Interfacial photochemistry of marine diatom lipids: Abiotic production of volatile organic compounds and new particle formation

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**Abstract**

The global importance of abiotic oceanic production of volatile organic compounds (VOCs) still presents a source of high uncertainties related to secondary organic aerosol (SOA) formation. A better understanding of the photochemistry occurring at the ocean-atmosphere interface is particularly important in that regard, as it covers > 70% of the Earth’s surface. In this work, we focused on the photochemical VOCs production at the air-water interface containing organic material from authentic culture of marine diatom *Chaetoceros pseudocurvisetus*. Abiotic VOCs production upon irradiation of material originating from total phytoplankton culture as well as the fraction containing only dissolved material was monitored by means of PTR-ToF-MS. Furthermore, isolated dissolved lipid fraction was investigated after its deposition at the air-water interface. All samples acted as a source of VOCs, producing saturated oxygenated compounds such as aldehydes and ketones, as well as unsaturated and functionalized compounds. Additionally, a significant increase in surfactant activity following irradiation experiments observed for all samples implied biogenic material photo-transformation at the air-water interface. The highest VOCs flux normalized per gram of carbon originated from lipid material, and the produced VOCs were introduced into an atmospheric simulation chamber, where particle formation was observed after its gas-phase ozonolysis. This work clearly demonstrates abiotic production of VOCs from phytoplankton derived organic material upon irradiation, facilitated by its presence at the air/water interface, with significant potential of affecting the global climate as a precursor of particle formation.

# **1. Introduction**

Unicellular marine microalgae are responsible for approximately half of the global net primary production (Field et al., 1998, Bryant, 2003). One of the ecologically most important groups of phytoplankton are diatoms, ubiquitous across aquatic environments, from tropical to polar regions, and from highly dynamic coastal and upwelling habitats to more stable oceanic waters (Malviya et al., 2016, Bruland et al., 2005), accounting for approximately 20% of the global primary production (Mock and Medlin, 2012). Diatoms thus play a major role in carbon biogeochemical cycles (Field et al., 1998), in part through volatile organic compounds (VOCs) production, as a way of enhancing abiotic stress resistance, transferring information, protecting against predators and playing allelopathic roles (Zuo, 2019).

Major components of diatoms are lipids, accounting for up to 25% of their dry weight (Levitan et al., 2014). Production of lipids in diatom varies depending on culture conditions (Yi et al., 2017), however, diatom lipid material contains nearly all lipid classes, including both polar (Guschina and Harwood, 2006) and neutral lipids, such as free fatty acids (FFA), sterols (ST) and triacylglycerols (TG). Biogenic lipids are integral parts of cellular membranes (digalactosyldiacylglycerol (DGDG), monogalactosyldiacylglycerol (MGDG), sulfoquinovosyldiacylglycerol (SQDG), phosphatidylglycerol (PG) and phosphatidylcholine (PC)) (Stonik and Stonik, 2015) and energy storage molecules (TG), especially during specific environmental conditions such as nutrient starvation (Levitan et al., 2015).

Phytoplankton presents a main source of marine lipids through activities such as direct release, metabolic processes and cell lysis, which then have a significant impact on various processes in the aquatic environment due to their amphiphilic nature and the ability to accumulate at interfaces. As the top millimetre of the sea surface, the sea surface microlayer (SML) represents the largest environmental interface where many important processes such as wind action, water transpiration, solar energy flux and atmospheric interactions take place (Blough, 2005; Wurl et al., 2017). The SML is enriched with surface active organic matter (or surfactants) which accumulates at the air-water interface and forms surface films (Wurl and Holmes, 2008; Wurl et al., 2011; Frka et al., 2009, 2012). Although lipid material presents only a small portion of the organic matter (OM) pool in the SML, its presence at the air-water interface is the result of competitive adsorption and segregation from other macromolecular constituents due to its extremely high surface affinity (Frka et al., 2012). Biogenic lipids are also known to play a role in the formation and stabilization of the SML, as well as the physico-chemical, and morphological properties of the marine surface films (Van Vleet and Williams, 1983). Such surface films were found to have considerable effects on marine biogeochemical and climate-related mechanisms by directly affecting processes such as exchange of trace gases (e.g., CO2) (Engel et al., 2017; Milinković et al., 2022; Barthelmeß et al., 2021). Additionally, OM can be emitted directly from the ocean to the atmosphere as a major component of primary marine aerosol (Cochran et al., 2016; Chingin et al., 2018; Frossard et al., 2019) by wind-driven processes and bubble bursting (Facchini et al., 2008; Frossard et al., 2014, 2019). In primary aerosol, organic compounds enriched in the SML have been identified, including surface active dissolved organic matter (DOM) like polymers incorporated into microgels, but also transparent exopolymer particles (TEP) and Coomassie stainable particles (CSP) (Kuznetsova et al., 2005; Orellana et al., 2011; Aller et al., 2017). Long chain saturated fatty acids, such as palmitic acid and stearic acid, corresponding to major constituents of the SML, were also detected in marine aerosols (Marty et al., 1979; Slowey et al., 1962; Wu et al., 2015; Hu et al., 2018; Kang et al., 2017). The transfer and enrichments of the biogenic lipid classes on aerosol particles may be related to the distribution of compounds within the bubble-air-water interface (Triesch et al., 2021, Ellison et al., 1999).

