

# Marine Organic Aerosols at Mace Head: Effects from Phytoplankton and Source Region Variability

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## 2. Abstract

Organic aerosols (OA) are recognised as a significant component of particulate matter (PM), yet, their specific composition and sources, especially over remote areas remain elusive due to the overall scarcity of high-resolution online data. In this study, positive matrix factorisation was performed on organic aerosols mass spectra obtained from high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) measurements to resolve sources contributing to ~~the~~ coastal PM. The focus was on a summertime period marked by enhanced biological productivity with prevailing pristine maritime conditions. Four OA factors were deconvolved by the source apportionment model. The analysis revealed primary marine organic aerosols (PMOA) as the predominant submicron OA at Mace Head during summertime, accounting for 42% of the total resolved mass. This was trailed by more oxidised oxygenated organic aerosols (MO-OOA) at 32%, methanesulphonic acid organic aerosols (MSA-OA) at 17%, and locally emitted peat-derived organic aerosols (Peat-OA) at 9% of the total OA mass. [Elemental ratios \(O:C-H:C\) were derived for each of these factors: PMOA \(0.66-1.16\), MO-OOA \(0.78-1.39\), MSA-OA \(0.66-1.39\) and Peat-OA \(0.43-1.34\). The specific O:C-H:C range for MO-OOA hints at aliphatic and lignin-like compounds contributing to more oxidised organic aerosol formation.](#) The total mass concentrations of primary organic aerosols and secondary organic aerosols were overall equal and almost exclusively present in the marine boundary layer, [in agreement](#) ~~consistently~~ with previous findings. This study reveals that OA not only reflects atmospheric chemistry and meteorology – as evidenced by the significant aging of summertime polar air masses over the North Atlantic, driven by ozonolysis under Greenland anticyclonic conditions - but also serve as indicators of marine ecosystems. This is evident from MSA-OA being notably associated with stress enzyme markers and PMOA showing the typical makeup of largely abacterial phytoplankton extracellular metabolic processes. This study also reveals distinctive source regions within the North Atlantic Ocean for OA factors. MSA-OA is primarily associated with the Iceland Basin, with rapid production following coccolithophore blooms (lag of 1-2 days), while diatoms contribute to a slower formation process (lag of 9 days), reflecting distinct oceanic biological processes. In contrast, PMOA is sourced from more variable ecoregions, including the Southern Celtic Sea, West European Basin, and Newfoundland Basin, with additional contributions from chlorophytes and cyanobacteria at more southerly latitudes. Overall, these findings emphasise the need for [further](#) longer-term investigations to [fully account for](#) [further map the influence of](#) phytoplankton [s](#)-taxa variability [influence](#) on aerosol composition, and the [if](#) broader impacts on aerosol-climate interactions.

**Keywords:** Submicron Marine Aerosols, Secondary Organic Aerosols, HR-ToF-AMS, Positive Matrix Factorisation, phytoplankton

## 1 Introduction

The marine environment plays a critical role in Climate regulation through sea spray and gas-phase emissions from the oceans, via direct and indirect solar radiation effects and aerosol-cloud interactions governed by ocean biology, sea spray physicochemical properties and secondary reactions (Cochran et al., 2017). However, aerosols in the atmospheric marine boundary layer (MBL) remain a significant source of uncertainty in radiative forcing estimates (Rosenfeld et al., 2019; Wang et al., 2020) primarily due to limited knowledge about aerosol mass, chemical composition, and particle number distributions (Carslaw et al., 2017). Radiative transfer implications from aerosol-cloud interactions alone range from -2.65 to -0.07 Wm<sup>-2</sup>, contrasting with the more well-established CO<sub>2</sub> radiative forcing estimate of 1.83±0.18 Wm<sup>-2</sup> (Etminan et al., 2016). Along with this, the origin of marine organic aerosols (OA), specifically whether formed by primary or secondary processes, requires further investigation as the respective impacts from primary sea spray (Fossum et al., 2018; Ovadnevaite et al., 2011b; Xu et al., 2021) and secondary aerosols (Mayer et al., 2020; Quinn et al., 2017) on clouds formation or Atmosphere optical properties (Bian et al., 2019; Kahnert and Kanngießer, 2023; Li et al., 2023) in pristine environments is still being debated.

Primary Marine Organic Aerosols (PMOA) are part ~~consist~~ of sea spray aerosols, produced by wave breaking ~~bursting bubbles, film, jet, and spume drops~~ (Ovadnevaite et al. 2014; Veron 2015; Villermaux et al. 2022) that ~~carry sea salt particles enriched in biogenic organic aerosols and made of biogenic organic matter~~ (O'Dowd et al. 2004; Facchini et al. 2008). The majority (80 %) of fine carbonaceous particles in the clean N.E Atlantic marine atmosphere has been shown to directly originate from phytoplankton activity as reported with dual carbon isotopes analysis (Ceburnis et al., 2011). The phytoplankton-OA link is particularly well-established, yet the topic is still highly debated as no clear full-picture consensus has been reached owing to widely changing temporal and geographic fluctuations (Lawler et al., 2024; Lewis et al., 2021; Seidel et al., 2022). In addition, the specifics of how phytoplankton control OA chemical composition (Behrenfeld et al., 2019; Facchini et al., 2008; O'Dowd et al., 2015), particle ~~numbers~~ flux (Markuszewski et al., 2024; Sellegrí et al., 2023), size (Croft et al., 2021; O'Dowd et al., 2004; Saliba et al., 2019), lifespan and surface tension (Lee et al., 2020; Ovadnevaite et al., 2017; Sellegrí et al., 2021) are all the focus of intense ongoing investigations.

In a warming world, following a high-emissions scenario (RCP8.5) trajectory, climate change is projected to drastically alter the geographic and seasonal variability of phytoplankton blooms in the N.E Atlantic (Asch et al., 2019). Furthermore, long term trends already show that the N.E Atlantic has experienced major changes in phytoplankton functional diversity over the last 60 years (i.e. -5% dinoflagellates decade<sup>-1</sup> whereas vs diatoms increased by +0.1% diatoms decade<sup>-1</sup>) due to rapid warming and various environmental transformations attributable to Climate change (Bedford et al., 2020; Holland et al., 2023; Mutshinda et al., 2024). All of this strongly supports the pressing needs for further investigations on phytoplankton-aerosol interactions as environmental stressors will result in significant non-linear effects and tipping points (Ban et al., 2022; Wolf et al., 2024) .

In contrast to PMOA, marine Secondary Organics Aerosols (SOA) in the remote MBL arise from new particle formation (NPF) and are governed by other subtle chemical mechanisms. These include gas-to-particle  
85 conversion (Peltola et al., 2022; Zheng et al., 2021), oxidation of volatile organic compounds and consequent volatility reduction that leads to condensation (Hallquist et al., 2009; Kroll et al., 2018), ion-induced nucleation of biogenic particles (Kirkby et al., 2016) and fission of organic biogels (Karl et al., 2013). SOA formation occurs through various processes such as homogeneous, heterogeneous and multiple phase reactions (Marais et al., 2016; McNeill, 2015) as well photochemistry reactions (Brüggemann *et al.* 2018). While various SOA  
90 molecular classes have been identified, the complexity of SOA, which consist of thousands of multifunctional compounds (Goldstein and Galbally, 2007) including high molecular weight species and oligomers from diverse sources underscores the pressing need for continued exploration. All of this can now be partly described thanks to continuous widespread progresses in aerosol mass spectrometry (DeCarlo *et al.* 2006; Laskin, and Nizkorodov 2012).

95 ~~The present study focuses on source apportionment, aiming to delineate the sources of marine OA, particularly distinguishing between SOA and PMOA sources.~~

~~Both SOA and PMOA serve as cloud condensation nuclei (CCN) (Mayer et al., 2020), impacting cloud albedo and lifetime, leading to uncertainties in global chemistry climate models (Bellouin et al., 2020). Radiative transfer implications from such interactions range from 2.65 to 0.07 Wm<sup>-2</sup>, contrasting with CO<sub>2</sub> radiative  
100 forcing estimate of 1.83±0.18 Wm<sup>-2</sup> (Etminan et al., 2016). In pristine environments, SOA nucleation events significantly shape CCN concentrations, altering cloud radiative forcing (Liu and Matsui, 2022) but so does the presence of primary sea spray (Fossum et al., 2018). Complementing this, previous literature shows that phytoplankton activity is related to emissions of organic and sulphate particle CCN precursors (O'Dowd et al., 2015; Sanchez et al., 2021). There is an ongoing debate over the respective impacts from primary sea spray  
105 (Ovadnevaite et al. 2011; King et al. 2012; Schwier et al. 2015; Xu et al. 2021) and secondary aerosols (Mayer et al., 2020; Quinn et al., 2017) on cloud formation in pristine environments. This study, thus, aims at identifying aerosol sources and quantifying elemental ratios which can serve as a proxy for ulterior parametrisations (e.g. as done in Han et al. 2022; Li et al. 2023). The present study focuses on source apportionment in a coastal environment, with the aim to separate primary OA (POA) and SOA into their respective sources. Marine SOA  
110 sources notably include methane sulphonic acid ~~which~~<sup>that</sup> is ~~primarily~~ formed through the oxidation of dimethyl sulphide (Becagli et al., 2019; Hodshire et al., 2019; Mansour et al., 2024), and oxidised OA aerosols (i.e. carboxylic acids; Kawamura and Bikkina, 2016) ~~which~~<sup>that</sup> are a complex mixture resulting from unsaturated fatty acids<sup>s</sup> oxidation found in very diverse locations (Crippa et al., 2013; Florou et al., 2024; Nøjgaard et al., 2022). On the other hand, POA sources not only include sea spray but also potential anthropogenic influences  
115 like local biomass burning (i.e. wood, peat or charcoal) or long-range continental transport (Lin et al., 2019; O'Dowd et al., 2014; Xu et al., 2020).~~

~~The source apportionment was performed with the positive matrix factorisation (PMF) model which has been widely adopted for more than two decades now (Paatero, 1999; Paatero and Tapper, 1994) and successfully used on a wide range of different instruments; HR-Tof-AMS (e.g. Aiken et al., 2008), ToF-ACSM (e.g.. Fröhlich et al., 2015), PTR-MS (e.g. Slowik et al., 2010), EESI-ToF (e.g. Tong et al., 2022), offline filters (e.g. Maykut et al., 2003), SMPS (e.g. Nursanto et al., 2023) as well as other matrix-based measurements. Several previous~~

remote ocean HR-ToF-AM PMF studies have been carried out in the Atlantic (Crippa et al., 2013; Huang et al., 2018), Arctic (Moschos et al., 2022; Nielsen et al., 2019; Nøjgaard et al., 2022), Mediterannean Sea (Florou et al., 2024; Mallet et al., 2019), Pacific Ocean (Loh et al., 2023, 2024) and Antarctica (Giordano et al., 2016; Paglione et al., 2024; Schmale et al., 2013) which facilitates cross-sites comparability.

