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Rontani, J-F

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6 Biotic and abiotic degradation of suspended particulate lipids along a transect in the Chukchi Sea Rontani, Jean-François^{1*}, Smik, Lukas², Ha, Sun-Yong³, Min, Jun-oh^{3,4}, Belt, Simon T.² ¹ Aix Marseille Université, Université de Toulon, CNRS/INSU/IRD, Mediterranean Institute of Oceanography (MIO), UM 110, 13288 Marseille, France. ² Biogeochemistry Research Centre, School of Geography, Earth and Environmental Sciences, University of Plymouth, Drake Circus, Plymouth, Devon PL4 8AA, UK. ³ Division of Ocean Sciences, Korea Polar Research Institute, 26 Songdomirae-ro, Incheon, 21990, Republic of Korea. ⁴ Department of Marine Science and Convergence Technology, Hanyang University, 55 Hanyangdaehak-ro, Ansan, 15588, Republic of Korea. * Corresponding author. Tel.: +33-4-86-09-06-02; fax: +33-4-91-82-96-41. E-mail address: jean-francois.rontani@mio.osupytheas.fr (J.-F. Rontani)

Abstract. Lipids and their degradation products were investigated in samples of suspended particulate matter (SPM) collected in summer 2015 from surface waters of a South-North transect (ca. 65-81°N) of the Chukchi Sea. This material appeared to be composed mainly of diatoms (dominated by Thalassiosirales) and copepod faecal pellet debris. The high solar irradiances measured in the surface waters (up to 500 W m⁻²) favour chlorophyll (sensitizer) photobleaching at the expense of Type II photosensitized oxidation of unsaturated lipid components of phyto- and zooplankton (photodynamic effect). The weak photoreactivity of wax esters of herbivorous zooplankton in these SPM samples contrasts with previous observations of strong photooxidation of these compounds in sinking particles, which suggests that the photodynamic effect should be favoured in large faecal pellets of herbivorous copepods sinking quickly in weakly irradiated zones. Autoxidation (free radical induced oxidation) processes operating in all oxic environments appeared to be particularly efficient in faecal pellets of omnivorous and carnivorous zooplankton and limited in those of herbivorous origin. These differences were attributed to the consumption of algal antioxidants (such as mycosporine-like amino acids and carotenoids) during the diet of omnivorous and carnivorous copepods, favouring the involvement of free radical oxidation processes.

Key words: Chukchi Sea; SPM; Lipids; Biotic and abiotic oxidation.

1. Introduction

The Arctic and Antarctic represent the most sensitive regions on Earth in terms of responses to climate change induced by global warming (IPCC, 2013). Sea ice is a particularly characteristic feature of the polar regions and has various roles in controlling local and global climate. During recent decades, there have been dramatic changes in Arctic sea ice cover and thickness (Comiso et al., 2008), and these likely impact the local, regional and global climate through complex coupling mechanisms within the Earth system. Sea ice is also critical for polar ecosystems, providing a host for phototrophic organisms at the base of the food chain, a source of nutrients for other phototrophs, which provide food to primary consumers, and a physical platform for apex marine predators such as polar bears.

Sea ice undergoes large seasonal and inter-annual variation. Having reached its maximum extent during late winter, the region of open ocean defined by the retreating ice edge in spring/summer is frequently referred to as the marginal ice zone (MIZ). The MIZ plays host to major spring phytoplankton blooms, stimulated by higher irradiance, elevated nutrient supply and water column stratification. Such spring primary production (PP) in the MIZ drives the Arctic ecosystem and also contributes significantly to the overall carbon cycle by exerting control over atmospheric CO₂ drawdown and subsequent carbon sequestration in marine sediments (Wassman et al., 2008). Increased spatial extent and duration of open waters due to sea ice loss favors phototrophic activity, and likely explains some of the recent increase in PP in the Arctic (Arrigo et al., 2015).

Some of the most dramatic sea ice decline is evident in the MIZ of the Barents Sea, the Chukchi Sea, the Sea of Okhotsk and Baffin Bay (Onarheim et al., 2018), with impacts on both climate and ecology (Serreze et al., 2007; Leu et al., 2011; Post et al., 2013; Renaut et al., 2018). Indeed, the Chukchi Sea is generally considered as one of the most biologically productive regions in the world's oceans (Springer and McRoy, 1993). The hydrography of the Chukchi

Sea is strongly influenced by the advection of water masses from the Bering Sea (Grebmeier et al., 2006), which is particularly strong in summer (Hunt et al., 2013). This inflow of warmer waters has increased in recent times as a response to global warming, with consequences for the spatial coverage of sea ice and timing of retreat (Grebmeier et al., 2006). It is thus very important to study the lower trophic levels in this region, in order to improve our understanding of the effects of sea ice reduction on marine ecosystems (Abe et al., 2020).

In the Southern Chukchi Sea, low grazing pressure (resulting from a temporal mismatch between zooplankton and phytoplankton production; Kitamura et al., 2017), fast sinking rates of large diatoms, and shallow bottom depths (Grebmeier et al., 1988, 2006a), result in strong pelagic-benthic coupling leading to increased efficiency of carbon sinking (Bates, 2006; Chen and Borges, 2009). However, these ecosystems are currently undergoing rapid and profound changes due to the effects of climate change, which can modify the ecosystem structure from pelagic-benthic to pelagic-pelagic type structures (Piepenburg, 2005). In the present work, we aimed to monitor phytoplanktonic diversity and degradation state in different suspended particulate matter (SPM) samples collected in summer surface waters across a South-North transect in the Chukchi Sea. For this, we analysed for various lipids and their biotic and abiotic degradation products, owing to their relative stability, specificity, and general suitability as tracers of the origin and fate of organic matter (OM) (Volkman et al., 1998). Moreover, the majority of studies dealing with the degradation of phytoplanktonic OM to date have focused on biotic degradation processes (Afi et al., 1996; Sun et al., 1999; Mäkinen et al., 2017), while the role played by photooxidation and free radical-mediated oxidation (autoxidation) processes in the degradation of lipid components during the senescence of phototrophic organisms has only very recently been investigated (Rontani, 2012; Rontani and Belt, 2020).

2. Material and methods

2.1. Sampling

Seawater sampling was carried out at 14 station over the Chukchi Sea (Table S1) during Korea research ice breaker R/V Araon cruise (ARA06B, August 1-22, 2015). A volume of 50-100 L of seawater was directly collected into 50 L carboy bottles from the vessels underway pumped system located at about 5 m below sea level. Filtration on GF/F filters (pre-combusted at 450°C for 6 h before use to avoid contamination) was performed at sea. The samples were frozen onboard (-20°C), freeze dried in the Korean laboratory and then shipped frozen to the UK and French laboratories.