Furthermore, recent field studies confirmed for the first-time previous laboratory observationssuggesting that irradiation of the air-water interface by sunlight presents a novel source of oxygenated VOCs to the marine boundary layer (Mungall et al., 2017). These emissions were exclusively attributed to the photochemical reactions facilitated by the presence of surfactants at the air-sea interface. Namely, experimental photosensitized reactions at the air-water interface by using humic acids as a proxy for DOM have led to chemical conversion of linear saturated fatty acids into unsaturated functionalized gas-phase products (Ciuraru et al., 2015a). Atmospheric photochemistry was even shown to take place in the absence of photosensitizers if the air-water interface is coated with a fatty acid (Rossignol et al., 2016). On a global scale, interfacial photochemistry has recently been suggested to act as an abiotic source of VOCs, comparable to their marine biological emissions (Brüggemann et al., 2018). VOCs have been shown to be important precursors to secondary organic aerosol (SOA) formation. Recent studies revealed that SOA, accounting for as much as 50-85% of the total organic aerosol burden (Jimenez et al., 2009), can strongly impact the radiation balance of the atmosphere, modify cloud microphysics, and participate in chemical transformations (Zhu et al., 2017). Marine SOA of biogenic origin could be especially important for understanding the cloud-mediated effects of aerosols on climate, because cloud properties respond to aerosols in a nonlinear way and are most sensitive to the addition of particles when the background concentration is low.

Although previous studies have demonstrated photoinduced VOCs production from artificial surfactants in laboratory grade water (Fu et al., 2015; Alpert et al., 2017; Bernard et al., 2016), saline solutions (Ciuraru et al., 2015a), and biofilm-containing solutions comprising a mixture of different microorganisms (Brüggemann et al., 2017), these experiments were typically conducted under far from ambient conditions or for a very limited number of authentic samples (Ciuraru et al., 2015b). To investigate the potentially underestimated VOCs source under more realistic conditions, in this study the biogenic OM was produced from an authentic diatom culture. Additionally, lipid material was isolated to further investigate phytoplankton lipid material as the main driver of surfactant release and, thus, abiotic photochemical VOCs production. Experiments were conducted in a photochemical reactor and an atmospheric simulation chamber, to study VOCs formation upon irradiation and implications for aerosol formation and growth, respectively. An improved chemical characterization of the surface-active organic compounds of marine origin and their interfacial photochemical processing is highly desirable to better understand the abiotic VOCs sources, and to improve our understanding of their subsequent impact on the climate.

# **2. Methods**

## 2.1. *Chaetoceros pseudocurvisetus* culture and sample preparations

Batch culture of C. pseudocurvisetus (105 cells) was inoculated in sterile VWR® Tissue Culture Flasks (VWR, Radnor, Pennsylvania) and placed in a temperature-controlled incubator (20 ℃) under illumination of 4500 lx on a 12h/12h light/dark cycle. The growth medium was F/2 medium (Guillard, 1975) with a lowered nitrogen content (NO3−: 42 μmol L−1), relevant for nutrient depleted oligotrophic waters, which represent 60% of the global ocean. The media was prepared in North Adriatic seawater rested for 6 months in the dark, filtered through sterile 0.22 μm filters (Merck Millipore Ltd.) and microwave boiled (Keller et al., 1988). Media amendments were added aseptically after sterilization. The cell number was controlled by a Fuchs-Rosenthal Chamber hemocytometer equipped with an Olympus BX51-P polarizing microscope. The growth was terminated on day eight, at the onset of the stationary phase.

At the onset of stationary phase growth (4 x 106 cells), an aliquot of the original non-filtered phytoplankton solution (hereafter, total solution, TS) was immediately frozen and kept at -20 ℃ until further experiments. Due to cell bursting caused by freezing and defrosting, the TS sample used for experiments contained dead cell material along with the phytoplankton produced dissolved organic matter. Additionally, an aliquot of the phytoplankton solution was immediately filtered through a pre-combusted (5 h at 450 ℃) 0.7 µm Whatman (Kent, UK) GF/F filter, in order to obtain phytoplankton derived dissolved organic matter sample (DS). Following filtration, the DS was immediately frozen and kept at -20 ℃ until further experiments. In order to isolate a sample of biogenic lipid material (LS) and analyse its lipid classes, 200 mL of the filtrate was collected into a chromic acid washed and ultrapure water (Merck Millipore, Burlington, Massachusetts, USA) rinsed glass bottle, after addition of an internal standard, 2-nonadecanone (purity ≥ 97%, Sigma Aldrich) (5 µg). The detailed procedure can be found elsewhere (Gašparović et al., 2017, 2015). Dissolved lipids were extracted by liquid-liquid extraction with dichloromethane (LiChrosolv®, Merck), twice at pH 8 and twice at pH 2, following hydrochloric acid addition. Extracts were evaporated until dry by rotary evaporation under a nitrogen atmosphere and stored at -20 ℃ until measurements.

## 2.2. Organic matter analysis

### 2.2.1. Particulate and dissolved organic carbon

For measurements of dissolved (DOC) and particulate (POC) organic carbon concentrations, the culture sample was filtrated through a pre-combusted (5 h at 450 ℃) 0.7 µm Whatman GF/F filter. Following filtration, the GF/F filter was stored at −80 ℃ until POC analysis. DOC analysis aliquots were collected in duplicates in 22 mL glass vials pre-combusted at 450 °C for 4 h. Sample preservation was done with mercury chloride (10 mg L−1) and samples were stored at +4 ℃ in the dark until analysis. Samples for DOC and POC were analysed according to methods EN 1484:2002 in a laboratory accredited according to the EN ISO/IEC 17025:2017 standard. A TOC-VCPH analyser (Shimadzu, Japan) equipped with a platinum silica catalyst and nondispersive infrared (NDIR) detector for CO2 measurements was used for DOC measurements. Concentration was calculated as an average of 3–5 replicates. The average instrument and Milli-Q blank were 0.03 mg L−1, with a high reproducibility (1.6%). POC was analysed with a solid sample module SSM-5000A associated to a Shimadzu TOC-VCPH carbon analyser calibrated with glucose. To remove inorganic carbonate fraction, filters were acidified with hydrochloric acid (2 mol L−1), and dried afterwards at 50 ℃ for 12 h. The samples were incinerated in a flow of oxygen at 900 ℃, and the CO2 produced was detected by an NDIR detector. POC concentrations were corrected based on the blank filter measurements. The average instrument and Milli-Q blanks correspond to 0.42 µmol L−1. The reproducibility of measurements determined by using a glucose standard was 3%.