Finally, ~~This study seeks to relate different aerosol sources and geographical regions to phytoplankton taxonomic group simulations (Rousseaux et al. 2013), as the effects from ocean biology being currently are not unaccounted for in current climate models (Sellegrí et al., 2021)~~ ~~this study seeks to relate different aerosol sources and geographical regions to phytoplankton taxonomic group simulations (Rousseaux et al. 2013)~~. This multi-faceted approach ~~allows to place places~~ the local measurements at Mace Head into the broader context of Ocean-Atmosphere interactions, ~~allowing and the explore exploration of~~ the potential influences of marine ecosystems on atmospheric aerosols ~~leading~~ over the North Atlantic region.

## 2. Materials and Methods

### 2.1 Site Description

Mace Head (MHD) atmospheric research station is located on the west coast of Ireland (53.33°N, 9.90°W) on a peninsula exposed to open ocean air masses. These air masses, originating from a nominal clean sector (between 190° and 300°; Grigas et al. 2017) are predominantly steered by westerlies ushered by the polar jet's low-pressure systems. Importantly, open ocean air mases are mostly devoid of anthropogenic influences, with over 60% of air masses arriving at MHD classified as pristine marine (Grigas et al., 2017; Sanchez et al., 2022).

However, the remaining 40% of all the other air masses from the 360 sector exhibit varying degrees of anthropogenic influences, particularly during or just after periods of continental outflow under high-pressure regimes (Jennings et al. 2003).

This study focuses on August 2015, a summertime period characterized by heightened biological activity (Behrenfeld et al., 2019) and predominant pristine marine conditions. This specific year is also marked by the onset of the *cold blob*, with the subpolar gyre region (The North Atlantic waters south of Greenland) reaching around 2°C lower than previous long-term average owing to the slowing down of the Atlantic Meridional circulation and Greenland Ice melt (Rahmstorf et al., 2015; Sanders et al., 2022). As such these specific conditions could serve as an indication for future measurements of aerosols-phytoplankton interactions during cold blob phenomena.

### 2.2 In-situ measurements

~~Ambient submicron non refractory aerosol major species~~ ~~Aerosol chemical composition was~~ ~~were~~ monitored using an Aerodyne high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) equipped with a standard tungsten vaporizer operated at 650°C. ~~HR-ToF-AMS sampling covers vacuum aerodynamic size range of ~ 35 nm to ~ 1.5 µm (DeCarlo et al., 2006)~~. The instrument working principles have been extensively described in the literature (Canagaratna et al., 2007; DeCarlo et al., 2006). The HR-ToF-AMS used a 5 min time resolution scan on the single-reflection highly sensitive V mode configuration (mass resolution up to 3000 m/Δm) while detection limits were estimated based on the approach described by Drewnick et al (2009). Ionisation efficiency (IE), particle velocity and inlet flow were determined following standard methods while applying standard ~~relative RIE (RIE) for species~~ (Nault et al., 2023; Xu et al., 2018). The particle transmission and detection efficiency expressed as the collection efficiency (CE; Huffman et al. 2005) was corrected for

detection losses due to particle bounce and lens efficiency by applying the composition-dependent collection efficiency (Middlebrook *et al.* 2012)..

The AMS data were analysed using SQUIRREL (SeQUential Igor data RetRiEvaL) v1.65B and PIKA (Peak Integration by Key Analysis) v1.25B software packages. Sea salt was estimated based on a scaling factor of 51 of the common sea salt ion  $\text{NaCl}^+$  (m/z 57.96) (Ovadnevaite *et al.* 2012) while MSA was quantified by upscaling the  $\text{CH}_3\text{SO}_2$  (m/z 79) ion (Ovadnevaite *et al.*, 2014). Interferences from MSA on  $\text{SO}_4$  and OA were accounted for as follows:

$$\text{SO}_4 \text{ corrected} = \text{SO}_4 - \frac{\text{CH}_3\text{SO}_2 * 12.48}{\text{RIE}_{\text{SO}_4}}$$

$$\text{OA} \text{ corrected} = \text{OA} - \frac{\text{CH}_3\text{SO}_2 * 15.86}{\text{RIE}_{\text{Orgs}}}$$

An improved Ambient (I-A) method was adopted for the mass spectra elemental ratio analysis of O:C, H:C, N:C, S:C, and the OM:OC (organic matter to organic carbon) ratio (Canagaratna *et al.*, 2015). High-resolution analysis was performed on each m/z in the mass range 12–130 m/z with ion fitting applied to difference between open and closed spectra. Based on their elemental composition (C, O, H, N, S), ions were then grouped into chemical families:  $\text{C}_x$ ,  $\text{C}_x\text{H}_y$ ,  $\text{C}_x\text{H}_y\text{O}_z$  ( $z = 1$ ),  $\text{C}_x\text{H}_y\text{O}_z$  ( $z > 1$ ),  $\text{C}_x\text{H}_y\text{N}_w$  ( $w = 1$ ),  $\text{C}_x\text{H}_y\text{N}_w$  ( $w > 1$ ),  $\text{C}_x\text{S}_j$ ,  $\text{H}_y\text{O}_z$ ,  $\text{N}_w\text{H}_y$ ,  $\text{N}_w\text{O}_z$ ,  $\text{S}_j\text{O}_z$ , and  $\text{C}_x\text{S}_i$  where the indices x, y, z, w, j represent the number of C, H, O, N, S atoms, respectively.

The concentration of equivalent black carbon (eBC) was measured by a multi-angle absorption photometer (MAAP, Thermo Fisher Scientific model 5012). The MAAP operated at a flow rate of 10 L min<sup>-1</sup> and a 5 min time resolution. The transmittance and reflectance of eBC-containing particles were measured by the MAAP at two different angles to derive optical absorbance as detailed in (Xu *et al.*, 2020).

Carbon monoxide (CO) measurements were also conducted using a model RGA-3 CO analyser (Trace Analytical, Inc., CA, USA), which operates on the principle of hot mercuric oxide reduction gas chromatography (Derwent *et al.* 1994).

Ozone ( $\text{O}_3$ ) was measured [at the station](#) with an UV  $\text{O}_3$  spectrometer (Model 8810, Monitor Labs San Diego, CA), the raw voltage output was converted to concentration values based on Automatic Urban/Rural Network (AURN) calibration audits (Derwent *et al.* 2018). Finally, meteorological data were continuously recorded at the station (including rainfall, [irradiance](#)[solar radiation](#), wind speed, wind direction, temperature, relative humidity and pressure) using standard meteorological instruments and retrieved using the *worldMet* R package (station ID: 039630-99999) from the NOAA ISD website (<https://www.ncdc.noaa.gov/isd>).

### 2.3 Source apportionment

The HR organic mass spectra was deconvolved using the Positive Matrix Factorisation (PMF; Paatero and Tapper 1994; Paatero 1999) source-receptor model to investigate the various source contributions to OA. A major advantage of using HR data over unit mass resolution (i.e. ToF-ACSM studies) is the distinct differentiation of multiple ions sharing the same nominal mass, thereby allowing for a more exact characterisation of the temporal fluctuations of different ion families (e.g.,  $\text{C}_x\text{H}_y^+$ ,  $\text{C}_x\text{H}_y\text{O}_z^+$ ). The information richness in the HR-ToF-AMS datasets, as a result of the improved chemical resolution, is advantageous for

restricting the PMF solutions, minimising rotational ambiguity and results in more reliable solutions and a larger number of interpretable OA factors.

The IGOR PRO Source Finder (SoFi v6.8.1) toolkit (Canonaco *et al.* 2013) was used to run the PMF algorithm. Solutions were assessed across 2 to 12 factors using the unconstrained factors rotational Fpeak tool. Factors were 200 explored for Fpeaks (rotations) between -1 and 1 (0.1 steps). A final solution consisting of 4 factors was retained as the optimal solution based on several considerations. These include its Q/Qexp ratio value (1.38), which is tested for a range of FPEAKS and scaled residuals distribution (as recommended by Zhang *et al.* 2011). The solutions were also investigated in regard to key diagnostic plots, diurnal profiles, correlations with meaningful external tracer time series and reference mass spectra (Canonaco *et al.* 2021) extracted from the aerosol mass 205 spectrometer database (Ulbrich *et al.*, 2009).

