (R²=61%, p<0.0001), 18:4n-3 (R²=68%, p<0.0001), 18:5n-3 (R²=68%, p<0.0001). At the opposite, depth explained significantly the variability of 16:0 (R²=68%, p<0.0001), 18:1n-9 (R²=56%, p<0.001), 428 20:1n-9 (R²=78%, p<0.0001), 22:1n-7 (R²=41%, p<0.0001), 22:1n-9 (R²=36%, p<0.001), and 22:5n-3 429 (R²=79%, p<0.0001), through positive linear regressions. The best fitting regressions for 18:4n-3, 430 431 18:5n-3, 20:1n-9 and 22:5n-3 as a function of depth are presented in Figure S1. Variations according 432 to depth concerned also LC-PUFA 20:5n-3 and 22:6n-3, and diatom C16 PUFA (the sum of 16:2n7, 16:2n-4, 16:3n-4 and 16:4n-1) as illustrated in Figure 6 for the three depth intervals considered.
Averaged proportions of 20:5n-3 and diatom C16 PUFA were significantly higher in the MLD150m and 150-300m depth intervals in comparison to the ML. For 22:6n-3, relative proportions
remained high throughout the upper water column (0-300m) ranging from 18 to 20% of TFA, and
were significantly higher in the deepest depth zone (150-300m).



Figure 7: Percentages (relative to TFA) of C18 PUFA (sum of 18:3n-3, 18:4n-3 and 18:5n-3), C16 PUFA (sum of 16:2n-7, 16:2n-4, 16:3n-4 and 16:4n-1), 22:6n-3 and 20:5n-3 according to stations in the ML. Letters a and b indicate statistical differences among conditions.

438 Considering the significant differences between depth intervals, we compared FA profiles among

- 439 stations within depth zones (Figure 7 for the ML). In the ML, the sum of 18:3n-3, 18:4n-3 and 18:5n-
- 440 3 proportions was significantly higher at M1 and M2 stations compared to M3 and M4 stations off
- 441 the plateau. The sum of diatoms C16 PUFA (16:2n-7, 16:2n-4, 16:3n-4 and 16:4n-1) and the

442 proportion of 20:5n-3 were significantly higher at station M4 than at stations M2 and M3. No
443 difference among stations were found within both MLD-150m and 150m-300m zones.

To attempt explaining difference of FA composition among stations, relationships between physico-chemical parameters and FA composition were explored. We only considered regressions with $R^2 > 40\%$ with p value <0.01. Diatoms C16 PUFA and 20:5n-3 proportions were statistically negatively correlated to temperature (R^2 =47% and 44%, respectively, with p <0.01, Figure S2). The sum of 18:3n-3, 18:4n-3 and 18:5n-3 was positively correlated to NH₄+ (R^2 =41%, p<0.01, Figure S2).

449 Diatom C16 PUFA and 20:5n-3



Figure 8: Correlation between diatoms C16 PUFA and 20:5n-3 proportions (% of total fatty acid-TFA) for the mixed layer (ML) (a) and for ML depth (MLD)-300m depth interval (b).

450 As diatoms C16 PUFA and 20:5n-3 were gathered in the PCA, we explored further this 451 relationship in Figure 8. For the ML (Fig. 8a), C16 PUFA and 20:5n-3 appeared highly related 452 regardless of stations (R²=0.945, p<0.001). The slope of the regression indicated an average ratio of 453 20:5n-3 to C16 PUFA of 3.6±0.2, substantially higher than the ratio obtained for four diatom classes 454 (Bacillariophyceae, Cosciondiscophycea, Fragilariophycea, and Mediophycea) gathered from literature (see Table S1), which ranged from 0.9 to 1.6. Considering the MLD-300m depth interval 455 456 (Fig. 8b), the correlation between C16PUFA and 20:5n-3 was lower but still significant (R²=0.487, p<0.001) and the average 20:5n-3 to C16 PUFA ratio decreased to 1.6±0.2. 457



Figure 9: Linear regressions between 18:3n-3 and 22:6n-3 (% total fatty acid-TFA) in the mixed layer for off-plateau stations (M1, M3 and M4) and on-plateau station (M2).

459 From the PCA, 22:6n-3 was poorly correlated to axis 1 (R²=12.5%, p<0.05) and not correlated to axes 2 and 3 when considering all stations and depths together. To further explore the 460 461 relationship with non-diatom markers, we plotted in Figure 9 18:3n-3 against 22:6n-3 proportions in the ML. Although the numbers of data points reduced its statistical strength, separating the 462 samples according to station locations (off-Plateau versus on-Plateau) allowed to obtain a 463 464 significant (R²=0.907, p<0.001) positive relationship between 18:3n-3 and 22:6n-3 for off-plateau 465 stations (M1, M3, and M4). For the on-plateau station M2, the relationship was not significant (p=0.17) although 18:3n-3 and 22:6n-3 were negatively correlated. 466



469 PUFA ratio and 22:5n-3 and 20:1n-9 ($R^2=0.874$ with p<0.0001 and $R^2=0.551$ with p<0.001, 470 respectively) are shown in Figure S3.