### 2.2.2. Surfactant activity

Phase sensitive alternating current (a.c.) voltammetry (out-of-phase signal, frequency 77 Hz, amplitude 10 mV) was used for the quantification of surface-active substances (SAS) in TS and DS samples before and after irradiation. The concentration of SAS is expressed as the equivalent amount of the selected standard of non-ionic surfactant tetra-octylphenolethoxylate (Triton-X-100; mg L−1) as described in details previously (Ćosović and Vojvodić, 1982, Frka et al., 2009). Electrochemical measurements were performed using 663 VA Stand with potentiostat/galvanostat µAutolab II and automatic Hg drop production (Metrohm Autolab B.V., The Netherlands) equipped with GPES 4.9 software (Eco Chemie B. V., Uttrecht, The Netherlands). An automated hanging mercury drop electrode (HMDE, Metrohm, Switzerland) was used as a working electrode, Ag/AgCl/3 mol L−1 KCl as the reference electrode and a Pt coil as the auxiliary electrode. SAS measurements of the LS samples before and after irradiation were performed by re-dissolving the dry lipid aliquot in 0.55 mol L−1 NaCl (Suprapur®, Merck, Germany) as the supporting electrolyte (see 2.3.2. Aqueous Phase analysis).

### 2.2.3. Lipid classes

Aliquots of the lipid extracts re-dissolved in dichloromethane were spotted onto silica-coated quartz thin-layer chromatography rods, where they were developed in a series of seven developing baths containing mixtures organic solvents of increasing polarity. Eighteen lipid classes were separated and analyzed by thin–layer chromatography flame ionization detection (TLC–FID) (Iatroscan MK–VI, Iatron, Japan), at a hydrogen flow of 160 mL min−1 and air flow of 2000 ml min−1, and quantified by external calibration with a standard lipid mixture. More experimental details can be found in Gašparović et al (2015, 2017). Limits of detection (LODs) were determined as the analyte concentrations corresponding to a signal-to-noise (S/N) ratio of 3. The sample was analysed in triplicate. Total lipid concentration was determined as a sum of all quantified lipid classes: sterol esters (SE); fatty acid methyl esters (ME); fatty ketone hexadecanone (KET, internal standard); triacylglycerols (TG); free fatty acids (FFA); fatty alcohols (ALC); 1,3-diacylglycerols (1,3 DG); sterols (ST); 1,2-diacylglycerols (1,2 DG); pigments (PIG); monoacylglycerols (MG); three classes of glycolipids (GL): mono- and di-galactosyldiacylglycerols (MGDG and DGDG) and sulfoquinovosyldiacylglycerols (SQDG), and three classes of phospholipids (PL): phosphatidylglycerols (PG), phosphatidylethanolamines (PE), and phosphatidylcholine (PC).

### 2.2.4. Chlorophyll *a*

Subsamples (500 mL) for the determination of chlorophyll *a* (Chl *a*) were filtered on 0.7 m Whatman GF/F glass filters and stored at −20 ℃ until further processing. Spectrometric determination of the Chl *a* concentration was determined according to ISO 10260:1992 method with minor modifications. Samples were extracted in 95% ethanol (10 mL) at 60 ℃ for 1 h in dark, centrifugated (Rotofix 32A, Hettich) for 10 min at 4000 rpm and absorbance was measured on a Turner TD-700 spectrophotometer at 750 nm and 665 nm before and after the addition of 10 L 2 mol L−1 HCl.

## 2.3. Photochemical reactor experiments

Phytoplankton culture samples TS and DS (7 mL) were placed into a round Pyrex reactor cell, thermostated to 20 ℃, and mounted with Quartz windows. The experiment with the lipid sample (LS) was conducted by spreading the hexane lipid extract aliquots, with a syringe (Hamilton, Nevada, USA), on the surface of a seawater model solution (0.55 M NaCl). The system was left for 15 min under very slow flow, in order for the hexane to evaporate, obtaining in this way a lipid film at the air-water interface. The surplus of lipid material is added to ensure a fully covered surface (near-monolayer coverage) due to losses on the cell walls. A Xenon lamp (150 W; LOT Quantum Design, France) placed at a distance of 13 cm from the reactor was used to mimic solar irradiation on the Earth’s surface (see Fig. S5 in the study of Ciuraru et al. (2015a)). A quartz water filter of 5 cm was mounted in front of the lamp to remove infrared irradiation, while short wavelengths (λ < 290 nm) were eliminated by a Pyrex filter positioned directly in front of the reactor. A flow of 200 – 300 mL min-1 of compressed, filtered and purified air was pushed continuously through the reactor. Typically, after the introduction of the samples and acquisition of a stable background signal, the samples were irradiated for 1 h. Experiments were stopped when signals reached background levels again, after switching off the light. All experiments were preceded by background experiments performed on an empty reactor containing ultrapure water. It should be noted that temperature effect can be excluded in our experiments as temperature control tests have been performed. VOCs fluxes were calculated according to Ciuraru et al. (2015b), where light intensity of the Xe lamp, assuming that only the UV-A fraction is responsible for the observed photochemistry, was 8 mW cm-2 for wavelengths below 400 nm. The mean solar flux used was 21 mW cm-2.

### 2.3.1. Gas Phase analysis

A selected reagent ion-proton transfer reaction-time-of-fight mass spectrometer (SRI-PTR-ToF-MS 8000, Ionicon Analytik GmbH, Innsbruck, Austria) was used to quantify the emitted VOCs using H3O+ as source reagent ions. More details about the setup and calibration of the PTR-MS were described in a previous study (Kalalian et al., 2020). Briefly, air was sampled at a constant flow of 50 ml min–1 at an inlet temperature of 333 K. Typically, a drift voltage of 550 V, a drift temperature of 353 K and a drift pressure of 2.2 mbar were used, resulting in an E/N-ratio of about 135 Td.