#### 2.4 Air masses trajectory analysis

Air masses back trajectories analysis was performed using the Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) (Stein *et al.*, 2015). Meteorological data were accessed from the Global Data Assimilation System (GDAS) archived by NOAA Air Resources Laboratory. HYSPLIT was used to calculate 210 72-hour back trajectories every 3 hours with arrival starting height set to 100m above ground level. To investigate potential source regions leading to total particle mass concentrations from each resolved source, the back trajectories were gridded to  $1^\circ \times 1^\circ$  grid cells and linked to particle concentrations using trajectory source contribution functions. While common source contribution functions assume that trajectories centrelines are accurate, we focused instead on the Simplified Quantitative Transport Bias Analysis (STQBA) method which 215 considers plumes transport bias along air mass trajectories as a more robust approach.

The Boundary layer height (BLH) was determined from the fifth generation ECMWF (European Centre for Medium-Range Weather Forecasts) atmospheric reanalysis (ERA5) dataset based on the bulk Richardson number (Guo *et al.*, 2021) by mapping HYSPLIT trajectories footprint along the gridded BLH data. This was used to find the fraction of time spent over the Ocean, within the Marine Boundary Layer (MBL; altitude 220 <BLH), in the Marine Free Troposphere (MFT; altitude >BLH) and in the planetary boundary layer over land (PBL; altitude < BLH). The R package *rnaturrearth* was also used to obtain a high-resolution land mask for Ireland allowing for identification of purely marine air masses (no advection over land for at least 3 days prior to being sampled at MHD) and aided in delineating lands from oceans.

Finally, NASA Ocean Biogeochemical Model (NOBM) taxonomic group simulations (Rousseaux *et al.*, 2013) 225 for *coccolithophores*, *diatoms*, *chlorophytes* and *cyanobacteria* were used to visualise phytoplankton geographic repartition estimates as well as for lags calculations with OA similarly to O'Dowd *et al* (2015).

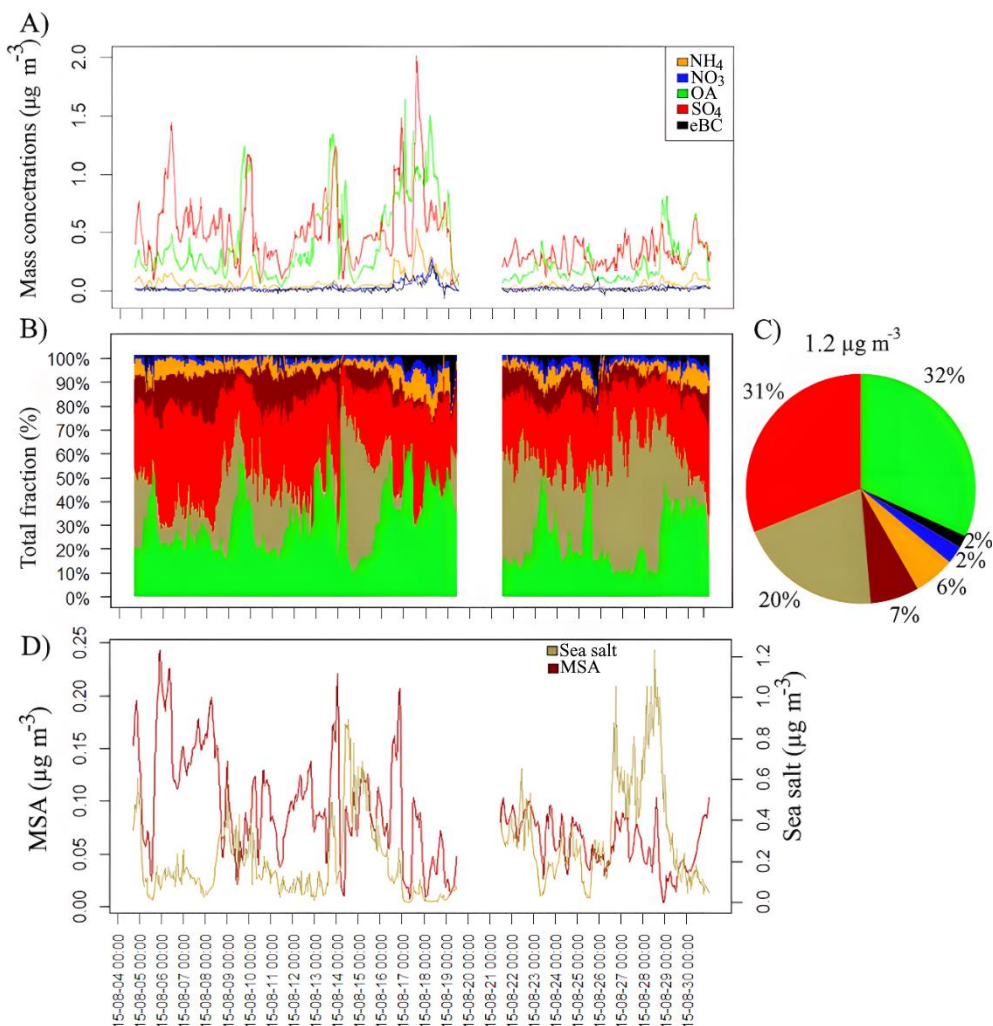
#### 2.5 Transfer entropy analysis

The R package RTransferEntropy (Behrendt *et al.*, 2019) was used to quantify the information flow between time series using the transfer entropy (TE) as previously done on recent SOA studies (Long *et al.*, 2023; Sinha *et al.*, 230 2024). Transfer entropy (TE) is a prediction model that quantifies the directional influence between two time series X and Y by determining how the past values of one series predict the future behaviour of the other (Schreiber, 2000). TE is calculated using Rényi entropy, a generalisation of Shannon entropy that offers enhanced robustness in the presence of tails effects. To account for spurious information transfer, the transfer entropy is also estimated from a shuffled version of the time series. This shuffled estimate called effective

235 transfer entropy (eTE) is used to correct for sampling bias, ensuring the validity of the results. Statistical significance is assessed with a bootstrapped Markov chain, where a p-value of less than 0.05 indicates a significant information transfer between X and Y. The reader is referred back to (Behrendt et al., 2019) for more details.

### 3. Results

240 **3.1 Submicron aerosol chemical composition overview**  
 The mass concentration time series of organic aerosols (OA), methane sulphonic acid (MSA), sulfate ( $\text{SO}_4^{2-}$ ), nitrate ( $\text{NO}_3^-$ ), ammonium ( $\text{NH}_4^+$ ) and sea salt measured by the HR-ToF-AMS as well as eBC from MAAP measurements are shown on Figure 1. The average chemical composition was dominated by OA (32%), followed by  $\text{SO}_4^{2-}$  (31%), sea salt (20%), MSA (7%),  $\text{NH}_4^+$  (6%),  $\text{NO}_3^-$  (2%) and eBC (2%) (Figure 1).



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**Figure 1.** A) OA,  $\text{SO}_4$ ,  $\text{NH}_4$ ,  $\text{NO}_3$  and eBC mass concentrations time series –  $\mu\text{g m}^{-3}$  B) Relative contributions to total PM1 C) Pie plot of total contributions to total PM1 D) shows MSA and sea salt –  $\mu\text{g m}^{-3}$ .

The total average bulk submicron aerosol mass was  $1.2 \mu\text{g m}^{-3}$  over the entire measurement period. These high 250  $\text{SO}_4^{2-}$  and OA relative contributions and overall low concentrations are common for coastal sites during summertime in the marine boundary layer as reported over the North & South Atlantic Ocean (Huang et al., 2018; Ovadnevaite et al., 2014b) as well as in the Arctic (Nielsen et al., 2019; Willis et al., 2017). MSA in

particular showed mass concentrations values of  $0.08 \pm 0.04 \mu\text{g m}^{-3}$  in the range of those previously reported at Mace Head ( $0.05 \pm 0.04$ ) (Ovadnevaite et al., 2011a) and more diverse locations such as the central Arctic (Dada et al., 2022) and the Atlantic Ocean from  $53^\circ\text{N}$  to  $53^\circ\text{S}$  where average mass concentrations of  $0.04 \pm 0.03 \mu\text{g m}^{-3}$  (Huang et al., 2018) were reported.

The low mass concentrations of  $\text{NH}_4^+$ ,  $\text{NO}_3^-$  and corresponding N:C ratio of  $0.006 \pm 0.002$  (Figure S1), indicate a limited presence of amino acids (below detection limit) from usual sources such as the North Atlantic oligotrophic gyre, ornithogenic emissions (i.e., birds), phytoplankton, bacteria, or in situ atmospheric processes (van Pinxteren et al., 2022; Schmale et al., 2013).

Following eBC thresholds established for the North-East Atlantic (Grigas et al., 2017), pristine conditions (eBC levels below  $0.015 \mu\text{g m}^{-3}$ ) were observed during 60.4% of the measurement period. Clean conditions (eBC levels between  $0.015$  and  $0.05 \mu\text{g m}^{-3}$ ) prevailed for 30.5% of the time, and moderately polluted conditions (eBC levels between  $0.05$  and  $0.3 \mu\text{g m}^{-3}$ ) occurred for 9.1% of the time with a significant pollution event spanning from August 17<sup>th</sup> to 19<sup>th</sup> 2015 onwards.

Likewise, CO mixing ratios were below 100 ppb for over 70% of the time, similarly to other pristine sites (Zhao et al., 2022). Winds advected through the clean sector ( $190\text{--}300^\circ$ ) for over 78% of the time. Finally mean wind speed was  $6.6 \pm 3.1 \text{ ms}^{-1}$  exceeding the whitecap threshold of  $4 \text{ m s}^{-1}$  (O'Dowd et al., 2014) for 77% of the time, hinting at strong- sea spray influences.