2.2. Lipid extraction

Filters were reduced at room temperature with excess NaBH₄ (70 mg) after adding MeOH (25 mL, 30 min) to reduce labile hydroperoxides (resulting from photo- or autoxidation) to alcohols, which are more amenable to analysis by gas chromatography (GC). A test carried out with methyl oleate confirmed that the conversion of esters to the corresponding *n*-alcohols during the NaBH₄ reduction step of the treatment was negligible (amount of octadec-9-en-1-ol formed < 1% of the initial methyl oleate) and could not interfere with *n*-alkan-1-ol quantification in the different samples. Water (25 mL) and KOH (2.8 g) were then added and the resulting mixture saponified by refluxing (2 h). After cooling, the mixture was acidified (HCl, 2 N) to pH 1 and extracted with dichloromethane (DCM; 3×20 mL). The combined DCM extracts were dried over anhydrous Na₂SO₄, filtered, and concentrated by rotary evaporation at 40°C to give total lipid extracts (TLEs).

2.3. Silvlation

122 Dry TLEs and standards were derivatized by dissolving them in 300 μ L pyridine/bis-123 (trimethylsilyl)trifluoroacetamide (BSTFA; Supelco; 2:1, v/v) and silylated in a heating block 124 (50 °C, 1 h). After evaporation to dryness under a stream of N₂, the derivatized residue was 125 dissolved in ethyl acetate/BSTFA (to avoid desilylation) and analysed by GC-QTOF.

127 2.4. Gas chromatography-EI quadrupole time-of-flight mass spectrometry

Accurate mass measurements were made in full scan mode using an Agilent 7890B/7200 GC/QTOF system (Agilent Technologies, Parc Technopolis - ZA Courtaboeuf, Les Ulis, France). A cross-linked 5% phenyl-methylpolysiloxane (Agilent Technologies; HP-5MS Ultra inert, $30 \text{ m} \times 0.25 \text{ mm}$, 0.25 µm film thickness) capillary column was used. Analyses were performed with an injector operating in pulsed splitless mode set at 270 °C. Oven temperature was ramped from 70 °C to 130 °C at 20 °C min⁻¹ and then to 300 °C at 5 °C min⁻¹. The pressure of the carrier gas (He) was maintained at 0.69×10^5 Pa until the end of the temperature program. Instrument temperatures were 300 °C for transfer line and 230 °C for the ion source. Nitrogen (1.5 mL min⁻¹) was used as collision gas. Accurate mass spectra were recorded across the range m/z 50–700 at 4 GHz with the collision gas opened. The QTOF-MS instrument provided a typical resolution ranging from 8009 to 12252 from m/z 68.9955 to 501.9706. Perfluorotributylamine (PFTBA) was used for daily MS calibration. Compounds were identified by comparing their TOF mass spectra, accurate masses and retention times with those of standards. Quantification of each compound involved extraction of specific accurate fragment ions, peak integration, and determination of individual response factors using external standards and Mass Hunter software (Agilent Technologies, Parc Technopolis - ZA Courtaboeuf, Les Ulis, France). Saturated (SFAs) and monounsaturated fatty acids (MUFAs) were quantified with structurally similar elaidic and nonadecanoic acid standards, respectively,

and the other lipids with an exact standard. The limit of detection (LOD) for each compound was based on a signal-to-noise ratio (S/N) greater than 5. The linear range was determined using values that met the standard analysis criteria of less than 15% deviation across the concentration range. Linear responses (depending on the nature of the silvlated analytes) were obtained over 2-3 orders of magnitude.

The recovery efficiency of polyunsaturated fatty acids (PUFAs) for the sample treatment was checked with a standard of $C_{20:5}$ fatty acid and compared to that obtained for the $C_{18:1\Delta9}$ (oleic) acid. Using this approach, we estimate the efficiency of $C_{20:5}$ fatty acid to be ca. 70% compared to that of oleic acid. The sample treatment employed thus induces some losses (likely thermal) of PUFAs.

2.5. Chlorophyll-a analyses

The chlorophyll-a (Chl-a) concentration was determined onboard using a Trilogy fluorometer (Turner Designs, USA) after 24 h extraction in 90 % acetone at 4°C (Parsons et al., 1984). Chl a was measured at 0 and 10 m and we took the mean value to represent the samples collected via the ship's underway system (ca. 5 m).

2.6. Standard compounds

Phytol, fatty acids, most of the sterols and 2,6,10,14-tetramethylpentadecanoic acid (pristanic acid) were purchased from Sigma-Aldrich (St. Quentin Fallavier, France). 6,10,14-Trimethylpentadecan-2-ol was obtained after oxidation of phytol with KMnO₄ in acetone (Cason and Graham, 1965) and subsequent NaBH₄ reduction of the resulting ketone. The syntheses of 3-methylidene-7,11,15-trimethylhexadecan-1,2-diol (phytyldiol), 3,7,11,15-tetramethylhexadec-2(Z/E)-en-1,4-diols and 3,7,11,15-tetramethylhexadec-3(Z/E)-en-1,2-diols

were described previously by Rontani and Aubert (2005). 4,8,12-Trimethyltridecanoic acid (4,8,12-TMTD acid) was synthesized from isophytol (Interchim, Montluçon, France) by a previously described procedure (Rontani et al., 1991). 3,7,11,15-Tetramethylhexadecanoic acid (phytanic acid) was produced in three steps from phytol as described previously (Rontani et al., 2003). 3,7,11,15-Tetramethylhexadecan-1-ol (dihydrophytol) was obtained by hydrogenation of phytol in methanol with Pd-CaCO₃ as a catalyst (Rontani et al., 2003). Type-II photosensitized oxidation of monounsaturated fatty acids and alken-1-ols was carried out in pyridine with hematoporphyrin as sensitizer, while autoxidation was achieved using Fe²⁺/ascorbate (Loidl-Stahlhofen and Spiteller, 1994). Subsequent reduction of the resulting hydroperoxides in methanol with excess NaBH₄ afforded the corresponding hydroxyacids.

181 2.7. Determination of the double bond position of monounsaturated fatty acids and n-alken-1182 ols

The double bond positions of monounsaturated fatty acids (MUFAs) were determined via analysis of the TMS derivatives of their oxidation products (allylic hydroxyacids), which are ubiquitous in environmental samples (for a review see Rontani and Belt, 2020). Silylated hydroxyacids affording di-functionalized fragment ions upon electron ionization (EI) (Fig. 2) were selected for this purpose. Monitoring of the accurate mass of these fragment ions (Table S2) allowed an unambiguous assignment of the double bond position of the MUFAs present in the samples investigated (an example is given in Fig. 2). It may be noted that these double bond positions may also be identified (although less clear) by GC-MS using unit masses. However, in the case of *n*-alken-1-ols, the use of accurate masses (Table S3) is needed (Fig. 3). Indeed, unit masses of the fragment ions arising from EI fragmentation of the TMS derivatives of n-alken-1-ol oxidation products (diols) interfere with more stable isobaric fragment ions arising

from the fragmentation of oxidation products of some MUFAs, which are present in higher proportions than *n*-alken-1-ols in environmental samples.