### 2.3.2. Aqueous Phase analysis

Immediately after irradiation experiments TS and DS sample aliquots were frozen at -20 ℃ for further DOC and surfactant analyses (described above). An aliquot of irradiated LS sample was extracted twice with dichloromethane. Extract was evaporated to dryness by rotary evaporation under a nitrogen atmosphere and stored at −20 ℃ until SAS measurements.

## 2.4. Atmospheric simulation chamber

Particle formation upon oxidation of the VOCs originating from the LS sample was investigated using a 2 m3 chamber made of fluorinated ethylene propylene (FEP) film. The chamber was surrounded by 12 UV-Vis lamps (OSRAM lamps, Eversun L80W/79-R) to mimic solar irradiation. When all chamber background levels were established, the filling of the chamber with VOCs produced from LS sample irradiation in quartz reactor cell was conducted. After 48 h of filling, the ozone, generated by UV photolysis of oxygen (Stable Ozone Generator 1, Ultra-Violet Products Ltd., USA), was injected in the dark, reaching a maximum of 300 ppbV. Gas-phase concentration of VOCs was monitored with a PTR-ToF-MS using H3O+ as source reagent ions. The resulting O3 concentrations in the chamber were continuously monitored by Ozone Analyser (49i, Thermo Scientific, USA). Particle number concentrations and corresponding size distributions were measured using an ultrafine condensation particle counter (UCPC, 3776, TSI, USA) and a scanning mobility particle sizer (SMPS, 3936, TSI, USA). After the particle concentrations reached an approximate steady state concentration, the UV lamps were turned on for 1 h to further study SOA ageing.

# **3. Results and discussion**

## 3.1. Initial characteristics of phytoplankton culture sample

At the onset of the stationary growth phase, the concentrations of POC, DOC, Chl *a*, dissolved lipids, and lipid carbon content of the dissolved fractions of the phytoplankton culture sample were determined and results are presented in **Table 1**.

**Table 1.** Initial properties of phytoplankton culture sample.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Chl *a*  mg L−1 | DOC  mg L−1 | POC  mg L−1 | DLip  mg L−1 | DLipC  mg L−1 |
| 25.9±3.1 | 2.9±0.1 | 6.2±0.2 | 1.2±0.1 | 0.8±0.1 |

The high POC concentration of 6.2 mg L−1 is due to abundant cell content (4 x 106 cells), which is also evident from the high Chl *a* concentration of 25.9 g L−1. Obtained DOC and POC concentrations were within the range of mean values typically detected in the sea surface microlayer of Adriatic Sea coastal area (DOC 1.5 mg L−1 and POC 3.5 mg L−1) (Milinković et al., 2022). Marine lipids are an important component of marine OM (Frka et al., 2011), with the dissolved fraction (DLip) originating from: primary production and release during life cycle, after cell death, or from dissolution of the particulate fraction (Yoshimura et al., 2009; Novak et al., 2018). Marić et al. (2013) reported that dissolved lipids in the northern Adriatic Sea ranged from 10.3 to 70.6 µg L−1, and were dominated by glycolipids, [phospholipids](https://www.sciencedirect.com/topics/earth-and-planetary-sciences/phospholipid) and [free fatty acids](https://www.sciencedirect.com/topics/agricultural-and-biological-sciences/free-fatty-acids). Increased sea temperatures facilitate marine DOC and DLip production, with more pronounced effects under nutrient limiting conditions (Novak et al., 2019, 2018). The amount of carbon in the extracted LS (**Table 1**) was determined based on the assumption that lipid carbon accounts for 70% of total lipid concentration (Novak et al., 2018). The contribution of DLip to DOC in the phytoplankton culture sample was approximately 28%. Dissolved lipid material was dominantly composed of polar lipids, i.e., phospholipids (PL) and glycolipids (GL) (**Figure 1**). The two groups accounted for 35 ± 6% and 21 ± 3% of the total DLip, respectively. Phosphatidylglycerols (PG) dominated within PL while monoacylglycerols (MDGD) contributed the most to GL. Dissolved PL are assumed to be a part of the non-living organic matter released from phytoplankton cells upon cell death since PL are essential components of membranes where they share a structural function with sterols. In algae and cyanobacteria, like in higher plants, GL are located predominantly in the photosynthetic membranes (thylakoids), where they are the most abundant type of lipids (Guschina and Harwood, 2009). It was found that phytoplankton adapted to low nutrient conditions enhanced MGDG synthesis because these molecules do not contain nitrogen or phosphorus (Frka et al., 2011). The most common glycolipids in plankton are MGDG, digalactosyldiacylglycerols (DGDG) and sulfoquinovosyldiacylglycerol (SQDG) although mannose, glucose and glucuronic acid are also found as constituent sugars in GL (Guschina and Harwood, 2009). High abundance of MGDG and PG indicates dominance of freshly produced OM, while the non-living material signifies the more re-worked lipid degradation indices (FFA, MG, 1,2- and 1,3-DG). FFA as indicators of lipid material degradation accounted for 12 ± 2% of lipids, while remaining lipid classes contributed by only 7% or less.



**Figure 1.** The composition of lipid classes in the dissolved fraction of *C. pseudocurvisetus* culture.

## 3.2. Photochemical transformation of biogenic material and VOCs production

Upon irradiation of biogenic material present in the three different samples originating from the phytoplankton culture, rapid VOCs release was detected. Since all samples contained only non-living material, direct biological mechanisms can be assumed to have had no effect on the VOCs production, meaning that all VOCs were formed through abiotic photochemical reactions. The total VOCs fluxes normalized per gram of carbon for TS, DS and LS samples are shown in **Figure 2**. Higher normalized VOCs emission flux measured in TS compared to DS shows that particulate material from the dead cells present in TS may be responsible for the enhanced release of VOCs precursors in addition to the dissolved biogenic material. This is consistent with previous findings showing that lysis of riverine microbial cells due to cell death was a significant source of surfactant compounds acting as VOC precursors (Brüggemann et al., 2017). Indeed, particles have a role in facilitating the accumulation and transport of surface active material, including lipid material, to the ocean-atmosphere interface. Therefore, the enrichment of the particulate organic fraction in the sea surface microlayer is typically greater than the enrichment of the dissolved fraction (Marty et al., 1988; Kuznetsova and Lee, 2002; Kuznetsova et al., 2005; Burrows et al., 2014, Milinković et al., 2022)(Marty et al., 1988; Kuznetsova and Lee, 2002; Kuznetsova et al., 2005; Burrows et al., 2014, Milinković et al., 2022).