There were also SOA influences were also consequent as revealed indicated by with the average OM/OC (organic matter to organic carbon ratio) value of  $2.10 \pm 0.14$  (Figure S1), aligning with the value of 1.9 previously reported for clean aged marine polar air masses at MHD (Ovadnevaite et al. 2014). Additionally, AMS derived OM/OC values in the high Arctic (Nielsen et al., 2019) also fall within the range of 1.96 to 2.42 for PMOA (primary marine organic aerosols) and MO-OOA (more oxidised organic aerosols) respectively, here median OM/OC value was 2.11 with minimum and maximum OM/OC values of 1.71 and 2.42 respectively. This indicates the presence of both POA (i.e., saturated hydrocarbons, unsaturated hydrocarbons and cycloalkanes) as well oxygenated SOA formed with photochemical processing during long range transport (Aiken et al., 2008; Simon et al., 2011).

To get a better sense of the aerosol sources, the respective contributions of the marine boundary layer (MBL), marine free troposphere (MFT) and boundary layer over land (BL) are shown in Figure S2. Overall, the measurement period was dominated by marine boundary layer influences (91% of the time), with minimal marine free troposphere influences (8%) and extremely low land-influences from the planetary boundary layer (1%) further hinting at evidencing pristine marine conditions.

### 3.2 Source apportionment

To accurately classify and categorise the diverse sources of OA that are present at Mace Head, source apportionment was performed utilising the Positive Matrix Factorization (PMF) method on the organic mass spectrum, which ranged from m/z 12 to m/z 130. The resulting chosen four-factors solution, as depicted in Figure 2, yielded a Q/Qexp ratio of 1.38 and accounted for up to 90% of the total OA mass. Solutions with a

higher number of factors introduced splitting and did not show additional emergent interpretable sources (Figure 290 S3, Text S1).

The following four factors, namely Methane Sulphonic Acid, More Oxidised Organics, Primary Marine Organics and Peat, were determined as the optimal representation of the aerosols at Mace Head:

295 Methane Sulphonic Acid Organic Aerosols (MSA-OA): Representing approximately 17.2% of the total OA mass, MSA displayed a distinct m/z fragment at m/z 78.98 ( $\text{CH}_3\text{SO}_2^+$ ), accounting for 36.3% of its total mass spectra signal intensity. The identification of specific methane sulphonic acid tracer ions further substantiated its origin. More details on all factors are provided below in sections 3.2.1-3.2.4.

300 More Oxidised Organic Aerosols (MO-OOA): Making up about 31.8% of the total elucidated PMF solutions, this factor exhibited prominent m/z fragments at m/z 27.99 and m/z 43.99 and showed significant correlations with reference mass spectra for MO-OOA ( $R = 0.97$ ) (Hu et al., 2015), SV-OOA ( $R = 0.76$ ) (Mohr et al., 2012), and was interpreted as MO-OOA after examining elemental ratios and correlations.

305 Primary Marine Organic Aerosols (PMOA): Comprising roughly 42.2% of the total resolved PMF solutions. This factor exhibited m/z fragments similar to MO-OOA (Schmale et al., 2013), but with higher contributions from aliphatics ( $\text{C}_x\text{H}_y$ ) such as alkyls (dominant in sea spray during phytoplankton blooms; Cavalli et al. 2004), alkenes (i.e. phenols or humic materials; Bahadur et al. 2010) and functional derivatives such as alcohols ( $\text{C}_x\text{H}_y\text{O}_z$ , where  $z=1$ ) as established in earlier studies (Ovadnevaite et al. 2011; Crippa et al. 2013).

Peat-OA: accounting for approximately 8.8% of the total PMF solutions, was characterised by saturated hydrocarbons ( $\text{C}_x\text{H}_{2y+1}$ ), unsaturated hydrocarbons ( $\text{C}_x\text{H}_{2y-1}$ ) and cycloalkanes ( $\text{C}_x\text{H}_{2y}$ ) ion series. This factor was identified as Peat-OA owing to its good correlation ( $R=0.75$ ) with the Peat-OA reference mass spectra (Lin et al., 2017). Additionally, its mass spectrum was marked by cellulose pyrolysis fragments typical for 310 levoglucosan (i.e.  $\text{C}_2\text{H}_4\text{O}_2^+$  at m/z 60 and  $\text{C}_3\text{H}_5\text{O}_2^+$  at m/z 73) and by the dominance of  $\text{C}_3\text{H}_7^+$  rather than  $\text{C}_2\text{H}_3\text{O}^+$  at m/z 43 which facilitated the distinction of peat emissions over wood or smoky coal emissions.

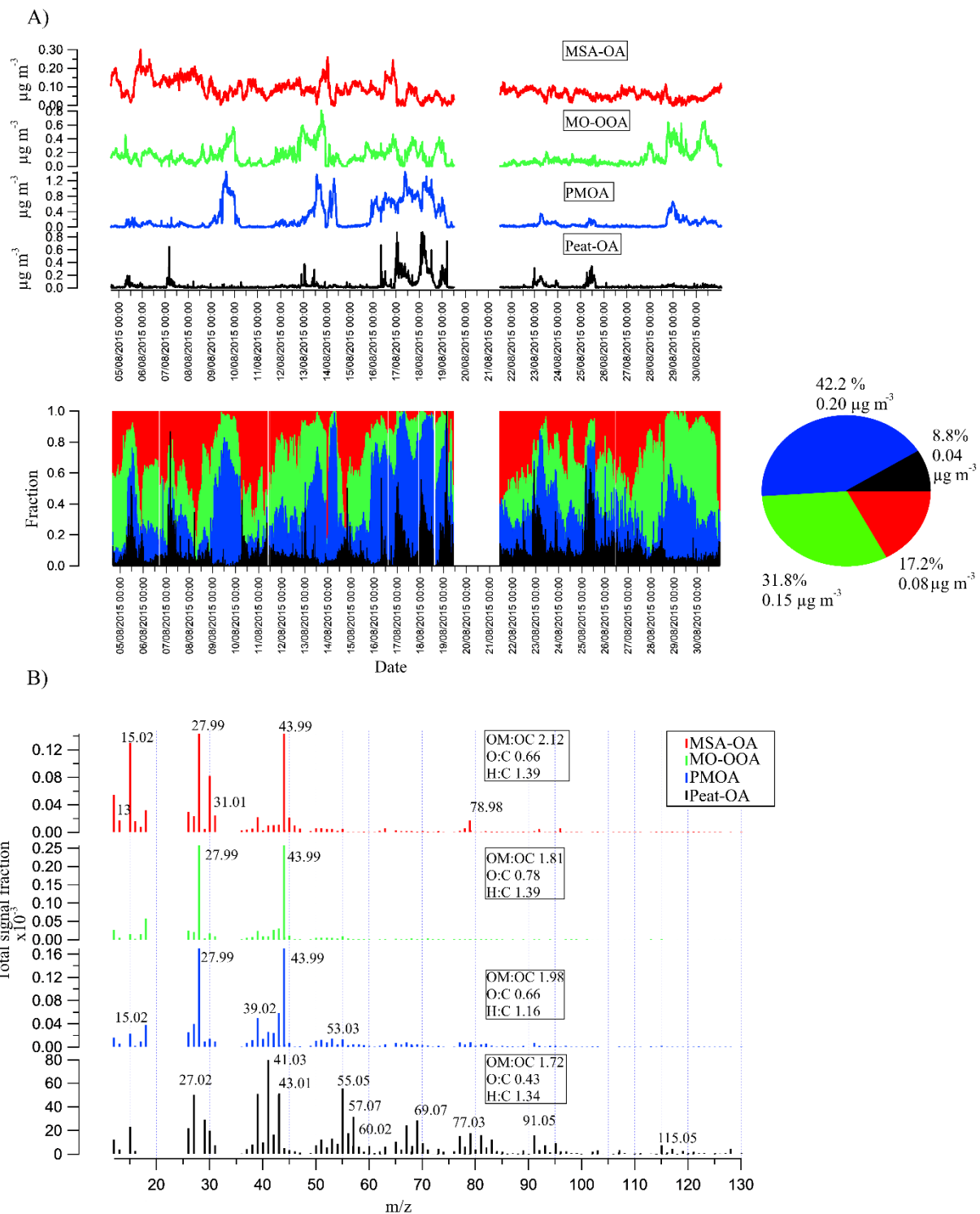


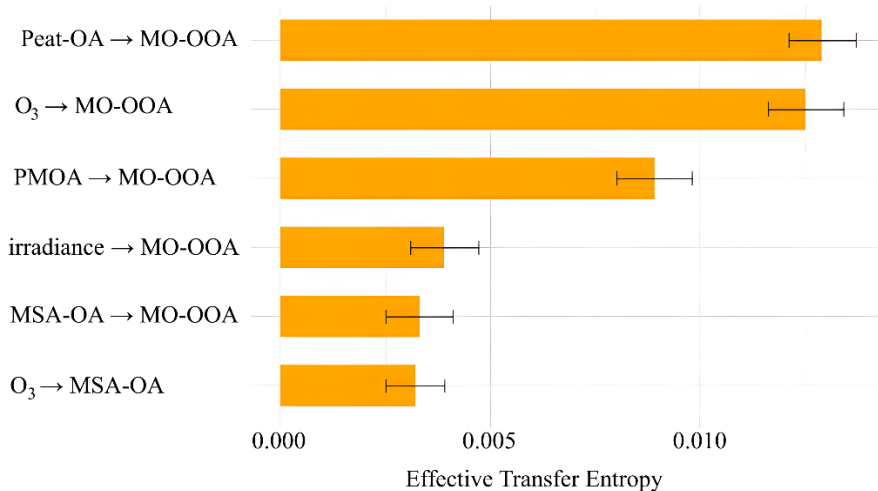
Figure 2. A) Factors time series (MSA-OA in red, MO-OOA in green, PMOA in blue, Peat-OA in black) and associated relative contribution time series and pie chart showing fractions and respective mass concentrations ( $\mu\text{g m}^{-3}$ ) for the whole period B) Factors mass spectra (MSA-OA in red, MO-OOA in green, PMOA in blue, Peat-OA in black) and associated improved ambient OM:OC, O:C and H:C ratios (Canagaratna et al., 2015).