This straightforward method for determining double bond positions only requires adding a simple NaBH₄ reduction step (to reduce unstable hydroperoxyacids to the corresponding hydroxyacids; Marchand and Rontani, 2001) prior to alkaline hydrolysis, which is commonly employed for the treatment of complex lipid extracts (Volkman, 2006).

2.8. Chlorophyll photooxidation estimates

Type II photosensitized oxidation of the chlorophyll phytyl side-chain leads to the production of 2-hydroperoxy-3-methylidene-7,11,15-trimethylhexadecan-1-ol, which, after NaBH₄ reduction, is converted to 3-methylidene-7,11,15-trimethylhexadecan-1,2-diol (phytyldiol) (Rontani et al., 1994). This diol constitutes a stable and specific tracer for the photodegradation of the chlorophyll phytyl side-chain (Cuny and Rontani, 1999). The molar ratio phytyldiol:phytol (Chlorophyll Phytyl side-chain Photodegradation Index, CPPI) has been previously proposed to estimate the extent of photodegradation of chlorophylls possessing a phytyl side-chain through use of the empirical equation: chlorophyll photodegradation % = (1- $[CPPI + 1]^{-18.5}$ × 100 (Cunv et al., 1999)' this equation was employed previously in natural marine samples from various latitudes (For reviews see Rontani and Belt, 2020; Rontani et al., 2021).

2.9. Lipid photo- and autoxidation estimates

Type-II photosensitized oxidation of Δ^5 -sterols produces mainly unstable Δ^6 -5 α -hydroperoxides with smaller amounts of stable and specific Δ^4 -6 α /6 β -hydroperoxides (Kulig and Smith, 1973). Sterol photooxidation percentage can be estimated on the basis of Δ^4 -

 $3\beta,6\alpha/\beta$ -dihydroxysterols (arising from NaBH₄ reduction of the corresponding hydroperoxides) concentration using the equation: sterol photooxidation $\% = (\Delta^4 - 3\beta, 6\alpha/\beta - 1)$ dihydroxysterol %) \times (1+0.3)/0.3 (Christodoulou et al., 2009), where the value (0.3) corresponds to the ratio Δ^4 -6 α/β -hydroperoxides/ Δ^6 -5 α -hydroperoxides found in biological membranes (Korytowski et al., 1992). Free radical autoxidation of Δ^5 -stenols yields mainly 10 222 unstable and unspecific 7α -and 7β -hydroperoxides and, to a lesser extent, specific $5\alpha/\beta, 6\alpha/\beta$ -epoxysterols and their stable hydrolysis products $3\beta_{2}5\alpha_{2}6\beta_{3}$ -trihydroxysterols (Smith, 1981). **225** The extent of sterol autoxidation can be estimated after alkaline hydrolysis using the equation: sterol autoxidation % = $(3\beta, 5\alpha, 6\beta$ -trihydroxysterols %) × 2.4 based on respective formation rate constants of the different autoxidation products obtained from incubation experiments (Christodoulou et al., 2010) and calculated previously by Morrisey and Kiely (2006). These equations have been employed previously to obtain reliable estimates of sterol photo- and autoxidation in various marine areas (For reviews see Rontani and Belt, 2020; Rontani et al., 2021).

Photo- and autoxidation of MUFAs and monounsaturated *n*-alken-1-ols yields mixtures of isomeric allylic hydroperoxyacids or hydroperoxyalcohols (Frankel, 1998; Rontani et al., 2012), respectively, which are converted to the corresponding hydroxyacids or diols after NaBH₄-reduction. The relative importance of photo- and autoxidation of these compounds can be readily calculated based on the proportion of *cis* isomers (which are produced specifically by autoxidation; Porter et al., 1995) and the temperature of seawater (Marchand and Rontani, 2001).

- 3. Results
- **232** 40 234 **239**

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3.1. Trophic environment across the Chukchi Sea transect

The main sterols: 24-norcholesta-5,22E-dien-3β-ol (24-norsterol), 27-nor-24-methylcholesta-5,22-dien-3 β -ol, cholest-5-en-3β-ol (cholesterol), cholesta-5,22E-dien-3B-ol (22 dehydrocholesterol), cholesta-5,24-dien-3β-ol (desmosterol), 24-methylcholesta-5,22E-dien-3β-ol (epi-brassicasterol), 24-methylcholesta-5,24(28)-dien-3β-ol (24-methylenecholesterol), 24-ethylcholest-5-en-3β-ol (sitosterol) and 24(E)-ethylcholesta-5,24(28)-dien-3 β -ol (fucosterol) were quantified across the transect to estimate the amount and nature of the algal material present in the SPM samples. The highest values of particulate sterol biomass were 20 250 observed at stations 03, 02, 01, 09, 13 and 20, with a maximum at Station 03 suggesting the presence of a bloom event (Table 1, Fig. 4B). In contrast, the other stations, and notably the under ice station 26, contained relatively low concentrations of such sterols, indicating low phytoplankton abundances. For the majority of samples, sterol profiles were dominated by 24-**254** norsterol, cholesterol, 22-dehydrocholesterol, epi-brassicasterol and 24-methylenecholesterol, the latter being particularly dominant at station 03. The proportion of sitosterol showed a marked increase at stations 21 and 22 (Fig. 4A).

Chl-a, fatty acid and phytol concentrations (Tables 2 and 3) confirmed the dominance of algal material at stations 03, 02, 01, 09, 13 and 20. The fatty acid profiles showed very low proportions of PUFAs (mainly C_{16:4}, C_{18:4} and C_{20:5}) along the transect, reaching less than 4% of total fatty acids at station 03. SFAs were dominated by the classical C_{14:0}, C_{16:0} and C_{18:0}, and MUFAs by $C_{16:1\Delta9}$ (palmitoleic acid), $C_{18:1\Delta9}$ (oleic acid) and $C_{18:1\Delta11}$ (vaccenic acid) (Table 2).

Long-chain saturated and monounsaturated primary alcohols (ranging from C_{14} to C_{22}) resulting from alkaline hydrolysis of zooplanktonic wax esters (Lee et al., 2006) were also detected, with highest concentrations at station 02 (Table 4). Interestingly, this station also exhibited the highest proportion of C_{20:1} and C_{22:1} *n*-alken-1-ols (Fig. 5A).

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3.2. Biotic and abiotic degradation of phyto- and zooplanktonic lipids

3.2.1. Chlorophyll

Chlorophyll photooxidation estimates ranged from 33% to 96% along the transect (Table 3). Several biotic (dihydrophytol, phytanic, pristanic and 4,8,12-TMTD acids) and autoxidative (3,7,11,15-tetramethylhexadec-2(Z/E)-en-1,4-diols, 3,7,11,15-tetramethylhexadec-3(Z/E)-en-1,2-diols, and isophytol) degradation products of the chlorophyll phytyl side-chain could be detected in all samples (Fig. 6A). The phytyl side chain appeared to be strongly degraded at all stations except station 03, this degradation mainly involving autoxidative and biotic processes (Fig. 6B). The percentage of autoxidation products ranged from 4% to 29% of total isoprenoid compounds (Fig. 6B) and from 10% to 190% of the residual phytyl chain. Autoxidation was particularly intense at stations 01, 11, 14, 18, 20, 21 and 26. With the exception of station 03, biotic degradation products of phytol were dominant, ranging from 36% to 78% of total isoprenoid compounds.