Figure 10: Linear regression between 22:6n-3 to n-3 C18 PUFA ratio according to depth (m).

472 FA-based nutritional quality of SPOM

The nutritional quality index (NQI) deduced from FA is presented in Figure 11. NQI ranged from 354 to 477 (average 394±37) and did not vary significantly according to depth zones. It has to be noted that NQI was more variable with increasing depth. When considering together the three depth zones, NQI was slightly higher on-plateau than off-plateau.



Figure 11: FA-based nutritional quality index of SPOM (1-50 μ m) according to stations and depth intervals.

477 D. Discussion

This study investigated the upper water column distribution of 36 FA in SPOM (1-50 μm) collected in the Indian sector of the Southern Ocean in the region of Kerguelen Islands. Principal component analysis (PCA) revealed contrasted FA profiles attributable to depth for both PC1 and 2. The PC3 also revealed significant differences between HNLC stations M3 and M4 and the naturally-iron fertilized station M2. In the following, we discuss first how the post-bloom phytoplanktonic community and depth impact the FA composition of SPOM in terms of SC-PUFA and MUFA. We then consider the potential sources of essential LC-PUFA, 20:5n-3 and 22:6n-3, in the ML and their fate in the upper mesopelagic. Finally, the FA-based nutritional quality of SPOMis discussed according to depth and stations location.

487

87 **Post-bloom phytoplankton community and FA profiles**

488 The MOBYDICK survey took place in late summer/early autumn coinciding with post-bloom 489 conditions on and off the Kerguelen Plateau. Chl-a was low in the ML (<0.6 µg.L-1), and POC and 490 TFA concentrations were <45 mg.L⁻¹ and <4.5 µg.L⁻¹, respectively. The phytoplanktonic 491 community, as described using 18S rDNA amplicon sequencing, pigments, flow cytometry, and 492 microscopic enumeration (Irion et al., 2021, 2020; Lafond et al., 2020), revealed a mixed 493 composition with three main groups, prymnesiophytes, diatoms and prasinophytes showing 494 variable proportions on- and off the Plateau. Nano-phytoplankton (prymnesiophytes) were the 495 most abundant group (37–53% and 59–70% of total Chl-a on- and off-plateau, respectively) 496 followed by diatoms (27-40% and 18-33% of total Chl-a on- and off-plateau, respectively), and 497 finally by pico-phytoplankton (prasinophytes) (Irion et al., 2021, 2020). This latter group was 498 found in low proportions at all stations except on the plateau where it reached up to 16% of total 499 Chl-a at the end of the survey (M2-3) (Irion et al., 2020). The mixed composition of the 500 phytoplankton community was well reflected in the FA profile of SPOM, especially when 501 considering chloroplastic C16 and n-3 C18 PUFA (Alonso et al., 1998; Guschina and Harwood, 502 2006), which derive mainly from autotrophic biomass. n-3 C18 PUFA (sum of 18:3n-3, 18:4n-3, and 503 18:5n-3) are characteristic of non-diatom phytoplankton including prymnesiophytes (24% of n-3 504 C18 PUFA) and prasinophytes (42% n-3 C18 PUFA), and were particularly abundant in the ML representing between 17 to 32% of TFA. n-3 C18 PUFA in the ML were significantly (p<0.05) 505 506 higher on the Plateau (30±4% on average at station M2) and downstream (26% at M1) in 507 comparison to HNLC off-plateau stations (19±4% on average at stations M3 and M4). On the 508 Plateau, n-3 C18 PUFA increased from 22% (M2-1) to 32% (M2-2 and M2-3) during the survey 509 probably illustrating the increasing proportion of Prasinophytes.

510 For diatoms, their abundance could be identified using C16 PUFA (sum of 16:2n-7, 16:2n-4, 511 16:3n-4 and 16:4n-1), which are highly specific to this phylum with an averaged proportion of 512 16±4% (see Table S1 for details). Diatom C16 PUFA were found at all stations in the ML. C16 PUFA 513 were the lowest on the Plateau at station M2 (1.2±0.2% on average) and upstream at station M3 514 $(1.3\pm0.2\%)$ on average) and were significantly (p<0.05) higher upstream at station M4 (2.3\pm0.6\%) on 515 average) compared to the other stations. Variability of C16 PUFA according to stations in the ML 516 could be compared to estimates of diatom abundance deduced from pigments (Irion et al., 2020), 517 diatom biomass contribution to POC as well as BSi (Lafond et al., 2020). For off-plateau stations 518 (M1, M3, and M4), C16 PUFA matched the trends in BSi concentrations (2.2 μ mol.L⁻¹ at M4 > 1.0 519 μ mol.L⁻¹ at M1 > 0.4 μ mol.L⁻¹ at M3) as well as diatom contribution to POC (24±6% at M4 > 16±1%) 520 at M1 > $6\pm 2\%$ at M3). On the Plateau, differences arose and C16 PUFA proportions were the lowest 521 while diatoms were found to be more abundant, 27-40% of total Chl-a (Irion et al., 2020) and 17 to 522 43% of POC (Lafond et al., 2020). This was especially the case at the last visit on the Plateau (station 523 M2-3), where a bloom of Corethron inerme was observed (Lafond et al., 2020) which contributed to 524 an increase of TFA and POC concentrations (2.1 to 2.9 µg.L⁻¹ and 27 to 40 µg.L⁻¹, respectively). In 525 contrast, this effect was not reflected in C16 PUFA, whose proportions stayed relatively stable 526 between station M2-2 (1.0%) and M2-3 (1.2%). This pattern could indicate that this late summer 527 diatom bloom was probably poor in chloroplastic C16 PUFA.