The highest normalized flux intensity was obtained for the LS sample. This implies that biogenic lipid material homogenously distributed at the air-water interface (i.e., on a fully covered surface), is primarily responsible for the photochemical reactions leading to the production of VOCs from diatom OM during irradiation. Since the LS lipids present a fraction of the DS and TS organic material, we can assume that due to experimental conditions (insufficient time, no bubbling), lipid material from bulk of TS and especially of DS, did not accumulate sufficiently at the air-water interface to form homogeneous surface films and enable more intense VOC production. Namely, it was observed previously that the presence of lipids at the ocean-atmosphere interface, i.e. the sea surface microlayer, is the result of its competitive adsorption and segregation from other macromolecular constituents (Frka et al., 2012), and that the formation of a lipid surface film, as well as its properties, are strongly affected by other organic material present (Kozarac et al., 2000).



**Figure 2**. VOC fluxes in TS, DS and LS normalized per gram of carbon. Yellow colour represents the period of irradiation.

Photochemical processing of the three samples exposed to light was followed by comparing their surfactant activity before and after irradiation (**Figure 3**). As observed, irradiation induced a significant increase in the surface activity of the organic material present in TS, DS and LS samples. An increase in surface activity upon irradiation of the SML and underlying water was recently reported by Rickard et al. (2022), showing that in addition to temperature, which is known to affect surfactant adsorption kinetics, irradiation independently contributes to enhanced interfacial surfactant activity due to photo-degradation of larger organic molecules. The less water-soluble substances i.e., surfactants present in the bulk of the TS and DS samples might spontaneously accumulate at the air-water interface and drive the observed photo-induced VOCs production. Furthermore, our results confirmed photo-induced increase of OM surface activity, which additionally facilitated OM accumulation at the air-water interface and further simultaneous formation of the detected VOCs.

The highest normalized flux intensity originated from the LS sample, also characterized by the highest surface activity, pointing to the importance of the photochemical processing of biogenic lipids as efficient surfactants in the VOCs formation in the marine environment.



**Figure 3**. Surface activity of TS, DS and LS samples measured before and after irradiation.

The notion of omnipresence of aerosol bound surfactants from ocean waters has been supported by recent laboratory studies linking aerosol production and fatty acid composition with marine biological activity (Alpert et al., 2015, 2017, Cochran et al., 2016). This is especially relevant because surfactants on water aerosol particles and droplets may undergo unique photochemical reactions upon irradiation (Rossignol et al., 2016, George et al., 2015), due to distinctly different physical and chemical properties of interfaces, compared to bulk and gas phases.

Another driving force of photochemical reactions includes photosensitizers. TS and DS contained photosensitizing organic molecules, as well as traces of photo-active transition metals such as Fe and Cu, which could have additionally initiated the observed photochemistry (Rossignol et al., 2016). Photosensitized reactions were also possible in the LS sample, where pigments that could act as photosensitizers, accounted for 5 ± 3% of the total lipids (**Figure 1**). More specifically, chlorophylls are very efficient photosensitizers under both PAR (Nelson, 1993) and UVR (He and Häder, 2002) since they absorb in the UVR range (Quesada and Vincent, 1997; Ehling-Schulz and Scherer, 1999) and enable type II (i.e. involving singlet oxygen) oxidation processes (Rontani, 2001). Christodoulou et al. (2010) showed that photodegradation of most of the unsaturated lipids of E. huxleyi cells (chlorophylls, sterols and monounsaturated fatty acids) involved type II photosensitized processes, induced by both UVR and PAR. UVR exposure also induced photosensitized stereomutation (*cis-trans* isomerization) of double bonds of some lipids, leading to the production of compounds sufficiently specific to act as tracers of UVR-induced photodegradation in the natural environment.

A decrease in VOCs concentration was observed prior to irradiation in TS sample, indicating a potential removal of gaseous compounds present, as the stream of air is flushing the area above the sample and introducing it into the PTR-ToF-MS. The potential de-gassing of the TS sample was particularly related to C3H6O (*m/z* = 59.048), attributable to **acetone,** whose concentration decreased prior to irradiation was followed by an immediate increase when irradiation started, which continued to increase for a while, before again showing a decreasing trend (**Figure S1).** C3H6O was not detected in DS, as particulate material producing it in TS might have been removed during the filtration of the sample. C3H6O formed from LS exhibited different behaviour compared to TS, indicating different processes or material present were responsible for C3H6O formation in different types of biogenic samples. Thus, constant concentrations prior to irradiation were followed by a gradual increase upon irradiation (**Figure S1**) indicating continuous production of C3H6O by abiotic processes originating from lipid material present at the air-water interface. Along with C3H6O, saturated oxygenated VOCs, such as aldehydes and ketones, as well as unsaturated and functionalized VOCs (e.g., alkenes, dienes, unsaturated aldehydes or ketones) were photochemically produced from the analysed samples (**Table 2**). Most of the VOCs were previously observed in photochemical studies of marine SML and biofilm samples and surfactant-containing aqueous solutions, and attributed to photochemistry at the air-water interface - photosensitized reactions as well as photo-activity of organic surfactants without the presence of photosensitizers (Brüggemann et al., 2017; Rossignol et al., 2016; Ciuraru et al., 2015a, 2015b; Chiu et al., 2017). Photochemical activity and VOCs production thus occur in an organic-enriched environment, typically found at the air-water interface of surfactant-containing aqueous solutions, since formation of mono- and poly-unsaturated compounds is not favoured in the bulk water.