3.2.2. MUFAs, n-alken-1-ols and sterols

Type II photosensitized oxidation of MUFAs appeared to be very limited except at station 01 where 24% of vaccenic acid was photooxidized (Fig. 7C). While palmitoleic acid was very weakly affected by autoxidative processes (Fig. 7A), autoxidation reached 73% and 64% in the case of oleic and vaccenic acids, respectively (Figs. 7B and 7C). Monounsaturated n-alken-1ols arising from the hydrolysis of zooplanktonic wax esters were only weakly oxidized (Fig. 8). Interestingly, in the first part of the transect, photooxidation was more intense in the case of $C_{20:1\Delta11}$ and $C_{22:1\Delta11}$ *n*-alken-1-ols than for $C_{16:1\Delta9}$ and $C_{18:1\Delta9}$ (Fig. 8). Due to their relatively low contribution, photo- and autoxidation products of sterols (sum < 10%) were not quantified.

4. Discussion

4.1. Trophic environment at the different stations

Sterols possess structural characteristics (double bond positions, nuclear methylation and patterns of side-chain alkylation) restricted to a few groups of organisms (for reviews see Volkman, 1986; 2003; Rampen et al., 2010), and are often used to estimate phytoplanktonic diversity (Taipale et al., 2016). The diatom *Thalassiosira* aff. antarctica and the dinoflagellate Gymnodinium simplex have been proposed as potential sources of 24-norsterol (Rampen et al., 2007), which was found to be present in high proportion (ranging from 15 to 50%) along all the Chukchi Sea transect (Fig. 4A). Sterols that are methylated at the C-4 position including 4α ,23,24-trimethyl- 5α -cholest-22*E*-en-3-ol (dinosterol) are often employed as tracers for the contribution of dinoflagellates in the marine environment (Robinson et al., 1984). The absence (below detection limit) of such sterols at the different stations therefore allowed us to exclude a significant contribution of dinoflagellates across the transect. T. antarctica, which is found at high latitudes but not in tropical waters (Hasle and Heimdal, 1968), seems thus to strongly contribute to the phytoplankton community along all the Chukchi Sea transect. This centric diatom also contains a high proportion of 24-methylenecholesterol (Rampen et al., 2007), which is present with 24-norsterol at all stations (proportion ranging from 5 to 23% of total sterols) (Fig. 4A). It seems thus that T. antarctica or similar species belonging to the order of Thalassiosirales are one of the main phytoplankton contributors to these SPM samples. This assumption is in good agreement with the dominance of T. antarctica and T. nordenskioeldii previously observed in surface sediments of the Chukchi Sea (Asthakov et al., 2015). Further, *epi*-brassicasterol, which is also present in significant proportions along all the transect (Fig. 4A), suggests the presence of prymnesiophytes and notably of *Phaeocystis pouchetii*, whose sterol content comprises almost entirely this sterol (Nichols et al., 1991), and is widely

distributed in the Arctic (Riisgaard et al., 2015). The increasing proportions of sitosterol observed between stations 11 and 22 could potentially be attributed to the presence of increasing proportions of the diatoms *Asterionella glacialis*, *Haslea ostrearia* or *Amphiprora hyalina* (Volkman, 2003). Finally, the relatively high proportions of cholesterol present in most of the samples (Fig. 4A) suggest the occurrence of zooplanktonic material. Indeed, it is well-known that herbivorous and omnivorous crustaceans convert dietary phytosterols to cholesterol (Grieneisen, 1994; Behmer and Nes, 2003).

The presence of a high proportion of zooplanktonic material in the different samples is supported further by the detection of *n*-alkan-1-ols (Table 4) arising from alkaline hydrolysis of wax esters, which are generally the main storage lipids of marine zooplankton in highlatitude species (Lee et al., 2006). The most common n-alkan-1-ols of the wax esters of herbivorous copepods that undergo diapause (common in the Arctic) are $C_{20:1\Delta 11}$ and $C_{22:1\Delta 11}$, while omnivorous or carnivorous zooplankton generally show a predominance of $C_{14:0}$ and $C_{16:0}$ *n*-alkan-1-ols (Lee and Nevenzel, 1979; Albers et al., 1996). The proportion of herbivorous and omnivorous/carnivorous zooplankton was thus estimated on the basis of *n*-alkan-1-ol composition. The results obtained (Fig. 5B) show highest herbivorous zooplanktonic contribution at station 02 (where Calanus glacialis and Eucalanus bungii appeared to be dominant, Kim et al., 2020), which exhibits the highest concentration of *n*-alkan-1-ols (Table 4, Fig. 5A). Further, the very low amounts of the C_{20:5} acid, a well-known component of numerous zooplankton species (Cass et al., 2014), suggests the presence of fecal pellets, where PUFAs are generally only present in low proportions (Harvey et al., 1987; Prahl et al., 1985) compared to zooplanktonic organisms. Fecal pellets generally settle out of the water column relatively quickly. Although coprophagy is usually the more important source of fecal pellet degradation (Iversen and Poulsen, 2007), they also experience fragmentation (coprorhexy)

leading to the formation of small slowly sinking particles (Smetacek, 1980), which could havebeen collected during the sampling.

The presence of PUFAs in only trace amounts is very surprising; indeed, these compounds are generally major components of high-altitude phytoplankton blooms (Parrish et al. 2005; Leu et al. 2006; Marmillot et al. 2020). We suggest that the relatively low proportions of PUFAs observed may be attributed to: (i) a highly detritic state of all the SPM samples investigated, (ii) a degradation of the more labile components of these samples (including PUFAs) between sampling and analysis (ca. 5 years) despite storage at -20 °C, or (iii) thermal losses during the treatment (see section 2.4). Although PUFAs appeared to be strongly degraded, the detection of residual amounts of the unusual fatty acids $C_{16:4}$ (synthesized by diatoms; Volkman, 2006) and C_{18:4}, a well-known acid component of copepod wax esters (Kattner, 1989; Graeve and Kattner, 1992), is in good agreement with the trophic environment described above. The C_{18:4} fatty acid is also present in high proportion in *Phaeocystis* spp. (Sargent et al., 1985), but in this case it is accompanied by $C_{18:5}$ and $C_{22:6}$ fatty acids, which are absent in the SPM samples. The main source of the C_{18:4} fatty acid in our SPM samples seems thus to be copepod wax esters.