528

Changes in FA profiles with depth

529 Below the ML, TFA and POC decreased rapidly and substantial changes were evidenced at all 530 stations with a consistent decrease in n-3 C18 PUFA (from $24 \pm 6\%$ in the ML to $16 \pm 6\%$ in the 531 MLD-150m down to 5 ± 1% in the 150-300 m depth zones) and an increase in C16 PUFA (from 1.6 532 \pm 0.6% in the ML to 3.5 \pm 1.1% in the MLD-150m and to 3.1 \pm 1.4% in the 150-300 m intervals). The observed pattern in phytoplankton-derived FA was in agreement with the vertical distribution of 533 pigments showing a rapid decrease in prymnesiophytes pigments below the ML and an 534 535 increasing contribution of diatom pigments, the latter representing up to 77-96% of total Chl-a at 536 250 m depth (Irion et al., 2021). In addition, we noted variable vertical trends of n-3 C18 PUFA 537 and C16 PUFA as observed with pigments (Irion et al., 2021). On the Plateau and downstream 538 (Station M2 and M1) n-3 C18 PUFA decreased rapidly below the MLD, while at station M3 C18 539 PUFA remained relatively high in the MLD-150m depth interval (22% on average) and decreased 540 only below 150 m (6% on average). For C16 PUFA and diatoms, the increase with depth was

541 particularly rapid on the Plateau at the first two visits (1.5 to 5.5% at M2-1 and 1.0 to 4.3% at M2-542 2 between the ML and the MLD-150m depth zone) indicating an important accumulation of 543 diatoms just below the MLD. Considering that biomass and associated FA increased below the 544 ML as a result of vertical transfer from the surface, the observed trends in n-3 C18 PUFA and C16 PUFA proportions may be consistent with a more efficient export of diatoms in comparison to 545 546 nano- and pico-phytoplankton below the ML. This feature is commonly observed in the SO (Salter 547 et al., 2012, 2007) and was documented in details on the Plateau (Blain et al., 2021; Rembauville et 548 al., 2015a, 2015b) and also in the surrounding HNLC areas (Rembauville et al., 2017).

549 Unexpectedly and along with n-3 C18 PUFA, the MUFA 17:1n-7 decreased linearly with 550 depth. As its distribution was concomitant with the one of chloroplastic n-3 C18 PUFA, it suggested its production was mostly autotrophic and occurred in the ML. 17:1n-7 is not listed in 551 552 the meta-analysis of Jónasdóttir (2019) and Cañavate (2019) and is only rarely cited in papers 553 analyzing FA in phytoplankton species. The cyanobacteria Synechococcus elongates, Microcystis 554 aeruginosa and Anabaena variabilis were reported to contain respectively 0.6%, 0.5% and 1.3 of 555 17:1n-7 (Bec et al., 2006; Martin-Creuzburg et al., 2008). Interestingly, the heterotrophic 556 nanoflagellates Paraphysomonas sp., concentrated this MUFA at 3.4% and 1.5%, when fed S. 557 elongates, M. aeruginosa, respectively (Bec et al., 2006). Indeed, these planktonic groups were 558 observed during the MOBYDICK survey. Heterotrophic nanoflagellates, pico and nano-plankton 559 varied between 3.3 and 4.7 106 cells.L-1, 0.8 and 2.0 106 cells.L-1, and 73 and 111 103 cells.L-1 560 respectively, in the ML (Christaki et al., 2020). However, while heterotrophic nanoflagellates were more abundant at M2 than at off-plateau stations (M3 and M4) (Christaki et al., 2020), no 561 562 significant difference was observed according to stations for both concentration (in µg.L-1) and 563 relative proportion of 17:1n-7.