### 3.2.1 More Oxidised Oxygenated Organic Aerosols (MO-OOA)

MO-OOA main contributing ions are associated with oxygenated compounds belonging to the COOH functional group (Figure S5), reflecting pronounced fragmentation of mono- and dicarboxylic acids into fragments with multiple oxygen atoms (Duplissy et al., 2011). Specifically,  $C_xH_yO_z$  ( $z > 1$ ) ions family accounts for 63.1% of the total mass spectra intensity, followed by  $C_xH_yO_z$  ( $z = 1$ ) ions family ( $m/z$  27.99,  $m/z$  43.02,  $m/z$  42.01...) contributing 13.68% to MO-OOA, adding up to a total contribution of 76.8%. Additionally,  $CO^+$  and  $CO_2^+$  each accounts for 25% of MO-OOA intensity which is typical for remote Ocean carboxylic acids (Dominutti et al., 2022).

In contrast,  $C_xH_y$  (aliphatics) ions family ( $m/z$  13.00,  $m/z$  15.02,  $m/z$  16.03...) contributes only 13.7% to MO-OOA total mass spectra intensity. Nitrogen-containing ions fragments constituted a very low portion of the signal (0.8%), similarly to previous remote ocean measurements (Ovadnevaite et al. 2011). The weak contribution from  $C_2H_3O^+$  (3.16%) which has been reported to be predominantly due to non-acid oxygenates (Ng et al., 2011a) suggests a considerable prevalence of aging/oxidation during transport over the North-East Atlantic Ocean. This is also further confirmed by the low  $m/z$  43:44 ratio of 0.12 hinting to MO-OOA rather than less oxidised species (Ng et al., 2011). This factor respective O:C ratio and H:C ratio of 0.78 and 1.17 further agree with MO-OOA reported at other similar locations (Figure S4). MO-OOA also has a strong contribution from  $CO_2^+$  (25.7%) which is assumed to originate mainly from acids or acid-derived compounds (Duplissy et al., 2011; Ng et al., 2011) that are known to be mostly water-soluble (Decesari et al., 2007) such as organic acids (e.g., mesotartaric acid, meso-erythritol, tartaric acid, oxalic acid) formed from oligomerization of small  $\alpha$ -dicarbonyls (e.g., glyoxal) (Cui et al., 2022).

MO-OOA formation through ozonolysis is postulated based on a robust hourly averaged correlation ( $R=0.67$ ) of MO-OOA to  $O_3$  across the entire observational period (Figure S6-B). Using the effective transfer entropy test (Behrendt et al., 2019) further reveals the non-linear dynamics between  $O_3$  and MO-OOA, indicating  $O_3$  as a significant reactant in the formation of MO-OOA from its precursors with a transfer entropy value (Figure 3) of 0.0144 and an effective transfer entropy value of  $0.0127 \pm 0.0009$  ( $p$ -value<0.05). In other words, there is a significant directional information flow between the two time series. Figure 3 also shows that MO-OOA is a mix of local (Peat-OA) and regional marine influence (PMOA, but also MSA-OA to a lesser extent) all eventually concurring to MO-OOA formation, with ozone contributing 3x more information to MO-OOA than irradiance does. Although, irradiance and ozone was measured onsite and may not be as directly related to conditions of aerosol during transport. This Yet, this finding aligns with studies showing  $O_3$  to be a strong oxidation driver during summertime in the marine environment (Ovadnevaite et al., 2011), where unsaturated aliphatic chains ( $C=C$  double bonds) react with ozone to form oxidised compounds (Decesari et al, 2011).



**Figure 3. Significant (p-value=0) effective transfer entropy flow values between PMF factors, ozone and irradiance.**

### 3.2.2 Methanesulphonic Acid Organic Aerosols (MSA-OA)

355 The mass profile of MSA-OA reveals that two oxygenated carbon families CHO (sum of  $C_xH_yO_k$  and  $C_xH_yO_w$  where  $k = 1$  and  $w > 1$ ) dominate 53.4% of the total mass spectra fraction followed by aliphatics (pure Hydrocarbon-like,  $C_xH_y^+$ ) whose fraction accounts for 33.3% (Figure S5). MSA-OA is clearly identified owing to its substantial contribution from the  $C_xS_y^+$  family (6%) over other sources, this is in line with the  $C_xS_y^+$  contribution (7%) for MSA-OA also reported by Huang *et al.* (2018). The excellent correlation ( $R=0.82$ )

360 between this factor and the  $C_xS_y^+$  family (Figure S6-D) also further highlights the organosulphur nature of MSA-OA as opposed to other factors.

Similarly, to results reported by Schmale *et al.* (2013), the correlation coefficient with the AMS database MSA-OA laboratory reference spectrum (Figure S6-A) is rather moderate ( $R=0.55$ ), although this factor spectra still allows for the precise identification of characteristic MSA ions at m/z 44.98 ( $CHS^+$ ), 47.00 ( $CH_3S^+$ ), 64.97 (365  $HSO_2^+$ ), 77.98 ( $CH_2SO_2^+$ ), 77.99 ( $CH_3SO_2^+$ ), and 95.99 ( $CH_4SO_3^+$ ). MSA-OA O:C and H:C ratios were 0.66 and 1.39 respectively, close to values (O:C: 0.54, H:C: 1.42) reported by Loh *et al.* (2022).

370 MSA-OA  $C_xH_y$  family also features a typical  $CH_3^+$  ion at m/z 15.02 that is absent from other factors. Similarly, the  $C_xH_yO_w$  ( $w=1$ ) family features the tracers ions  $CH_2O^+$  (8.2%) and  $CH_3O^+$  (2.4%) which are heat stress related marker (Faiola *et al.* 2015) attributed to methyl jasmonate (MeJA) and possibly acrylic acid (Van Alstyne and Houser, 2003) or other oxylipins stress enzymes (Aguilera *et al.*, 2022; Koteska *et al.*, 2022) which are known to be emitted by kelp (Saha and Fink, 2022) or phytoplankton species (Koteska *et al.*, 2022).

375 The  $C_xS_y^+$  fragment family was dominated by  $CHS^+$ (25.9%),  $CH_3SO_2^+$  (20.2%),  $CH_2S^+$  (12.2%),  $CH_4SO_3^+$  (7.5%),  $CH_3SO^+$  (7.2%),  $CH_2SO_2^+$ , (6.9%),  $CH_4SO_2^+$  (5.45),  $CH_3S^+$  (6.41%),  $C_2H_4SO_2$  (5.4%) and  $CH_2SO^+$  (2.5%) which are common MSA ions found in the literature (Moschos *et al.*, 2022). Overall, the  $C_xH_y^+$  and  $C_xS_y^+$  fragment ions families indicate a clear MSA fragmentation pattern with a characteristic high  $CH_3^+$  relative intensity (13%) typical for marine SOA in line with recent findings (Huang *et al.*, 2018; Moschos *et al.*, 2022).

Finally, MSA-OA correlated moderately (Figure S6-B) with particulate sulphate ( $R=0.51$ ) which is expected since dimethyl sulphide, released by phytoplankton, can be oxidized to either form MSA or sulphur dioxide and then to sulphuric acid, leading to their partitioning into the particulate phase (Mungall *et al.*, 2018). Although MSA is often found in clusters with amine, dimethylamine or trimethylamine (Bork *et al.*, 2014; Chen *et al.*, 2016; Paglione *et al.*, 2024), the MSA spectrum minor methylamine contributions such as m/z 30 ( $\text{CH}_4\text{N}^+$ ), m/z 41 ( $\text{C}_3\text{H}_5^+$ ), m/z 42 ( $\text{C}_2\text{H}_4\text{N}^+$ ) (Malloy *et al.*, 2009) were too sparse to assume involvement.

### 3.2.3 Primary Marine Organic Aerosols (PMOA)

High-resolution mass spectrum of this factor reveals that two CHO oxygenated carbon families (sum of  $\text{C}_x\text{H}_y\text{O}_k$  and  $\text{C}_x\text{H}_y\text{O}_w$  where  $x = 1$  and  $w > 1$ ) dominate 61.5% of the total mass spectra followed by aliphatics (pure Hydrocarbon-like,  $\text{C}_x\text{H}_y$ ) whose fraction accounts for 36.2% of the total mass spectra signal (Figure S5) aligning with previous findings reported by Ovadnevaite *et al.* (2011). The  $\text{C}_x\text{H}_y\text{O}_w$  ( $w=1$ ) family features ions series (m/z 55.02, 69.03, 83.05, etc...) related to alkenyl groups, diunsaturates, cyclic alcohols, and ethers. Such functional groups repartition is consistent with previous reports of water-insoluble organics being formed in sea spray (O'Dowd *et al.* 2004; Ovadnevaite *et al.* 2011). Additionally, this factor mass spectrum closely resembles ( $R=0.99$ ) marine organic aerosols (MOA) mass spectra (Ovadnevaite *et al.* 2011) (Figure S6-A) and its O:C ratio of 0.66 and an H:C ratio of 1.16 respectively are close to literature O:C values for sea spray (Ovadnevaite *et al.* 2011; Flerus *et al.* 2012; Willoughby *et al.* 2016) (Figure S4).