4.2. Biotic and abiotic degradation of phyto- and zooplanktonic lipids

4.2.1. Chlorophyll

It was demonstrated recently that high solar irradiances favour photobleaching of chlorophyll (sensitizer) at the expense of Type-II (i.e. involving singlet oxygen) photosensitized oxidation of lipids (the photodynamic effect) (Amiraux et al., 2016; Rontani et al., 2021). Since the SPM samples investigated in the present work were collected near to the surface (5 m), thereby receiving highest solar irradiance (up to 500 W m⁻², Dr. Ha, unpublished results), the elevated

366 CPPI-based chlorophyll photodegradation estimates are not surprising (Table 3). Further, we 367 note that even at station 03, 45% of chlorophyll was photooxidized, suggesting the diatoms 368 were in a poor physiological state, despite an apparent bloom event.

Isoprenoid compounds resulting from the degradation of the chlorophyll phytyl sidechain were not only useful for quantifying chlorophyll photodegradation (see section 2.7), but also for identifying the importance of autoxidative and biotic alteration of primary production. Autoxidation of the phytyl chain affords mainly isophytol (Rontani and Galeron, 2016), 3,7,11,15-tetramethylhexadec-2(Z/E)-en-1,4-diols and 3,7,11,15-tetramethylhexadec-3(Z/E)en-1,2-diols (Rontani and Aubert, 2005) (Fig. 9), which are specific tracers of these degradative processes. The detection of significant proportions of these compounds in some SPM samples (Fig. 6) thus demonstrates the important role played by autoxidation in the degradation of primary producers.

Pelagic crustaceans assimilate the chlorophyll phytyl chain when feeding herbivorously (Prahl et al., 1984; Harvey et al., 1987). During this metabolism, phytol is successively hydrogenated to dihydrophytol, oxidized to phytanic acid, and then cleaved to pristanic and 4,8,12-TMTD acids by way of classical α - and β -oxidation sequences (Fig. 9) (for a review see Rontani and Volkman, 2003). Such isoprenoid compounds have been detected previously in diff erent *Calanus* species (Avigan and Blumer, 1968; Prahl et al., 1984). As such, while phytol logically dominates isoprenoid compounds at station 03 due to a declining bloom event mainly composed of weakly altered diatoms, the other stations exhibit a strong dominance of phytanic and pristanic acids (Fig. 6B), which confirms a strong contribution of zooplanktonic material.

4.2.2. Unsaturated fatty acids

Photodegradation rates of unsaturated fatty acids increases, logically, with their degree of unsaturation (Frankel, 1998), such that PUFAs are very reactive towards Type-II

photosensitized oxidation processes (Frankel, 1998; Rontani et al., 1998). Unfortunately, the resulting oxidation products cannot be used as quantitative tracers of Type-II photooxidation processes in the environment due to the instability of the primary oxidation products formed (Rontani and Belt, 2020). However, it was demonstrated previously that PUFAs are photodegraded approximately 5 times faster than MUFAs in dead cells of diatoms (Rontani et al., 2011). Based on the very weak photooxidation percentage of MUFAs in our SPM samples (Fig. 7; mean value $1.9 \pm 4.2\%$), the contribution of Type-II photosensitized oxidation processes to the degradation of PUFAs thus appeared to be very weak (~ 10%). This very weak photooxidative alteration of PUFAs and MUFAs results likely from the limitation of the photodynamic effect under high solar irradiances (Amiraux et al., 2016; Rontani et al., 2021).

Whereas autoxidation of palmitoleic acid was limited to 25% in SPM samples (Fig. 7A), it reached 73% in the case of oleic acid (Fig. 7B). These differences of reactivity may be attributed to the origins of these MUFAs. Indeed, although palmitoleic and oleic acids are not unambiguous biomarkers of phytoplankton and zooplankton inputs (Wakeham and Canuel, 1988), they are often used to obtain relative indication of the predominance of algal vs. zooplanktonic sources (Tolosa et al., 2004). Palmitoleic acid, which is a major fatty acid of diatoms (Volkman et al., 1989; Dunstan et al., 1993), is also present in several bacteria (e.g., de Carvalho and Caramujo, 2014), and may be found in high proportion in storage lipids of zooplankton feeding on diatoms (Lee et al., 2006). In contrast, omnivorous and carnivorous zooplankton species are characterized by wax esters with high amounts of oleic acid (Lee et al., 2006). However, we note that *Phaeocystis* spp. may also contain elevated levels of oleic acid (Dalsgaard et al., 2003). On the basis of high proportions of phytanic and pristanic acids observed in all SPM samples (Fig. 5), we consider that the major source of oleic acid to be omnivorous zooplankton (more precisely euphausiids or Metridia pacifica detected in the samples analysed, Kim et al., 2020) rather than prymnesiophytes. Autoxidation seems thus to

be stronger in faecal pellets of omnivorous and carnivorous zooplankton than in diatoms or
herbivorous zooplankton feeding on diatoms. Further, we note that the highest autoxidation
percentages of oleic acid were observed at stations 01, 11, 14, 18, 20, 21 and 26, which also
exhibit the highest autoxidation of the chlorophyll phytyl side-chain (Fig. 6B).

Based on: (i) pseudo-first order autoxidative degradation rates of palmitoleic and $C_{20:5}$ acids previously measured during incubation of dead diatom cells $(1.5 \times 10^{-3} \text{ and } 2.7 \times 10^{-2} \text{ h}^{-1}$, respectively; Rontani et al., 2014), and (ii) the mean autoxidation percentage $(14.4 \pm 7.7\%)$ of palmitoleic acid observed in the different SPM samples (Fig. 6A), an important contribution of autoxidation to the disappearance of PUFAs in these samples may be expected. However, it is well known that PUFAs are also significantly reduced relative to total fatty acids during passage through the gut of copepods (Harvey et al., 1987; Prahl et al., 2009). Consequently, due to the important contribution of zooplanktonic faecal pellets to the SPM samples analysed, an important degradation of PUFAs during copepod grazing is very likely. Note that a significant degradation during the storage and the treatment cannot be totally excluded.

Surprisingly, vaccenic acid, well-known to be specific to bacteria (Lambert and Moss, 1983; Sicre et al., 1988), also appeared to be strongly autoxidized (Fig. 7C). Autoxidation states of vaccenic and oleic acids are not significantly different (Kruskal-Wallis, n=26 p = 0.817 > 0.05), while the autoxidation state of vaccenic acid is significantly different to that of palmitoleic acid (Kruskal-Wallis, n=26 p = 0.00025 < 0.05). Autoxidation of vaccenic acid seems thus to mainly intervene in omnivorous or carnivorous zooplanktonic faecal pellets, rather than in heterotrophic bacteria associated to diatom phytodetritus or present in faecal pellets of herbivorous zooplankton feeding on diatoms. Interestingly, the lipid composition of bacterivorous ciliates resembles that of their prey (Harvey et al., 1987; Boëchat and Adrian, 2005). Recently, the presence of high proportions of *trans* MUFAs in SPM samples collected under sea ice in Baffin Bay was attributed to the ingestion of bacteria stressed by salinity in

internal brines of sea ice by sympagic ciliates, and the direct incorporation of these highly
isomerized dietary fatty acids (Burot et al., 2021). A trophic link between copepods and ciliates
has been well established (Pierce and Turner 1992), and some authors have shown that lipids
of bacterial origin can be transferred up the food web without modification to copepods that
consume bacterivorous ciliates (Ederington et al., 1995). The presence of both vaccenic and
oleic acids in copepod wax esters, as in the case of some euphausiids (Lee et al., 2006), could
thus explain their similar autoxidative reactivity (Figs. 7B and 7C).