The long chain MUFA 20:1n-9, 22:1n-7 and 22:1n-9, which are mainly produced by zooplankton such as calanoid copepods and other crustaceans (Brett et al., 2009; Dalsgaard et al., 2003; Kattner and Hagen, 1995; Mayzaud et al., 2007; Parrish, 2013; Wilson et al., 2010) can be used to detect potential interactions with zooplankton via the production of fecal pellets and their presence into SPOM (Mayzaud et al., 2007; Sheridan et al., 2002; Wilson et al., 2010). Fecal pellets

569 are known to play an important role in the vertical transfer of OM in the Kerguelen region 570 (Ebersbach and Trull, 2008; Laurenceau-Cornec et al., 2015), particularly on the Plateau during the post-bloom period (Rembauville et al., 2015a). In our samples, long chain MUFA 20:1n-9, 22:1n-7 571 572 and 22:1n-9 were detected at all stations, their proportions were low in the ML (sum of 20:1n-9, 573 22:1n-7, and 22:1n-9 ranging between 0.11 and 0.25%) and increased linearly with depth (R²=0.683, 574 p<0.0001) reaching the highest values in the 150-300m depth zone (range: 0.21-0.61%). Although 575 particularly low, these proportions still appeared comparable to those measured in fecal pellets 576 of copepods and euphausiids (range: 1.3-2.2%) sampled in the Subantarctic zone of the SO 577 (Mayzaud et al., 2007). This could indicate that a fraction of the SPOM sampled during the post-578 bloom period was composed of fecal material excreted by zooplankton, which increased with 579 depth. Moreover, and as outlined in the PCA, the PUFA 22:5n-3 was clearly associated with long 580 chain MUFA. Indeed, a similar linear increase of 22:5n-3 with depth ($R^2=0.79$, p<0.001) was 581 reported suggesting a common origin. This PUFA can be formed from the 20:5n-3 through 582 elongation and could be a synthesis intermediate of 22:6n-3 in crustaceans including copepods 583 (Kabeya et al., 2021; Monroig and Kabeya, 2018; Nielsen et al., 2019). This PUFA was found in 584 three zooplankton species Themisto libellula, Calanus marshallae/glacialis (Calanus spp.), and 585 Thysanoessa raschii collected in the Bering Sea and ranged between 0.5-0.6% of TFA (Wang et al., 2015). As part of MOBYDICK, 22:5n-3 accounted on average for 0.77% of total phospholipids 586 587 extracted from 70 zooplankton samples (Puccinelli E. in prep) and its abundance in SPOM reached 588 1.3±0.1% on average in the 150-300m depth interval.

589 Origin and fate of 20:5n-3

590 Synthesis of the LC-PUFA 20:5n-3 is assumed to occur essentially via autotrophy and can be 591 performed by several phytoplankton phyla such as Diatoms, ochrophytes, haptophytes, and 592 rhodophytes (see Table S1 for details). In the ocean and in particular the SO, the main suppliers 593 of 20:5n-3 are generally considered to be diatoms (Ericson et al., 2018; Hellessey et al., 2020). In 594 the case of MOBYDICK and for the Kerguelen area, this assumption seems to be true. 20:5n-3 was 595 clearly associated with diatom C16 PUFA in the PCA and furthermore in the ML, proportions of 596 20:5n-3 appeared tightly related to C16 PUFA through a positive linear correlation (R²=0.945,

597 p<0.05) regardless of stations. Abundance of 20:5n-3 accounted for a significant contribution to 598 TFA in the ML (13±2% on average), highest proportions were found at station M4 (15±2% on 599 average), significantly (p<0.001) different in comparison to stations M2 and M3 (12±1% on average 600 at both stations) and to a lesser extent to station M1 (14% on average). Measured proportions in 601 1-50 µm SPOM were comparable to literature data for diatoms ranging from 15 to 20% of 20:5n-3 602 (see details in Table S1). However, the literature range corresponds to diatoms monoculture and 603 differs from the MOBYDICK situation. Because diatoms were a minor group in the ML (18-40% 604 of total Chl-a) (Irion et al., 2020), proportions of 20:5n-3 in SPOM are expected to be lower 605 compared to literature data if only provided by diatoms. Based on the highly significant linear 606 relationship (p<0.001) between 20:5n-3 and diatoms C16 PUFA in the ML, we calculated a 20:5n-607 3/C16 PUFA ratio of 3.6±0.2, which appeared to be ~2 fold higher compared to literature data 608 (average of 1.5 calculated from Table S1). This difference suggests that the diatom community of 609 the Kerguelen region, both on and off-Plateau, had higher proportions of 20:5n-3 than cultured 610 diatoms. Such a case was previously reported by Vaezi et al. (2013), who found that the cold-water 611 Fragilariopsis cylindrus was able to produce up to 31.4% of 20:5n-3, approximately twice the 612 average value usually found in the literature (Table S1). Additionally, we reported a significant 613 negative regression between 20:5n-3 and temperature in the ML (R²=44%, p<0.01). Indeed, 20:5n-614 3 proportion was the highest at station M4 where temperature was the lowest. Based on literature 615 data, Hixson and Arts (2016) established a negative linear regression between temperature and 616 20:5n-3 in diatoms. Thus, it can be speculated that the high 20:5n-3 content in diatoms of the 617 Kerguelen region partially reflected homeoviscous adaptation to low temperatures.