PMOA  $\text{C}_x\text{H}_y$  mass spectra family is dominated by ion series  $\text{C}_x\text{H}_{2y-3}$  (m/z 39.02, 53.03, 67.05 etc...) indicating dienes, alkynes, and cycloalkenes contributions, which is further confirmed by the presence of  $\text{C}_x\text{H}_{2y-1}$  ions series (m/z 27.02, 41.04, 55.05 etc...) while the  $\text{C}_x\text{H}_{2y+1}$  family (m/z 43.05, 57.07, 83.08 etc...  $y>2$ ) indicative for anthropogenically influenced refined hydrocarbons is absent from this factor mass spectra. The marine biogenic origin of this factor is also indicated by the absence of alkanes ( $\text{C}_x\text{H}_{y+2}$ ; m/z 16.03, 58.08, 72.09) which are typical for continental air masses (Lewis *et al.*, 2021) and by its lack of correlation with eBC ( $R=0.17$ ) thereby excluding contribution from fossil hydrocarbons to PMOA. The  $\text{C}_x\text{H}_y$  family is also marked by alkyls ( $\text{C}_x\text{H}_{2y+1}$ ; m/z 15.02, 29.03, 37.00 etc...) which have been reported to be dominant in sea spray during phytoplankton blooms as a possible result of phosphate cycling (Cavalli, 2004; Meador *et al.*, 2017).

Prior atmospheric measurements have shown that PMOA containing a large fraction of alkenes and oxygenated functional groups (ie. alcohols, ethers, aldehydes, ketones) are dominated by insoluble organic colloids and aggregates (Facchini *et al.*, 2008; Rinaldi *et al.*, 2020) composed of microgels derived from phytoplankton extracellular metabolic extraction and adsorption organic pool rather than exopolymers produced from bacteria, with abacterial microgels aerosols being quite common and possibly accounting for 50-90% of phytoplankton derived organics (Bigg and Leck 2008; Bates *et al.* 2012; Liu *et al.* 2023). These bacterial exopolymers would follow the makeup of ordinary bacterial cell fragments, which comprise approximately 55% nitrogen-containing organics and 10% carbohydrates (Schmale *et al.*, 2013). The latter are accounted for by summing up pure carbohydrates (i.e.; glucose, saccharose, mannitol and glycogen) identified by typical fragments (Schmale *et al.*, 2013; Schneider *et al.*, 2011) at m/z 56.03 ( $\text{C}_3\text{H}_4\text{O}^+$ ), m/z 60.02 ( $\text{C}_2\text{H}_4\text{O}^+$ ), m/z 61.03 ( $\text{C}_2\text{H}_5\text{O}^+$ ) and m/z 85.03 ( $\text{C}_4\text{H}_5\text{O}^+$ ) only amounting for about 1.3% of the total PMOA aerosols mass. Similarly, contributions from other bacterial tracers such as glycogen; m/z 55.01 (1.36%), mannitol; m/z 56.02 (0.4%) and polysaccharide species; m/z 97.02 ( $\text{C}_5\text{H}_5\text{O}_2^+$ ) and m/z 125.02 ( $\text{C}_6\text{H}_5\text{O}_3^+$ ) (Glicker *et al.*, 2022) tracer ions were also relatively poor

(0.7%). All of this paired with below detection limit amino acids thus implicates that PMOA organic pool was largely shaped by abacterial processes. However, bacterial influence cannot be ruled out entirely as carbohydrates might have been processed by enzymes or acidity during air masses transport and subsequent aging (Zeppenfeld et al., 2023). [A potential important tracer for this activity could be lactic acid which has been observed before in sea spray owing to microorganisms fermenting sugars \(Miyazaki et al., 2014; Paglione et al., 2024\), although lactic acid itself still remains understudied in HR-ToF-AMS studies.](#)

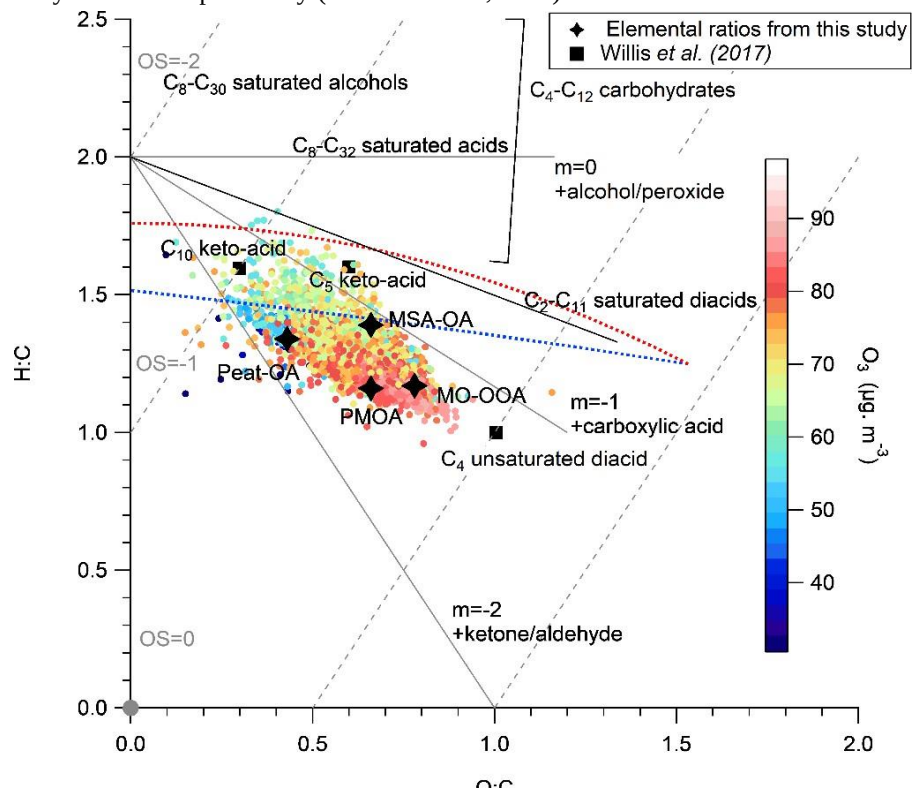
### 420 3.2.4 Peat Related Organic Aerosols (Peat-OA)

425 Although the measurement period is largely dominated by pristine ocean air masses, some residential heating influence is still observed owing to local peat burning. Peat-OA mass spectra is largely dominated by saturated Alkanes- $C_xH_{2y+1}^+$  (76.92%) and Alkenes- $C_xH_{2y-1}^+$  (Figure S5) which is typical for peat. This factor mass spectrum correlates well ( $R=0.86$ ) with previous measurements of Peat-OA in Galway city (Lin et al., 2017) (Figure S6-A). More specifically, the presence of aromatic ion series at m/z 77.03 ( $C_6H_5^+$ ) and m/z 91.0.5 ( $C_7H_7^+$ ) (Cubison et al., 2011) and the ratio between m/z 55.05 ( $C_4H_7^+$ ) and m/z 57.07 ( $C_4H_9^+$ ) of 1.74 as well as the ratio between m/z 43.05 ( $C_3H_7^+$ ) and m/z 44.01 ( $C_2H_3O^+$ ) of 1.03 all allow for the clear distinction of peat 430 burning over other sources (Lin et al., 2017). Peat-OA was freshly emitted as evidence by the pollution wind rose (Figure S7-E, S7- F) and concurrent increase along with eBC ( $R=0.72$ ) indicate that both were locally co-emitted within the PBL.

### 435 3.3 Elemental ratios -Van Krevelen diagram

440 The Van Krevelen (VK) diagram (Heald et al., 2010) provides valuable information on chemical evolution of OA as demonstrated by subsequent marine aerosols studies (Ovadnevaite et al. 2014; Willis et al. 2017; Dada et al. 2022). The VK plot of the PMF factors identified in this study superimposed with bulk OA O:C and H:C values is depicted in Figure 4. Overall, the bulk OA slope of -1.18 and  $\overline{Osc}$  values spanning over -1.8 to 0.8 in the carbon oxidation state space indicates that higher levels of oxidation involving the generation of carboxylic acids, and the subsequent breakdown of the carbon backbone are prevalent over the measurement period which is consistent with MO-OOA functional groups (Heald et al. 2010, Ng et al., 2011). The O:C ratios for MO-OOA, PMOA and MSA-OA all fall within the range of 0.64–1.15 reported for diverse OOA factors from previous studies (Aiken et al., 2008; Jimenez et al., 2009). All PMF factors have H:C values lower than 2 which indicate that they all contain unsaturated carbons capable of reacting with  $O_3$  (Ovadnevaite et al. 2011). This is evidenced 445 by effective transfer entropy flow analysis (Behrendt et al., 2019) between Peat-OA, PMOA, MO-OOA, MSA-OA and  $O_3$  values (Figure 43) which indicates that Peat-OA had the highest information transfer flow, making it the most susceptible to ozonolysis, closely followed by PMOA and, to a lesser extent, MSA-OA. Both Peat-OA (O:C=0.43, H:C=1.34) and PMOA (O:C=0.66, H:C=1.16) VK positions broadly fall in the area consistent with lignin-like compounds ( $H/C = 0.6–1.5$ ,  $O/C = 0.1–0.6$ ; Park et al. 2022) which have been largely associated with 450 terrestrial origin OA (Jang et al., 2022) and found to be high in Arctic Ocean air masses as well (Choi et al., 2019) with authors reporting ~30% of the total assigned molecular formulae as marine lignin-like compounds. These lignin-like compounds are also known to oxidise and form Humic-like molecules, characterized by polar carbonyl (keto and carboxyl) functional groups alongside hydrophobic aliphatic chains (Cavalli, 2004) which broadly agrees with MO-OOA functional groups.