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4.2.3. Monounsaturated n-alkan-1-ols

Although Type II photosensitized oxidation of monounsaturated n-alkan-1-ols was also relatively weak (Fig. 8), likely due to the strong chlorophyll photobleaching, it is interesting to note that C_{20:1Δ11} and C_{22:1Δ11} n-alkan-1-ols (specific to herbivorous copepod wax esters; Lee et al., 2006) were more photooxidized in the first five stations (01, 02, 03, 09, and 11) than the non-specific $C_{16:1\Delta9}$ and $C_{18:1\Delta9}$ *n*-alken-1-ols (Kruskal-Wallis, n=20 p = 0.00516 < 0.05). Previously, a strong photooxidation of $C_{20:1\Delta 11}$ and $C_{22:1\Delta 11}$ *n*-alken-1-ols was observed in sinking particles collected in summer at 100 m in the Beaufort Sea (Rontani et al., 2012), and attributed to a very high efficiency of Type-II photosensitized oxidation processes in faecal pellets of herbivorous copepods. A very efficient transfer of singlet oxygen from more or less digested diatoms to the wax esters present in the lipid droplets trapped in faecal pellets (Najdek et al., 1994) was also proposed. The strongest photooxidation observed in the Beaufort Sea likely results from a fast transfer of faecal pellets to deep waters where the intensity of solar irradiance should be sufficiently weak to favour the photodynamic effect at the expense of chlorophyll photobleaching. The weak and selective photooxidation of $C_{20:1\Delta 11}$ and $C_{22:1\Delta 11}$ *n*-alken-1-ols observed during the present study confirms: (i) the involvement of this process in faecal pellets of herbivorous copepods, and (ii) the key role played by solar irradiance and

sinking rates in the efficiency of the photodynamic processes. Due to the intensity of solar
irradiance during summers, only the chlorophyll of senescent diatom cells and the debris of
herbivorous zooplankton faecal pellets (which are suspended in surface waters of the Chukchi
Sea), is thus strongly affected by photooxidation processes.

Monounsaturated *n*-alkan-1-ols also appeared to be weakly affected by autoxidative processes in our Chukchi Sea samples (Fig. 8). Since autoxidation of monounsaturated long-chain MUFAs and *n*-alkan-1-ols involves attack of the allylic positions of their isolated double bonds (Frankel, 1998), their degradation rates should be very similar and only weakly impacted by esterification. The differences in reactivity observed between these alcohols and oleic and vaccenic acids (Figs. 7B and 7C) are thus very surprising. Autoxidation of wax esters of omnivorous or carnivorous copepods containing high amounts of oleic acid and of the unreactive C_{14:0} and C_{16:0} alcohols (Albers et al., 1996) seems thus to be strongly favoured (Fig. 7B), while wax esters of herbivorous copepods dominated by $C_{20:1\Delta 11}$ and $C_{22:1\Delta 11}$ *n*-alken-1-ols and C_{14:0}, C_{16:0}, C_{20:1} and C_{22:1} fatty acids (Lee et al., 2006) are only weakly affected (Fig. 8). It was previously observed that herbivorous amphipods are somewhat resistant to oxidative stress due to the presence of high concentrations of antioxidants (i.e. mycosporine-like amino acids and carotenoids) ingested from their algal diet (Abele and Puntarulo, 2004; Obermüller et al., 2005; Obermüller, 2006), while carnivorous species are very sensitive to this stress due to their lower accumulation of algal antioxidants via the food chain (Obermüller et al., 2005). The differences in autoxidative reactivity observed in the present work between oleic acid (main acid counterpart of omnivorous or carnivorous copepod wax esters; Albers et al., 1996) and C_{20:1Δ11} and C_{22:1Δ11} *n*-alken-1-ols (main alcohol counterparts of herbivorous copepod wax esters; Lee et al., 2006) thus complement these previous findings.

It is generally considered that sinking of zooplanktonic faecal pellets to the seafloor is strongly limited by coprophagy (Paffenhöfer and Knowles, 1979) and bacterial decomposition

491 (Smetacek, 1985). In the case of faecal pellets of carnivorous or omnivorous zooplankton,492 autoxidation processes seem thus to also contribute to their degradation.

4. Conclusions

Analysis of the lipid content of SPM samples collected in surface waters of the Chukchi Sea enabled us to confirm: (i) their detritic character, (ii) the dominance of diatoms (Thalassiosirales) among primary producers, and (iii) the presence of high proportions of zooplanktonic (copepods) faecal pellet debris during the summer period. Type II photosensitized oxidation of phyto- and zooplanktonic lipids (photodynamic effect) appeared to be relatively limited, likely due to the enhancement of photobleaching of chlorophyll (sensitizer) by the high solar irradiance observed near to the surface. The weak photooxidation of zooplanktonic wax esters observed in these SPM samples contrasts with the strong photooxidative alteration of these compounds previously observed in sinking particles collected from the Beaufort Sea in summer (Rontani et al., 2012). The photodynamic effect seems thus to be favoured in large faecal pellets of herbivorous copepods sinking quickly in weakly irradiated zones. In contrast, autoxidative processes, which can operate in all oxic aquatic environments (Schaich, 2005) and potentially affect all unsaturated lipids (Rontani, 2012; Rontani and Belt, 2020), acted more intensively in faecal pellets of omnivorous or carnivorous copepods than in those of herbivorous counterparts. Such differences of reactivity were attributed to the consumption of algal antioxidants during the diet of omnivorous or carnivorous copepods, which should favour the establishment of free radical oxidation processes.

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FIGURE CAPTIONS

Figure 1. Map of the study area with location of the stations investigated in Chukchi Sea. The line corresponds to the limit of ice at the time of sampling.

Figure 2. GC-QTOF determination of the double bond position of MUFAs with the TMS derivatives of their oxidation products in the SPM sample collected at station 02.

Figure 3. GC-QTOF determination of the double bond position of monounsaturated *n*-alkan-1-ols with the TMS derivatives of their oxidation products in the SPM sample collected at station 02.

Figure 4. Relative proportions (A) and concentrations (ng L^{-1}) (B) of sterols in SPM samples collected at the different stations investigated.

Figure 5. Concentrations (μ g L⁻¹) of *n*-alkan-1-ols (A) and proportions of herbivorous and omnivorous or carnivorous zooplankton (B) in SPM samples collected at the different stations investigated.