In the upper mesopelagic, relative proportions of 20:5n-3 increased, reaching 17±2% in the 618 619 MLD-150m zone and 19±3% in the 150-300m zone, and were significantly (p<0.001) different 620 compared to the ML. Linear relationship between C16 PUFA and 20:5n-3 was still significant 621 below the MLD (R²=0.487, p<0.001) with an average ratio 20:5n-3/C16 PUFA of 1.3. This ratio was 622 lower than in the ML and closer to the literature value (1.5, calculated from Table S1). This 623 suggests that the vertical distribution of 20:5n-3 was still mostly determined by the fate of diatoms 624 in the mesopelagic with no significant difference among stations. Accordingly, trends in 20:5n-3 625 proportions were likely a result of the increasing proportion of diatoms observed below the MLD.

626 However, according to the low 20:5n-3/C16 PUFA ratio, the FA composition of diatoms has 627 probably been modified with depth, and this could be related to several factors. For instance, 628 diatoms assemblages exhibited consistent variations according to stations and depth (Lafond et 629 al., 2020). This was particularly the case at the plateau station M2, where in the ML diatom biomass 630 was dominated by the large centric Corethron inerme (59 to 83% of diatom biomass), while below 631 the ML, Eucampia antarctica at the first visit (M2-1), and Chaetoceros resting spores and Chaetoceros 632 spp. at the following visits (M2-2 and M2-3) were the dominant contributors (Lafond et al., 2020). 633 At other stations, the community structure was more diverse and included Actinocyclus spp., 634 Thalassiosira spp., Chaetoceros atlanticus and Chaetoceros resting spores (Lafond et al., 2020). 635 Considering that the amount of 20:5n-3 produced varies at a species level, diatom diversity were 636 thus likely to influence the FA content available. Other factors could also involve the physiological 637 state of diatom cells, which greatly evolved in the upper mesopelagic zone as a consequence of 638 grazing, parasitic infection and mortality (Lafond et al., 2020; Sassenhagen et al., 2020). Indeed, 639 abundance of detrital cells in the form of empty, broken or crushed frustules rapidly increased 640 below the ML and this was particularly pronounced at off-plateau stations M1, M3, and M4 in 641 comparison to on-plateau station M2, where up to 48% of cells were still intact between 175-200 642 m depth (Lafond et al., 2020). Finally, the relative proportions of 20:5n-3 as well as 20:5n-3/C16 643 PUFA ratio could have been modified as a result of the increasing proportion of fecal material 644 observed in the mesopelagic zone. The FA composition of fecal pellets reflects the composition of 645 the zooplankton organism (Hamm et al., 2001; Mayzaud et al., 2007) and can contain high and 646 variable proportions of 20:5n-3 (up to 25-30%) as observed in krill (Hellessey et al., 2020), with 647 substantial differences in comparison to the phytoplankton food source.

648

Origin and fate of 22:6n-3

The LC-PUFA 22:6n-3 is also produced at the basis of the marine food web by unicellular eukaryotic organisms (Dalsgaard et al., 2003; Parrish, 2013) and it is produced by several phytoplankton taxa (as detailed in Table S1). The taxa with the highest 22:6n-3 proportions, grown in autotrophic conditions, include Dinophyceae (17.5% in average), Coccolithophyceae (11.6% in average) and Prymnesiophyceae (10.7% in average) and exclude diatoms, which produce only 654 small amounts of 22:6n-3 (<2.5%). In this study and considering first the ML, where autotrophic 655 production was likely dominant, the 1-50 µm SPOM was found to be rich in 22:6n-3 at all stations representing 18±1% on average. The highest proportions were found off the Plateau at stations 656 M1 and M3, with 19.1% and 19.5±0.5%, respectively, which were slightly higher to the upstream 657 station M4 (17.0±1.0% in average) and the on-plateau station M2 (16.4±0.8% in average). 658 659 According to the composition of the post-bloom phytoplankton community (Irion et al., 2021, 660 2020), it can be hypothesized that Prymnesiophycea, as the dominant group, were the most 661 important suppliers of 22:6n-3 in the ML both on and off-Plateau. Autotrophic dinoflagellates 662 (Dinophyceae), which are generally considered as important producers in the SO (Dalsgaard et 663 al., 2003; Ericson et al., 2018; Falk-Petersen et al., 2000) were likely marginally involved as their 664 contribution to the phytoplankton community was very modest (Irion et al., 2020). In addition to Prymnesiophyceae, contribution of pico-phytoplankton (Prasinophyceae) to 22:6n-3 proportions 665 was also probable, especially on the Plateau where they were more abundant. Prasinophyceae are 666 667 able to produce 22:6n-3 but in much lower proportions (7.0%) compared to Prymnesiophyceae. In 668 order to test the assumption that Prasinophyceae and Prymnesiophyceae contributed to 22:6n-3 669 production, we considered the relationship between 22:6n-3 and 18:3n-3 in the ML. 22:6n-3 670 appeared strongly related to 18:3n-3 only at off plateau stations (M1, M3, and M4) through a 671 positive correlation (R²=0.907, p<0.05). The slope of the regression corresponding to the averaged 672 22:6n-3/18:3n-3 ratio found in off Plateau SPOM was 7.2 and was substantially higher than the 673 ratio deduced from literature data for Prymnesiophycea (2.1) and Prasinophycea (0.6). The fact 674 that the SPOM 22:6n-3/18:3n-3 ratio was closer to that of Prymnesiophyceae may indicate that 675 22:6n-3 was at least partially produced by Prymnesiophyceae rather than Prasinophyceae at these 676 off-Plateau stations. The situation was markedly different on the Plateau at station M2, where 677 22:6n-3 and 18:3n-3 were negatively related but not significantly, suggesting a more complex 678 pattern. The observed trend in SPOM could be consistent with a multiple origin of 22:6n-3 with 679 some from Prymnesiophyceae having a high 22:6n-3/18:3n-3 ratio and some from Prasinophyceae 680 having a low 22:6n-3/18:3n-3 ratio. Different contributions of Prymnesiophycea and 681 Prasinophyceae on and off Plateau seemed to be confirmed by the decrease of 22:6n-3/18:3n-3 ratio 682 in SPOM observed at station M2 according to visits (7.9 at M2-1, 5.7 at M2-2, and 5.1 at M2-3)