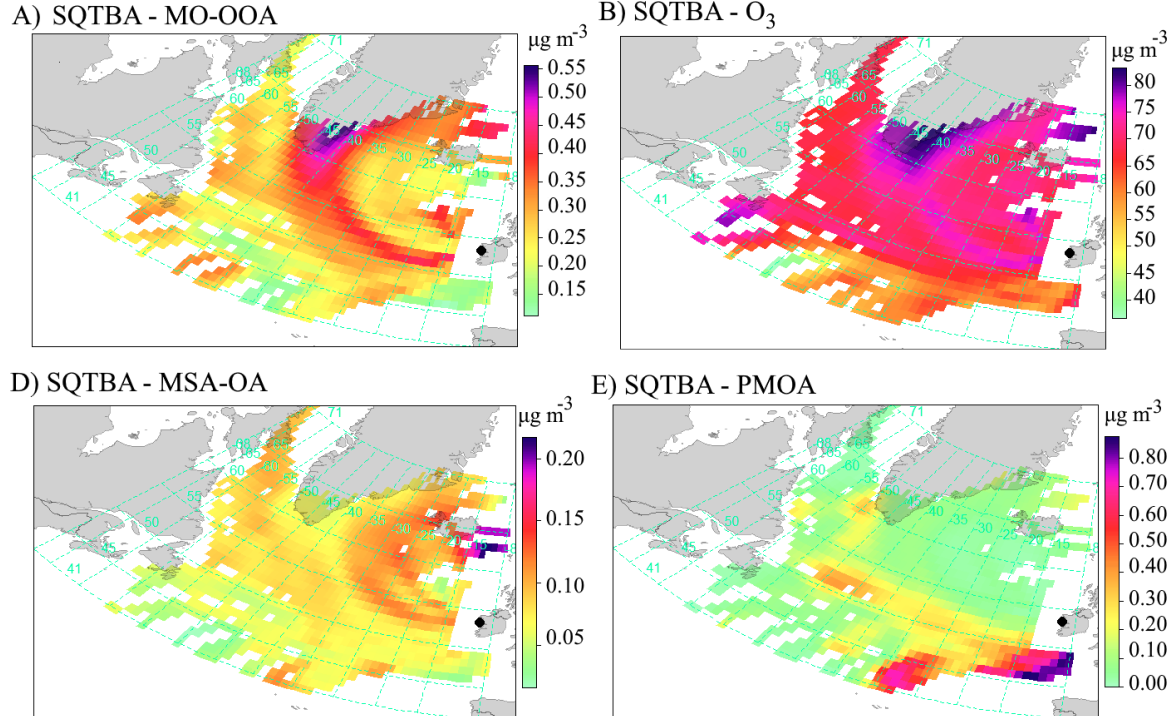
455 MSA-OA ( $O:C=0.66$ ,  $H:C=1.9$ ) is then also examined by colouring the VK scatter plot (Figure S8) with the MSA-OA/ $SO_4$  ratio, a proxy for biological marine sources contributions from DMS (Chen et al., 2021) with values ranging from 0.001 (ubiquitous anthropogenic influences) to 0.354 (significant contribution from biological marine sources) with an average value of 0.102 in line with pristine conditions (Huang et al., 2018). Figure S8 shows that high MSA-to-sulphate ratio were consistent with VK region for  $C_2$ - $C_{12}$  saturated diacids and inconsistent with  $C_4$ - $C_{12}$  carbohydrates and derivatives (trehalose, erythritol, arabitol, mannitol, sucrose, galactose, glucose, fructose etc...) similarly to results reported for summertime Arctic (Willis et al., 2017).  
 460 However, as opposed to Arctic aerosols,  $H:C$  ratios being higher, we report no association with VK areas for  $C_4$  unsaturated diacids (e.g maleic and fumaric acid) nor with  $C_{10}$  and  $C_5$  keto-acids (levulinic and pinonic acid) which are aqueous photochemistry tracers from isoprene and  $\alpha$ -pinene oxidation (Kołodziejczyk et al., 2019; Rapf et al., 2017). This is in line with the absence of other isoprene tracers;  $C_4H_5^+$  (0.49%) at  $m/z$  53.03 and  $C_5H_6O^+$  (0.18%) at  $m/z$  82.04 (Hu et al., 2015; Robinson et al., 2011) and monoterpenes tracers (Boyd et al., 465 2015), namely  $C_5H_7^+$  at  $m/z$  67.05 (0.15%) and ( $C_7H_7^+$ ) at 91.05 (0.15%). This is also supported with-by the lack of covariance ( $Cov[X, Y] \approx 0$ ) between bulk  $CO_2^+$ ,  $CO^+$  and  $C_2H_3O^+$  time series which also denotes the absence of non-acid carbonyls (naCO) (Yazdani et al., 2022) which are known to be derived from isoprene and 470 monoterpenes (Russell et al. 2011). The reasons behind the absence of isoprene and monoterpenes influence on OA in these findings are currently unclear although processes such as surface ocean consumption or unexplored oxidations pathways could be a possibility (Benavent et al., 2022).



475 **Figure 4. Relationship between the ToF-AMS estimated hydrogen-to-carbon (H/C) and oxygen-to-carbon (O/C) ratios of organic species [Canagaratna et al., 2015] coloured by  $O_3$  mixing ratio, all observations above ToF-AMS detection limits are shown for the entire period. Grey lines represent the ambient range of O/C and H/C observed by Ng et al. [2011] while dashed line represent the average carbon oxidation state ( $OS_c \approx 2 \times O : C - H : C$ ) (2011) superimposed on the Van Krevelen diagram (Ng et al. 2011, Kroll et al., 2011). Elemental composition of  $C_8$ - $C_{30}$  saturated alcohols,  $C_8$ - $C_{32}$  saturated acids,  $C_2$ - $C_{11}$  saturated diacids,  $C_4$**

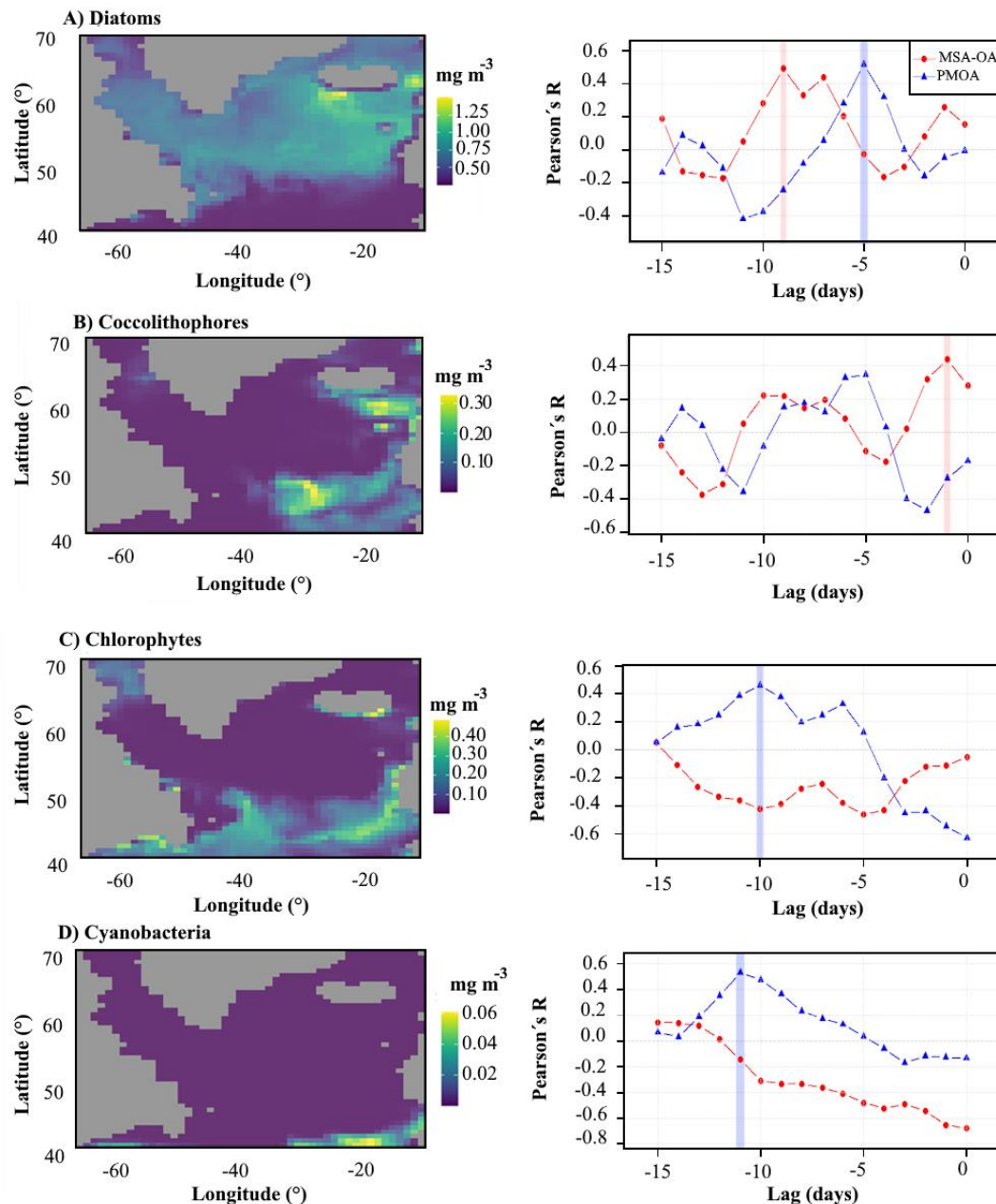
unsaturated diacid (maleic and fumaric acid), C<sub>4</sub>–C<sub>12</sub> carbohydrates (e.g., trehalose, erythritol, arabitol, mannitol, sucrose, galactose, glucose, and fructose), and C<sub>5</sub> and C<sub>10</sub> ketoacids (levulinic and pinonic acid, respectively) are shown for reference (Willis *et al.*, 2017).