Figure 6. Concentrations (ng L⁻¹) of chlorophyll phytyl side-chain and its main degradation products (A) and estimates of the proportion of its biotic, autoxidative and photooxidative degradation (B) in SPM samples collected at the different stations investigated.

Figure 7. Relative percentage of intact, autoxidized and photooxidized palmitoleic (A), oleic (B) and vaccenic (C) acids in SPM samples collected at the different stations investigated.

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1 2 3	825	Figure 8 . Relative percentage of intact, autoxidized and photooxidized $C_{16:1\Delta9}$, $C_{18:1\Delta9}$, $C_{20:1\Delta11}$
4 5 6	826	and $C_{22:1\Delta 11}$ <i>n</i> -alkan-1-ols in SPM samples collected at the different stations investigated.
7 8 9	827	
10 11	828	Figure 9. Biotic and abiotic degradation of the chlorophyll phytyl side-chain (compounds in
12 13 14	829	blue were quantified during the present work).
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Oxidation products of Δ^9 *n*-alken-1-ols





Stations

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Stations



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C_{18·1A9} *n*-alken-1-ol





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Table 1

Concentrations (ng L^{-1}) of the main sterols in the samples investigated.

Sterol	St 01	St 02	St 03	St 09	St 11	St 13	St 14	St 15	St 18	St 20	St 21	St 22	St 26
24-norcholesta-5,22 <i>E</i> -dien-3 β -ol	14.8	37.7	54.1	7.5	2.0	9.0	2.9	1.9	0.8	9.1	1.6	1.6	1.6
Cholesta-5,22 <i>E</i> -dien-3 β -ol	2.4 6.7	6.5 13.1	16.0 24.6	5.7 6.1	0.7 1.3	0.9 6.3	0.3 1.3	0.9 1.0	0.4 0.5	0.1 2.9	0.4 0.5	0.7 0.9	0.1 0.2
Cholest-5-en-3β-ol	6.4	28.1	23.5	13.2	2.6	8.7	2.9	3.1	1.0	2.3	1.2	1.2	0.8
24-methylcholesta-5,22 <i>E</i> -dien-3 β -ol	13.1 2.4	17.1	33.8 5.9	8.1 2.4	1.2	4.1 0.6	1.1 1.4	1.0 nd ^a	0.3 nd	1.7	0.3 nd	0.7 nd	0.2 nd
24 -methylcholesta-5,24(28)-dien-3 β -ol	2. 4 9.1	6.0	47.1	2. 4 5.8	1.2	3.9	0.6	0.4	0.3	3.0	0.2	0.4	0.2
24-ethylcholest-5-en-3β-ol	1.0	1.4	2.5	1.3	0.4	2.2	0.7	0.6	0.3	0.6	0.7	2.2	0.1
24(<i>E</i>)-ethylcholesta-5,24(28)-dien-3β-ol Total	nd 55.9	nd 113.2	1.2 208.8	1.3 51.4	0.1 9.8	0.5 36.1	0.3 11.6	0.2 9.0	0.1 3.6	nd 20.0	nd 4.9	nd 7.6	nd 2.2

^a nd = not detected (< 0.1 ng L^{-1})

Table 2

Concentrations (μ g L⁻¹) of saturated and monounsaturated fatty acids in the samples investigated. PUFAs present in very weak proportion (reaching less than 4% at station 03) were not included in this table.

Station	C _{14:0}	C _{16:1Δ9}	C _{16:1Δ11}	C _{16:0}	C18:149	C _{18:1Δ11}	C _{18:0}	C _{20:1Δ9}	C _{20:1Δ11}	C _{20:1Δ13}	C _{22:1Δ9}	C _{22:1Δ11}	C _{22:1Δ13}	Total
01	2 67	1 09	0.08	8 58	0 33	0.15	0.30	0.02	0.06	0.04	0.01	0.13	0.01	13.5
02	6.36	9.20	0.86	14.41	3.75	1.81	1.88	0.02	0.35	0.07	0.23	0.13	0.01	39.5
03	12.61	17.95	2.93	18.52	3.78	2.19	1.21	0.05	0.12	0.03	0.03	0.23	0.06	59.7
09	2.90	3.28	0.55	8.18	1.63	0.95	1.33	0.02	0.23	0.05	0.08	0.57	0.03	19.8
11	0.55	0.58	0.03	2.47	0.32	0.10	0.21	0.01	0.11	0.03	0.03	0.29	0.02	4.7
13	2.51	6.53	0.18	6.31	3.43	0.71	0.62	0.02	0.14	0.06	0.03	0.25	0.03	20.8
14	0.46	1.60	0.02	3.08	0.80	0.11	0.42	0.01	0.08	0.02	0.01	0.17	0.01	6.8
15	0.75	0.88	0.03	3.35	1.12	0.19	1.79	nd	0.02	0.01	nd	0.12	0.02	8.3
18	0.17	0.57	nd ^a	1.56	0.17	0.04	0.30	nd	0.03	0.01	0.01	0.30	0.01	3.3
20	1.47	8.97	nd	7.08	0.53	0.25	0.38	0.02	0.06	0.04	nd	0.12	0.02	18.9
21	0.52	1.10	0.02	2.58	0.25	0.05	0.27	nd	0.03	0.01	nd	0.18	0.01	5.0
22	1.57	5.64	0.10	4.69	2.14	0.27	0.89	nd	0.08	0.02	0.02	0.03	0.01	15.5
26	0.27	0.52	0.02	2.06	0.17	0.04	0.33	nd	0.05	0.01	0.02	0.04	0.01	3.5

^a nd = not detected (< 0.01 μ g L⁻¹)

Table 4

Concentrations (μ g L ⁻¹) of <i>n</i> -alkan-1-o	ls in the	samples	investigated.
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Compounds	St 01	St 02	St 03	St 09	St 11	St 13	St 14	St 15	St 18	St 20	St 21	St 22	St 26
C14:0	0.24 0.12	0.40 0.36	0.57 0.45	0.18 0.09	0.15 0.10	0.24 0.41	0.10 0.36	0.06 0.07	0.02 0.07	0.12 0.63	0.05 0.11	0.10 0.32	0.09 0.10
$C_{16:1\Delta 11}$	0.04	0.07	0.24	0.02	0.01	0.01	0.01	0.01	nd ^a	0.01	0.01	0.01	nd 0.09
$C_{16:0}$ $C_{18:1\Delta9}$	0.20	0.20	0.87	0.06	0.13	0.40	0.18	0.00	0.03	0.08	0.07	0.22	0.09
$C_{18:1\Delta 11}$ $C_{18:0}$	0.04 0.13	0.11 0.27	0.07 0.16	0.01 0.13	0.05 0.08	0.10 0.13	0.06 0.11	0.02 0.09	0.01 0.05	0.03	0.01 0.07	$\begin{array}{c} 0.04 \\ 0.08 \end{array}$	0.02 0.06
C _{20:1} Δ9 C _{20:1} Δ11	0.01 0.02	0.77 0.46	0.41 0.42	0.02 0.28	nd 0.09	nd 0.19	nd 0.13	nd 0.01	nd 0.02	nd 0.01	nd 0.01	nd 0.17	nd 0.11
$C_{20:1\Delta_{13}}$	nd nd	0.04 0.02	nd 0.01	0.02 0.02	0.01 nd	0.02 0.01	0.02 0.01	nd nd	nd nd	nd 0.01	nd nd	0.01 0.01	0.01 0.01
C _{22:1} Δ11	0.01	1.18	0.11	0.19	0.05	0.07	0.06	nd	0.02	nd	0.01	0.08	0.05
C _{22:1Δ13} Total	0.96	5.16	3.56	1.76	0.01	2.22	1.24	0.43	0.26	1.09	0.38	1.13	0.58