which fits with the increasing proportion of Prasinophyceae deduced from pigments and reachingup to 16% of total Chl-a at M2-3 (Irion et al., 2020).

685 Considering Prymnesiophycea, it is worth noting that this group was mostly composed by a single species, Phaeocystis antarctica (Irion et al., 2020), whose FA profile remains to be 686 687 documented. As a comparison, data available for other species of *Phaeocystis* (pouchetti and globosa) 688 exhibit contrasted FA profiles, with some studies reporting only trace or minor amounts of 22:6n-689 3 (maximum 6.2% TFA) (Claustre et al., 1990; Nichols et al., 1991; Skerratt et al., 1995), while others 690 reported much higher proportions, up to 15% (Hamm et al., 2001; Sargent et al., 1985; Virtue et 691 al., 1993). In our case, Phaeocystis antarctica was necessarily high in 22:6n-3 to match the high 692 proportions observed in the ML (18±1% on average) and further work is needed to confirm this 693 hypothesis.

694 In addition to autotrophic origin, trophic upgrading by heterotrophic protists such as 695 dinoflagellates and ciliates, which composed the microzooplankton, may also have contributed to 696 the observed 22:6n-3 content in the ML (Broglio et al., 2003; Lund et al., 2008). Heterotrophic 697 dinoflagellates and ciliates were documented at both on and off-plateau stations and ranged 698 between 0.29 and 2.3 103 cells L-1 (Christaki et al., 2021). Dinoflagellates, dominated by 699 Gymnodinium sp., were more abundant than ciliates at all stations and grazed actively on 700 phytoplankton with grazing rates exceeding phytoplankton growth rates (Christaki et al., 2021). 701 Heterotrophic dinoflagellates can contain especially high proportions of 22:6n-3, >40% as reported 702 by Lim et al. (2020) for Gymnodinium smaydae, and up to >50% for Crypthecodinium cohnii (Jiang 703 and Chen, 2000). Interestingly for this latter species, 22:6n-3 proportions were dependent on 704 cultivation temperature, increasing from 17% to 53% with decreasing temperature from 30°C to 705 15°C (Jiang and Chen, 2000). Considering the low temperature environment of the Kerguelen 706 region, this may have favored high 22:6n-3 proportions in heterotrophic protists. Another 707 characteristic of heterotrophic dinoflagellates is their low content in chloroplastic n-3 C18 PUFA 708 and as a consequence, their high 22:6n-3/n-3 C18 PUFA ratio (Lim et al., 2020; Lund et al., 2008). 709 As an example, *Gymnodinium smaydae* had a 22:6n-3/18:3n-3 ratio >80, much higher than the ratio 710 of its prey *Heterocapsa rotundata* of 5.4 (Lim et al., 2020). This may be one explanation for the higher 22:6n-3/18:3n-3 ratio found in the ML (7.2) in comparison to autotrophic producers (0.6-2.1) and
could be consistent with a small contribution (<10%) of heterotrophic protists to 22:6n-3 content.