The highest number of VOCs originated from the LS (**Table 2**), where along with C3H6O, C2H4O (*m/z* = 45.033) and CH4O (*m/z* = 33.033), attributable to acetaldehyde and methanol, respectively, significantly contributed to oxygenated VOC fluxes. These compounds are ubiquitous in the atmosphere, affecting the ozone budget of the troposphere, acting as precursors to peroxy acetyl nitrate and presenting significant sinks for hydroxyl radical and the oxidizing capacity of the lower atmosphere, especially in remote marine areas (Lewis et al., 2005; Singh et al., 1995). Photochemical processes, including photodegradation of chromophoric dissolved organic matter (CDOM) are responsible for the formation of acetaldehyde and acetone (Brüggemann et al., 2017; Mopper and Stahovec, 1986; Zhou and Mopper, 1997), with estimations in coastal upwelling locations ranging between 16% and 68% for acetaldehyde and 48 - 100% for acetone, whose production was shown to be higher in light conditions (Dixon et al., 2013). On the other hand, photochemical production of methanol is considered insignificant and mostly of biological origin, based on measurements and modelling (Beale et al., 2015; J. L. Dixon et al., 2011a; Dixon et al., 2011b; Millet et al., 2008; Mincer and Aicher, 2016). However, our data indicate that alongside C2H4O(acetaldehyde) and C3H6O (acetone), phytoplankton derived lipid material present at the air-water interface could also be a source of photochemically produced CH4O(methanol). Formation of several other aldehydes was also observed during the irradiation of LS, namely CH2O (*m/z* = 31.018), C4H8O (*m/z* = 73.063) and C6H12O (*m/z* =101.096), attributable to formaldehyde, butanal and hexanal respectively. Kieber et al. (1997) observed aldehydes in fatty acid/triglyceride-containing seawater samples exposed to sunlight, attributing their origin to photooxidation of linoleic acid and trilinolein. Photooxidation rate for the unsaturated fatty acid and triglyceride was determined to be over 10 times greater compared to the mono- unsaturated species, while photodegradation of e.g. palmitic acids was considered negligible - potentially due to its low concentration at the air-water interface (experimental details do not mention whether or not an interface was present). According to Zhou et al. (2014), the heterogeneous reaction of linoleic acid with O3 produces n-hexanal (C6H12O) and 3-nonenal (C9H16O), which is then degraded tomalondialdehyde (C3H4O2) and further to glyoxal (C4H2O2). C6H12O, C9H16O and C3H4O2, detected during the irradiation of LS, indicate that these highly reactive dicarbonyls might potentially originate as secondary products from *C. pseudocurvisetus* fatty acid photodegradation, as FFA present 12% of lipid material of LS samples. FFA are often found as one of the dominant lipid classes in dissolved lipid fraction, and are considered breakdown indices (Novak et al., 2018; Penezić et al., 2010).

C5H8 (*m/z* = 69.069), potentially attributable to *isoprene*, bearing in mind all possible interferences, was detected during irradiation of all samples studied. This assignment was made with respect to our previous work where different analytical tools were used (Ciuraru et al., 2015b). Isoprene has been determined in marine air across the world’s oceans (Yu and Li, 2021 and references therein), and although its terrestrial production (400–750 TgC yr−1) exceeds the marine one (0.1–1.9 TgC yr−1) (Arnold et al., 2009; Palmer and Shaw, 2005, Arneth et al., 2008; Guenther et al., 2006, 2012; Carslaw et al., 2000), isoprene of marine origin has a significant impact on SOA formation, especially during phytoplankton blooms in remote oceans (Hu et al., 2013). This is due to isoprene relatively short atmospheric lifetime (approximately 30 min – 1 hour), making its transport from terrestrial sources to marine areas insignificant. In coastal areas of the eastern Mediterranean, an increase in the concentration of isoprene and other VOCs has been observed during the daytime and with increasing temperature (Kameyama et al., 2014, Dayan et al., 2020), suggesting both biological and abiotic photochemical processes involvement. C5H8 fluxes during irradiation of phytoplankton-produced OM in this study were on average 5.7 x 109 molecule cm−2s−1 for the TS sample and 3.1 x 109 molecule cm−2s−1 for the DS sample. The C5H8 flux from the LS sample was the highest observed, and showed a constant increase upon being irradiated, from 7 x 109 molecule cm−2s−1 to 15 x 109 molecule cm−2s−1, with an average of 14 x 109 molecule cm−2s−1 during irradiation. In-situ measured isoprene fluxes ranged between 0.017 – 6 x 109 molecule cm−2s−1 (Shaw et al., 2010), while an authentic SML sample studied in laboratory conditions produced isoprene in the range between 7 – 230 x 109 molecule cm−2s−1 (Ciuraru et al., 2015b). Additionally, studies showed that biotic isoprene production rate by different diatom species can range between 1.12 – 28.48 μmol (g Chl a)−1 day−1 (Booge et al., 2016 and references therein), which is close to the average C5H8 production rate of 33 ± 3 µmol g Chla−1 day−1, obtained in our study for the TS sample. As the VOCs production in our samples can be considered exclusively abiotic, these results indicate the significant contribution of the biogenic lipid film processing to the overall isoprene production in the marine environment.

Along with C5H8, C10H16 (*m/z* = 137.129), attributable to another terpene, was detected originating from the LS sample. Monoterpenes, whose environmental concentrations are lower than those of isoprene, can have a disproportionately significant impact on marine aerosol, due to the higher yield of SOA (Yu and Li, 2021). Reaction of monoterpenes with molecular oxygen, ozone, hydroxyl radicals, NOx species and chlorine atoms leads to their atmospheric transformation to condensed-phase carbonyls, alcohols, esters, halogenated hydrocarbons and peroxynitrites within hours (Marmulla and Harder, 2014). It was experimentally observed that α-pinene and β-pinene had a SOA yield of 32% and 41% respectively (Luo and Yu, 2010; Yu, 2018), however their significance, especially considering abiotic processes for the global SOA production needs to be studied further (Yu and Li, 2021, Spracklen et al., 2008). Furthermore, C2H4O2 (*m/z* = 61.027) and C4H6O2 (*m/z* = 87.042), attributable to acetic and methacrylic acid, respectively, were also detected during LS sample irradiation, potentially due to monoterpene oxidation in the presence of OH radicals (Friedman and Farmer, 2018), which could have been formed by direct dissociation with C9 aldehydes (Rossignol et al., 2016).