### 3.4 Air masses and source apportionment



485 **Figure 5. Simplified Quantitative Transport Bias Analysis (SQTBA) -Gaussian Air masses dispersion for PMF sources (PMOA, MO-OOA, MSA-OA) and O<sub>3</sub>.**

MO-OOA (Figure 5-A) strongest sources can be traced back northward along a cyclonic gradual crescent shape spreading from Greenland Seas South of Cape Farewell (See Figure S9 for Ocean areas toponymy). This 490 culminates further with air masses origins spanning over the East Greenland Current (Denmark Strait) upwards to the Iceland Sea, [south-south](#) of Jan Mayen. MO-OOA is otherwise ubiquitous and shows contributions over the Newfoundland, Labrador and Iceland basins as well as other areas. O<sub>3</sub> (Figure 5-B) shared similar origin as MO-OOA further confirming its role in MO-OOA formation. Overall, we observe aged polar air masses eventually flowing from Greenland to MHD. The sustained blockade and aging of air masses over Greenland is 495 known and attributed to summertime high-pressure systems surrounding this region influenced by Arctic amplification (Pettersen *et al.*, 2022; Preece *et al.*, 2023) where [the Irminger current](#) also acts as a hotspot for turbulent eddies and heat transport which might contribute to aerosol nucleation (Semper *et al.*, 2022). Here the presence of a blocking anticyclone transition (Figure S10) leading to reduced cloud cover and warm air advection might ultimately have contributed to an increase in aged SOA at the southern tip of 500 Greenland possibly owing to its orography.



505 **Figure 6. Time averaged maps (0.67 x 1.25 deg) over 2015-Aug, Region 59W, 37N, 34E, 82N of dominant**  
**phytoplankton groups from NOBM Model data (Rousseaux *et al.* 2017; Buchard *et al.* 2017) and**  
**corresponding lagged crossed correlations for MSA-OA (red) and PMOA (blue) against A) Diatoms B)**  
**Coccolithophores, C) Chlorophytes and D) Cyanobacteria. The blue and red shaded areas correspond to**  
**maximum significant crossed-correlations extracted from the autocorrelation function (ACF)- 95%**  
**criteria.**

510 MSA-OA (Figure 5-C) main sources include the Iceland basin and more specifically the Iceland-Faroe Ridge. This is consistent with literature highlighting the diversity of eukaryotic phytoplankton in the Icelandic marine environment with the haptophyte coccolithophore *Emiliania huxleyi* being dominant during summertime (Cerfonteyn *et al.*, 2023) owing to nutrients transport by the North Atlantic Current acceleration (Oziel *et al.*, 2020) and findings (O'Dowd *et al.* 2015; Mansour *et al.* 2023) indicating concomitant MSA concentrations uptick during summertime. MSA-OA also spans along the East Greenland Current (Denmark Strait) where wind-driven coastal upwelling (Håvik and Våge, 2018) might result in increased DMS emissions (Edtbauer *et*

al., 2020). Likewise, MSA-OA extend moderately over diverse regions such as the North-Western European Basin, the Newfoundland basin (where intense DMS fluxes have been reported; Bell *et al.* 2021) and the Labrador Sea.

520 PMOA (Figure 5-D) on the other hand strongly extends over the South of the Celtic Sea and West of the Bay of Biscay as well as West European basin waters and are otherwise diffused all over the North Atlantic Ocean with moderate intensity hotspots over the Newfoundland basin (Davis strait) possibly owing to an inflated subpolar gyre (Hátún *et al.*, 2016).

525 Examination of NOBM model data (Figure 6) further reveals distinct MSA-OA and PMOA patterns. MSA-OA overlap with coccolithophores dominated ecoregions as well as diatoms ones. Similarly, diatoms also seem to contribute to PMOA sources, which is in line with recent results hypothesising that diatoms have a greater atmospheric significance than other eukaryotes due to their observed enrichment in PMOA (Alsante *et al.*, 2021) whereas association with coccolithophores appears much weaker than for MSA-OA. Another distinction lies in PMOA overlapping with chlorophytes (*flagellates, Phaeocystis spp*) over the Western European basin. This geographic area hosts more than 512 chlorophyte species (Narayanaswamy *et al.*, 2010) with recent reports of 530 chlorophytes being one of the key contributors to marine productivity (Landwehr *et al.*, 2021), further research is warranted to fully understand their role along other phytoplankton in this region during summertime. Likewise, cyanobacteria (combination of *Synechococcus*, *Prochlorococcus*, and nitrogen fixers such as *Trichodesmium*) might also contribute to PMOA more sparsely, especially at lower latitudes in the North Atlantic Ocean as previously reported (Baer *et al.*, 2017).

535 Here, calculated lagged correlations (Figure 6) further pointed at MSA-OA being directly associated with coccolithophores (with a lag of -1 day) as well as diatoms (lag of -9 days), however no significant correlations were observed for either cyanobacteria or chlorophytes. As opposed to MSA-OA, the association between coccolithophores and PMOA doesn't appear as meaningful (their autocorrelations are not statistically significantly different from zero). PMOA on the other hand are also associated with diatoms (lag of -5 days) and 540 show unique associations with chlorophytes (lag of -10 days) as well as cyanobacteria (lag of -11 days).

Overall, association between OA enriched sea spray time series and phytoplankton groups remains controversial owing to a wide range of governing mechanisms as highlighted by previous studies using chl- $\alpha$  as a proxy to calculate cross correlation time lags over the North Atlantic which were found to vary between 8 days (Rinaldi *et al.*, 2013) and 24 days (O'Dowd *et al.*, 2015) depending on the period and length of measurements.

545 Late summer measurements (Mansour *et al.*, 2020) show partially comparable lags to this study with a reported oceanic biological activity affecting aerosol properties within the order of 10-20 days. This delay roughly spans over the full blooming to decaying phase transitions of an algal bloom (Lehahn *et al.*, 2014) and is linked to the release of SSA-transferable organic matter in surface seawater by the interaction with marine viruses causing the demise of phytoplankton blooms (O'Dowd *et al.*, 2015).

550 Here, by focusing on the lagged correlations between PMF factors and specific phytoplankton groups rather than bulk-OA and chl- $\alpha$ , this study's findings indicate that PMOA is formed on such timescale from cyanobacteria and chlorophytes (lags of -11 and -10 days respectively) owing to atmospheric transport from the Western European basin whereas overwhelming diatoms influence results in a much shorter lag of -5 days. Additionally,

MSA-OA is rapidly produced from coccolithophores blooms in 1-2 days. This reflects stressed, senescent, 555 grazed, or virus-infected phytoplankton releasing high quantities of DMSP which rapidly oxidises to form MSA-OA (Mansour et al., 2020).

Finally, the interpretation of diatoms' role on either MSA-OA or PMOA remains ambiguous. The as the -5 days lag with PMOA could hint at lipase activity with -concurring to self-aggregation and formation of free fatty acids during the bloom phase, -during bloom potentially followed by a post-bloom phase (at a lag of -9 days with 560 MSA-OA) with significantly different taxa. Alternatively, the -9 day lag with MSO-OA could have been or caused by simply air advection from remote eco-regions further closer to the near the Arctic which that have been reported to host rich MSA producing diatoms communities as opposed to more not present in the southerly latitudes (Becagli et al., 2016).

#### 4. Conclusions

565 This study leverages high-resolution online aerosol mass spectrometry source apportionment to investigate the chemical composition and sources of submicron organic aerosols representing marine environment during a summertime period marked by phytoplankton blooms. The results emphasise balanced mass contributions from POA (PMOA and Peat-OA) and SOA (MO-OOA and MSA-OA), each category accounting for approximately 50% of the total submicron organic aerosol mass, with distinct chemical compositions reflective of their varied 570 origins.

One of this study's key finding is that summertime polar air masses undergo significant ozonolysis over the 575 remote ocean which happens to be largely driven by the ageing of Greenland blocking blocked air masses aging and anticyclonic conditions. Transfer entropy is introduced here to explain the dynamics of ozonolysis in this context revealing significant information transfer to MO-OOA during unsaturated aliphatic chains (C=C double bonds) breakdown of PMOA as well as MSA-OA to a lesser extent. However, this transfer entropy approach additionally shows that MO-OOA is also being formed locally from Peat-OA oxidation, as such, further studies will aim at exactly delineating open ocean versus locally produced MO-OOA.

Another essential takeaway is that OA not only reflects atmospheric chemistry and meteorology but may also 580 serve as an indicator of marine ecosystems (i.e. MSA-OA enzymes stress makers and PMOA phytoplankton extracellular metabolic processes markers). Air masses trajectory analysis also show reveal the source aerosols-phytoplankton ecoregions contributions with MSA-OA contributions are traced to the Iceland Basin and the Iceland-Faroe Ridge, with a rapid production burst (lag of 1-2 days) following coccolithophore blooms.

Whereas relationship with diatoms show much longer lag (9 days) indicating fundamentally different oceanic 585 biological processes. In contrast, PMOA is sourced from more diverse ecoregions (Southern Celtic Sea, West European Basin, and Newfoundland Basin), with additional chlorophytes and cyanobacteria influences from more southerly latitudes. All of this suggests that different phytoplankton taxa contributions to OA lead to specific m/z tracers and functional groups repartition (i.e. sulphides as coccolithophores tracers, aliphatics as tracers for diatoms) though further investigation is needed to explore the biological processes and ecoregions specificities influencing this relationship. Overall, this study demonstrates the complex aerosol chemistry and 590 diverse geographic origins influencing POA and SOA formation in the Northeast Atlantic marine environment.

Our findings emphasise the need for further long-term investigation to fully account for the various precursors and pathways contributing to OA, given their significant impacts on aerosol-climate interactions.

*Competing interests.* The authors declare that they have no conflict of interest.

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600 *Data availability.* Data available upon request. [The NASA Ocean Biogeochemical Model output data was obtained with the Giovanni online data system, developed and maintained by the NASA GES DISC. The ERA5 boundary layer data are available at the Copernicus Climate Data Store portal \(Hersbach et al., 2020\).](#)

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