713 Below the ML in the upper mesopelagic, 22:6n-3 concentrations (in µg.L-1) decreased 714 substantially and similarly to TFA concentrations (R²=0.975, p<0.05) it was maintained at high 715 relative proportions with 17±2% in the MLD-150m depth interval comparable to the ML, and up 716 to 20±2% in the 150-300m depth interval, significantly (p<0.001) higher in comparison to the upper 717 layers. This vertical trend was in a marked contrast compared to n-3 C18 PUFA, which decreased 718 substantially both in concentration and proportion below the ML. This contrasted pattern was 719 initially outlined by the PCA, where 22:6n-3 was not correlated to any of the three axes and furthermore, not associated to n-3 C18 PUFA when considering the whole dataset. These different 720 721 trends clearly indicate that the fate of 22:6n-3 in the mesopelagic could not be governed by the 722 vertical transfer of non-diatom phytoplankton which was initially involved in its production in 723 the ML, and most probably relied on another pathway. This pathway is further supported by the 724 linear increase of 22:6n-3 to n-3 C18 PUFA ratio according to depth (R²=0.726, p<0.0001) indicating 725 a progressive enrichment of 22:6n-3 compared to n-3 C18 PUFA in SPOM between the surface and 726 300 m depth. Moreover, this increase correlates fairly well with the heterotrophic FA 22:5n-3 (R²=0.874, p<0.0001) and also but to a lesser extent with the MUFA 20:1n-9 (R²=0.551, p<0.001). 727 728 Variations of 22:5n-3 and 20:1n-9 were attributed earlier in the discussion to an increasing 729 abundance of fecal material at depth. Accordingly, it can be postulated that 22:6n-3 in SPOM may 730 be at least partially associated with this fecal material in the upper mesopelagic and thus its fate 731 may be controlled by interactions with the heterotrophic food web. It is worth noting that 732 correlations were made regardless of stations, indicating that this potential transfer pathway 733 seemed to affect both iron-fertilized and HNLC regions. Such an association of 22:6n-3 with fecal 734 material has already been documented in some regions of the world ocean, including the high 735 latitude North Pacific where fecal OM and high proportions of 22:6n-3 (range 8-17% of TFA) were 736 evidenced throughout the water column (Sheridan et al., 2002; Wilson et al., 2010) and also in the 737 Arctic, where it was demonstrated that 22:6n-3 rich OM (~20% of TFA) derived from Phaeocystis 738 pouchetii, was exported below the euphotic zone via krill fecal strings (Hamm et al., 2001).

739 In the present case, the processing of 22:6n-3 from phytoplankton to suspended fecal material 740 is likely to be complex and could involve a number of heterotrophic intermediates. Among 741 grazers, mesozooplankton such as copepods and other large size crustacean zooplankton require 742 to be considered as C20-C22 MUFA are representative of fecal material excreted by these 743 organisms (Brett et al., 2009; Dalsgaard et al., 2003; Kattner and Hagen, 1995; Mayzaud et al., 2007; 744 Parrish, 2013; Wilson et al., 2010). Mesozooplankton were documented at all stations and 745 exhibited significant variations according to sites and visits from 207 ind.m⁻³ at M2-1 to 1636 746 ind.m⁻³ at M4–1 (Christaki et al., 2021). Moreover, grazing experiments showed that copepods 747 were not grazing directly on phytoplankton but were feeding primarily on microzooplankton 748 controlling their abundance during the post-bloom period (Christaki et al., 2021). This feeding 749 strategy on heterotrophic protists as well as the selective storage of 22:6n-3 as an essential LC-750 PUFA for mesozooplankton (Arendt et al., 2005) may have favoured its accumulation relative to 751 n-3 C18 PUFA and as a result, impacted the composition of fecal material produced by these 752 organisms. In addition to mesozooplankton, other grazers such as salps may also be involved in 753 the channelling of 22:6n-3. Salps are important grazers of small phytoplankton (Moline et al., 2004) 754 and flagellates (Pakhomov and Hunt, 2017; von Harbou et al., 2011), and produce easily 755 fragmented tabular fecal pellets (Iversen et al., 2017). Salps population (Salpa thompsoni) were 756 particularly abundant on the Plateau at station M2 and downstream at station M1 representing 757 up to 40% of total micronekton biomass (Henschke et al., 2021).

758 Nutritional food value according to production regimes

759 The nutritional quality of OM as food for higher trophic levels is partly determined by the 760 abundance of the essential omega-3 LC-PUFA 20:5n-3 and 22:6n-3 and the essential omega-6 761 20:4n-6. In the present study, LC-PUFA essentially composed of omega-3 LC-PUFA were found 762 in high proportions, making up 27-44% of TFA, indicating a high nutritional value of SPOM both 763 in the ML and upper mesopelagic. Following the approach developed by Cañavate et al. (2019), 764 the NQI of SPOM was also high and ranged from 354 to 403 in the ML and from 333 to 477 in the 765 MLD-300m depth interval. As a comparison, much lower NQI were obtained in the estuary of 766 Guadalquivir River, with values always <200 (Cañavate et al., 2021), in the northern part of the

767 Indian sector of the ACC, with NQI of 193 and 242 in surface SPOM of the subtropical and 768 Subantarctic zone, respectively (Mayzaud et al., 2007), in the Northern Pacific, NQI was <300 in 769 >51 µm SPOM of the Subarctic area (Wilson et al., 2010) as well as in the coastal Antarctic zone, 770 NQI was <260 for a diatom sea-ice community of McMurdo Sound (Nichols et al., 1993). This 771 comparison confirms that quality of SPOM in the Kerguelen area was especially high in the post-772 bloom period. High NQI values were observed over the whole area and seemed unrelated to the 773 contrasted productivity regimes that took place earlier in the season during the bloom. In the ML, 774 the high nutritional value of SPOM was mainly attributable to the mixed phytoplanktonic 775 community relying both on diatoms producing 20:5n-3 and small phytoplankton producing 776 22:6n-3. This mixed production of both LC-PUFA substantially promoted the nutritional quality 777 of SPOM and offered a valuable source of food for secondary consumers that could be of 778 importance for the ecosystem functioning.