**Table 2**. Detected VOCs originating from TS, LS and DS samples.

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| (*m/z*), assigned molecular formula | TS | DS | LS | (*m/z*), assigned molecular formula | TS | DS | LS |
| ppbv | | | ppbv | | |
| (31.018) CH2O | – | – | 0.3 | (57.069) C4H8 | – | 0.9 | 5.8 |
| (47.012) CH2O2 | – | – | – | (73.063) C4H8O | – | – | 0.9 |
| (46.028) CH3NO | – | – | – | (71.083) C5H10 | 0.3 | – | 1.9 |
| (33.033) CH4O | – | – | 2.4 | (87.077) C5H10O | – | – | 0.8 |
| (43.017) C2H2O | – | – | 1.7 | (69.069) C5H8 | 0.7 | 0.3 | 1.0 |
| (42.032) C2H3N | – | – | – | (85.063) C5H8O | 0.4 | – | 1.6 |
| (44.012) C2H3O | – | – | – | (101.056) C5H8O2 | 0.3 | – | – |
| (45.033) C2H4O | 5.1 | 7.1 | 8.4 | (81.067) C6H8 | 0.6 | – | 0.5 |
| (61.027) C2H4O2 | – | – | 1.1 | (83.083) C6H10 | 0.4 | – | 0.6 |
| (60.049) C2H5NO | – | – | 2.1 | (85.099) C6H12 | – | – | 1.4 |
| (63.040) C2H6O2 | – | – | 0.2 | (97.063) C6H8O | – | – | 0.1 |
| (41.038) C3H4 | 1.0 | 0.6 | 2.2 | (99.078) C6H10O | 0.4 | – | 0.5 |
| (57.034) C3H4O | – | – | 0.4 | (101.096) C6H12O | – | – | 0.3 |
| (73.027) C3H4O2 | – | – | 0.2 | (97.099) C7H12 | – | – | 0.6 |
| (43.054) C3H6 | 1.7 | 0.9 | 4.3 | (109.065) C7H8O | 0.2 | – | – |
| (59.048) C3H6O | 125.9 | – | 60.9 | (111.077) C7H10O | – | – | 0.1 |
| (75.042) C3H6O2 | – | – | – | (111.115) C8H14 | – | – | 0.4 |
| (58.072) C3H7N | – | – | 0.4 | (127.1100 C8H14O | – | – | 0.2 |
| (101.021) C4H4O3 | – | – | – | (141.124) C9H16O | – | – | 0.5 |
| (54.046) C4H6 | 0.7 | – | – | (137.129) C10H16 | – | – | 0.3 |
| (71.047) C4H6O | 0.3 | – | 0.5 | (151.107) C10H14O | 0.2 | – | – |
| (87.042) C4H6O2 | – | – | 0.2 | (155.140) C10H18O | – | – | 0.5 |

## 3.3. Atmospheric simulation chamber experiment

The highest concentration and number of produced VOCs was observed during irradiation of LS sample, leading to further investigation of the impact of biogenic lipids on the atmosphere’s oxidation potential, aerosol particle formation and growth. LS sample was irradiated within the photochemical reactor for a period of 48 h, during which the produced VOCs were directly introduced into a dark atmospheric simulation chamber. After 48 h, the irradiation was stopped and VOCs in the chamber were analysed by PTR-ToF-MS, revealing a smaller number of compounds compared to those previously determined in the LS sample (**Table S1**). This is possibly due to wall losses as well as chemical reactions that took place during the filling of the chamber, reducing the number of compounds and leading to the formation of new ones such as CH2O2, CH3NO, C2H3N, C2H3O, C3H6O2, C4H4O3, C4H8O, C5H8O2.

Following introduction of ozone, a sharp increase of several VOCs was observed (**Figure 4a**), most pronounced for C2H4O (acetaldehyde) and C3H6O (acetone), likely due to ozonolysis of organic unsaturated compounds (Conrad et al., 2021; Deng et al., 2018; Millet et al., 2010; Singh et al., 1994). The reaction between ozone and unsaturated organic compounds allows cleavage of unsaturated carbon bonds, facilitating formation of a carbonyl component and a carbonyl oxide as one of Criegee intermediates, and even carboxylic acids (Hammes et al., 2018; Wegener et al., 2007). The Criegee intermediates formed by ozonolysis are oxidative species (Conrad et al., 2021) which react, among others, with organic acids in the atmosphere, leading to formation of secondary organic aerosols (Chhantyal-Pun et al., 2018). Following O3 introduction, a potential increase in reactive Criegee intermediates could have caused different reactions among the already present and the newly produced VOCs, leading to significant particle formation within approximately 40 min after a maximum O3 concentration of around 300 ppb was reached (**Figure 4b**).



**Figure 4**. Concentration of volatile organic compounds in the multiphase atmospheric simulation chamber originating from the dissolved lipid sample at the air-water interface, with irradiation period indicated by the yellow colour (a) and the size distribution and number of particles formed within the chamber (b).

A steady increase in particle numbers continued for another 40 min, reaching maximum values. After an initial growth phase when small-sized particles with a diameter of around 10 nm were observed, larger particles began to form, reaching average diameters of around 25 nm. According to previous studies, the formation of such small and stable aerosol particles possibly indicated the formation of very stable and low volatile acid–base clusters, or even reaction products (Tang et al., 2013; Qiu et al., 2013; Duporté et al., 2016). This is further supported by the fact that nitrogen-containing compounds are among the VOCs produced and may act as bases in such reactions (**Table S1**). To study the effect of OH-oxidation and SOA aging, UV lights were turned on after 160 min, and an immediate particle growth was observed (**Figure 4b**). The UV light facilitated the oxidation of the yet unreacted VOCs as well as further low-volatile compound formations, which condensed on the already existing particles, causing particles to increase in size.