 $a nd = not detected (< 0.01 \ \mu g \ L^{-1})$

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Table S1

Station	Sampling date	Latitude (°N)	Longitude (°W)
01	02/08/2015	65.173	168.690
02	02/08/2015	66.630	168.688
03	02/08/2015	67.333	168.834
09	03/08/2015	69.167	168.668
11	04/08/2015	71.430	168.670
13	04/08/2015	73.581	168.283
14	05/08/2015	74.798	167.899
15	05/08/2015	75.250	171.901
18	07/08/2015	75.761	177.152
20	08/08/2015	76.007	173.608
21	08/08/2015	73.999	169.997
22	09/08/2015	77.006	173.617
26	13/08/2015	80.774	172.849

Coordinates and sampling dates of the different stations sampled.

Table 3

Concentrations of isoprenoid compounds, chlorophyll and chlorophyll photooxidation estimates in the samples investigated.

	St 01	St 02	St 03	St 09	St 11	St 13	St 14	St 15	St 18	St 20	St 21	St 22	St 26
3.7.11.15-tetramethylbexadec-2-en-1-ol (Phytol) ^a	286	762	3253	351	34	104	40	3	8	77	9	39	6
4.8.12-trimethytridecanoic acid (4.8.12-TMTD acid) ^a		28	20	35	33	17	19	4	6	18	10	20	8
2.6.10.14-tetramethylpentadecanoic acid (Pristanic acid) ^a		585	113	20	4	17	4	3	1	11	3	7	2
2,6,10,14-tetramethylhexadecanoic acid (Phytanic acid) ^a		745	553	346	64	143	26	20	10	169	30	77	23
3,7,11,15-tetramethylhexadec-1-en-3-ol (Isophytol) ^a		27	113	7	28	3	12	nd ^e	9	27	15	1	7
3-methylidene-7,11,15-trimethylhexadecan-1, 2-diol ^a		39	134	10	3	5	1	1	1	18	2	5	1
3,7,11,15-tetramethylhexadecan-1-ol (Dihydrophytol) ^a		24	46	17	4	7	1	2	1	10	2	4	1
3,7,11,15-tetramethylhexadec-3-en-1,2-diols (Z and E) ^a		29	61	10	2	5	2	1	1	5	1	2	1
3,7,11,15-tetramethylhexadec-2-en-1,4-diols (Z and E) ^a		77	156	28	5	15	5	1	2	10	1	6	1
6,10,14-trimethylpentadecan-2-ol ^a	23	30	59	18	6	12	4	3	2	12	3	6	2
Total	951	2346	4508	842	183	328	114	38	41	357	76	167	52
CPPI ^b	0.191	0.041	0.033	0.022	0.072	0.041	0.028	0.173	0.071	0.187	0.180	0.096	0.104
Chlorophyll ^c	2.50	2.01	7.17	0.68	0.27	0.29	0.16	0.04	0.06	1.01	0.09	0.13	0.11
Chlorophyll photooxidation estimate (%) ^d		52.7	45.3	33.0	72.2	52.8	40.5	94.8	71.8	95.8	95.4	81.5	84.0

^a (ng L⁻¹)

^b Chlorophyll Phytyl side-chain Photooxidation Index (molar ratio 3-methylidene-7,11,15-trimethylhexadecan-1,2-diol /phytol)

 $^{c}(\mu g L^{-1})$

^d Estimated with the empirical equation: chlorophyll photodegradation $\% = (1 - [CPPI + 1]^{-18.5}) \times 100$ (Cuny et al. 2002).

^e nd = not detected ($< 1 \text{ ng } L^{-1}$)

Table S2

Parent fatty acid m/z, *m/z*, m/zm/z 199.1518^{a} 213.1675^{a} 329.1968^b 343.2125^b C_{16:1}29 371.2437 171.1206 185.1363 357.2280 $C_{16:1\Delta11}$ 399.2749 143.0748 157.1051 385.2592 C_{16:1Δ13} 227.1830 241.1987 329.1968 343.2125 C_{18:1} 199.1518 213.1675 357.2280 371.2437 $C_{18:1\Delta11}$ 385.2592 399.2749 171.1206 185.1363 $C_{18:1\Delta13}$ 255.2139 329.1968 343.2125 269.2295 C20:149 357.2280 227.1830 241.1987 371.2437 $C_{20:1\Delta11}$ 385.2592 199.1518 213.1675 399.2749 $C_{20:1\Delta13}$ 283.2451 297.2607 329.1968 343.2125 C_{22:1}Δ9 255.2139 269.2295 357.2280 371.2437 $C_{22:1\Delta11}$ 227.1830 241.1987 385.2592 399.2749 $C_{22:1\Delta13}$

Accurate masses of the main fragment ions produced during EI fragmentation of silylated monounsaturated fatty acid autoxidation products.

^a Fragments containing the terminal methyl group.

^b Fragments containing the trimethylsilyl ester group.

Table S3

Accurate masses of the main fragment ions produced during EI fragmentation of silylated monounsaturated *n*-alken-1-ol autoxidation products.

Parent <i>n</i> -alkenol	<i>m/z</i> ,	m/z.	<i>m/z</i> .	m/z.
C16:140	199.1518 ^a	213.1675 ^a	315.2170 ^b	329.2327 ^b
C16:1A11	171.1206	185.1363	343.2483	357.2640
$C_{16:1\Delta13}$	143.0748	157.1051	371.2796	385.2953
C _{18:1}	227.1830	241.1987	315.2170	329.2327
C _{18:1Δ11}	199.1518	213.1675	343.2483	357.2640
C _{18:1Δ13}	171.1206	185.1363	371.2796	385.2953
C _{20:1} 29	255.2139	269.2295	315.2170	329.2327
C _{20:1Δ11}	227.1830	241.1987	343.2483	357.2640
C _{20:1Δ13}	199.1518	213.1675	371.2796	385.2953
C22:1Δ9	283.2451	297.2607	315.2170	329.2327
C _{22:1Δ11}	255.2139	269.2295	343.2483	357.2640
C _{22:1} Δ13	227.1830	241.1987	371.2796	385.2953

^a Fragments containing the terminal methyl group.

^b Fragments containing the terminal trimethylsilyl ether group.