779 In the upper mesopelagic, LC-PUFA in SPOM were maintained at similar, and even higher, 780 proportions in comparison to the ML (34±2% and 40±3% on average in the MLD-150m and 150-781 300m depth intervals, respectively). Similarly, NQI remained high until 300 m depth (333-477) 782 even though TFA concentrations in seawater (in µg.L-1) decreased substantially with depth. This 783 preservation of food quality in the upper mesopelagic was rather unexpected because LC-PUFA 784 are generally considered as a labile fraction of OM that could be easily degraded/consumed by 785 heterotrophic organisms during their transfer from the surface (Budge et al., 2006; Conte et al., 786 2003, 1995; Wakeham, 1995). This is illustrated by the low NQI values that can be deduced from 787 sediment trap data, with values between 90 to 180 in trap samples collected between 50-100 m depth in the North Atlantic (Budge and Parrish, 1998) and NQI <30 in samples collected at 300 m 788 789 depth in Breid Bay, Antarctica (Hayakawa et al., 1996). For the Kerguelen region, sediment trap 790 data obtained on the Plateau at station M2 (Rembauville et al., 2018) do not allow for NQI 791 estimates, however considering LC-PUFA concentrations per unit mass of OC, ranging from 0.03 792 to 3.4 µg.mgOC⁻¹ in the trap samples and from 14 to 44 µg.mgOC⁻¹ in our SPOM samples, it 793 appears that sinking material from traps was of much lower quality than suspended particles 794 collected on and off the Plateau. In the present case, the high quality of SPOM in the upper 795 mesopelagic could represent a valuable nutritional option for heterotrophic organisms residing in the part of the water column and feeding on suspended particles such as zooplankton,coprophages, detritivores, and bacteria.

798 E. CONCLUSION

799 In this study, FA distribution including essential LC-PUFA 20:5n-3 and 22:6n-3 was 800 documented in the upper water column of the Kerguelen Islands Region several weeks after the bloom period in late summer/early autumn. Our results illustrate the important role of the mixed 801 802 phytoplankton community, composed of nano- and pico-size phytoplankton and diatoms, in 803 providing high proportions of both LC-PUFA to surface waters. Diatoms were identified as the likely suppliers of 20:5n-3 and small phytoplankton (prymnesiophytes and prasinophytes) 804 805 supplying 22:6n-3. Elevated proportions are unrelated to past productivity regimes and were 806 observed both inside the iron-fertilized area on the Kerguelen Plateau (station M2) and 807 downstream (station M1), as well as outside in HNLC waters located upstream of the Plateau 808 (stations M3 and M4). As a consequence, and despite reduced productivity and biomass 809 abundance during the post-bloom period, phytoplankton-derived OM reveal an especially high 810 nutritional value that could be relevant for secondary consumers such as micro- and 811 mesozooplankton.

812 In the upper mesopelagic, both LC-PUFA were maintained at high relative proportions in 813 SPOM suggesting an efficient and probably fast vertical transfer from the surface. This vertical 814 transfer seems to proceed via distinct pathways according to LC-PUFA. For 20:5n-3, its fate in the 815 mesopelagic seems to mainly follow the one of diatoms and could be associated with the export 816 of large intact cells, diatoms aggregates as well as resting spores. For 22:6n-3, its fate with depth 817 appears not to be simply related to small phytoplankton but is most likely due to a complex 818 channeling through the heterotrophic food web resulting in its association with fecal material at 819 depth. Processing of 22:6n-3 could involve different intermediates such as heterotrophic protists 820 with dinoflagellates and ciliates (microzooplankton), which probably grazed on small 821 phytoplankton as well as larger zooplankton organisms such as copepods and salps, which 822 presumably fed on microzooplankton and produced fecal pellets rich in 22:6n-3. The complex 823 pathway of 22:6n-3 cannot be easily resolved using only SPOM data and additional evidences are 824 needed to better understand this pattern. Despite these contrasted pathways, FA-based 825 nutritional quality of SPOM remained especially high in the upper mesopelagic and could 826 represent an interesting food source for suspension feeders residing there. Finally, these results 827 provide a detailed framework of FA abundances that can be used in trophic and diet studies 828 dedicated to the Subantarctic islands.

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