## 3.4. Environmental relevance

In this study we showed that biogenic lipids of phytoplankton origin, as highly surface-active compounds present at the air-water surface, could be exclusively responsible for the photochemical production of VOCs in the marine atmosphere. The VOCs fluxes determined from the lipid sample were within the range of previous field observations and laboratory experiments, suggesting that the observed processes could occur to a similar extent in the marine environment, particularly within the SML directly exposed to the sunlight. It should be noted that such VOCs emissions have previously been attributed to processes driven by biology, artificial surfactants in the laboratory, and biofilm-containing solutions with different microorganisms. In contrast, the observed results suggest that VOCs released during irradiation of exclusively biogenic lipid material of phytoplankton origin accumulated at the ocean-atmosphere interface might lead to similar emissions of saturated and unsaturated VOCs and SOA precursors, impacting atmospheric budgets of cloud condensation nuclei, especially important in clean marine environments. On the other hand, the injection of surfactant films present at the sea-atmosphere interface into the atmosphere in an enriched form as part of the marine aerosol produced by bursting bubbles could occur in real marine conditions. This provides a mechanism for selective transfer of biogenic fatty acids and other marine lipids from the sea to the atmosphere, where photochemistry occurring on the surface of marine aerosol particles may also be a source of VOCs as well as SOA precursors, which could help close the gap for calculating organic aerosol concentrations which are consistently being under-predicted in marine environments. Moreover, such atmospheric processing could result in a change of aerosol surfactant content, directly affecting cloud activation and droplet number, especially important in clean marine atmosphere (Kroflič et al, 2018). There is also increased evidence of the importance of biogenic surfactant films in the air-sea exchange of climate-active gases. The viscoelastic behavior of the air-sea interface, a key parameter affecting air-sea exchange of gases is strongly dependent on naturally occurring adsorbed surfactant material. Namely, natural surfactant films reduce the air-sea gas transfer velocity (kw) of CO2 and other gases by up to 50% (Frew, 2005; Salter et al., 2011; Pereira et. al., 2016; 2018; Milinković et al 2022), and photochemical changes to lipid-like surfactant material can modify kw as well as marine boundary layer chemistry.

As a consequence of anthropogenic activity and climate change, many properties of the oceanic environment are changing rapidly. Phytoplankton as a primary producer is highly dependent on changing environmental conditions because of its adaptive evolution strategies aimed at survival. Important adaptation mechanism involves changes in their lipid composition. For example, enhanced accumulation of storage lipids (TG) was observed during nitrogen stress for the marine diatom (Parrish and Wangersky, 1987), during bloom decay (Parrish, 1987) and during oligotrophication in the NW Mediterranean (Bourguet et al., 2008). Increasing oligotrophy in the northern Adriatic Sea led to an increasing number of phytoplankton taxa, enhanced lipid biosynthesis and increasing synthesis of glycolipids (MGDG), as these molecules do not contain nitrogen or phosphorus (Frka et al., 2011). Similarly, phytoplankton may overcome phosphorus deficiency by synthesis of sulfoquinovosyldiacylglycerol, a lipid with sulfur and sugar in place of phosphate (Van Mooy et al., 2006). Moreover, intensive wildfire episodes which are expected to increase in future qualitatively and quantitatively modify lipid production of surface layers in the coastal environment (Milinković et al 2022). Therefore, we may expect that future scenarios would result in qualitative and quantitative changes of biogenic lipid pool, its accumulation and photo-transformation at the air-water interface with global consequence. This study thus clearly points toward a global impact of the biogenic lipid photochemistry at the air-sea interface, with in depth studies of the lipid remodelling, caused by environmental changes, present the next essential step in understand the drivers behind the difference in their VOCs production, as well as their impacts on the atmospheric processes.

# **4. Conclusions**

For the first time, photochemical processing at the air-water interface containing organic material from an authentic culture of the marine diatom *Chaetoceros pseudocurvisetus* was studied. Upon irradiation, the total and dissolved phytoplankton material, and in particular its isolated lipid fraction present at the air-water interface, acted as a source of abiotic production of saturated oxygenated compounds as well as unsaturated and functionalized VOCs, while OM photochemical transformation was followed by changes in surfactant activity of exposed biogenic material. Our findings confirm that particulate material from dead cells may play a role in facilitating the accumulation of biogenic surfactants at the air-water interface. The highest normalized flux intensity determined for the lipid material suggests that biogenic lipids, as less water-soluble substances, i.e., efficient surfactants present at the air-water interface, are primarily responsible for the photochemical reactions leading to the production of VOCs from diatom OM exposed to irradiation. The VOCs fluxes determined from the lipid sample were within the range of previous field observations and laboratory experiments, suggesting that the observed processes could occur to a similar extent in the marine environment, particularly in the sea surface microlayer. It should be noted that such VOCs emissions have previously been attributed to processes driven by biology, artificial surfactants in the laboratory, and biofilm containing a mixture of different microorganisms. In contrast, the observed results suggest that VOCs released during irradiation of exclusively biogenic lipid material accumulated at the ocean-atmosphere interface might lead to similar emissions into the atmosphere. Additionally, we demonstrated that the emitted VOCs can significantly affect the oxidation potential of the atmosphere and promote aerosol formation and growth. Oxidation by dark ozonolysis and subsequent particle formation clearly demonstrate the formation and release of unsaturated VOCs. Given that marine biogenic lipids as common natural surfactants can be considered ubiquitous at air-water interfaces, such as on rivers, lakes, oceans, aerosol particles, and cloud droplets, photo-induced abiotic VOCs production can be expected to occur globally, influencing ocean-atmosphere exchange processes as well as atmospheric chemistry on a large scale.

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