A one-year ACSM source analysis of organic aerosol particle 2 contributions from anthropogenic sources after long-range transport 3 at the TROPOS research station Melpitz

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14 Abstract

15 Atmospheric aerosol particles are a complex combination of primary emitted sources (biogenic and anthropogenic) and secondary aerosol resulting from the aging processes such as condensation, coagulation, and cloud processing. To better 16 17 understand their sources, investigations have been focused on urban areas in the past, while rural-background stations are 18 normally less impacted by surrounding anthropogenic sources. Therefore, they are predisposed for studying the impact of long-19 range transport of anthropogenic aerosols. Here, the chemical composition and organic aerosol sources of submicron aerosol 20 particles measured by an aerosol chemical speciation monitor (ACSM) and a multi-angle absorption photometer (MAAP) were investigated at Melpitz from September 2016 to August 2017. The location of the station at the frontier between Western and 21 22 Eastern Europe makes it the ideal place to investigate the impact of long-range transport over Europe. Indeed, the station is 23 under the influence of less polluted air masses from westerly directions and more polluted continental air masses from Eastern 24 EuropeTo better understand their sources, investigations have been focused on source identification in urban areas in the past. 25 while rural background stations are normally less impacted by surrounding anthropogenic sources. Therefore, they are 26 predisposed for studying the impact of long range transport of anthropogenic aerosols. Moreover, long term measurements 27 ean help to study the potential temporal changes in the sources. Here, the chemical composition and organic acrosol sources 28 of submicron aerosol particles were investigated at the Central European rural background research station, Melpitz, using a one yearlong dataset determined by an aerosol chemical speciation monitor (ACSM) and a multi-angle absorption photometer 29 30 (MAAP) from September 2016 to August 2017. Melpitz represents due to its location the Central European aerosol. It is an 31 ideal location to investigate the impact of long-range transport, since the location is influenced by less polluted air masses from 32 westerly directions and more polluted continental air masses from Eastern Europe. The organic aerosol (OA) dominated the 33 submicron particle mass concentration and showed strong seasonal variability ranging from 39 % (in winter) to 58 % (in

summer). It was followed by sulphate (15 % and 20 %) and nitrate (24 % and 11 %). The OA source identification was 34 35 performed using the rolling positive matrix factorisation (PMF) approach to account for the potential temporal changes in the 36 source profile (SoFi Pro). It was possible to split OA into five -factors with a distinct temporal variability and mass spectral 37 signature. Three were associated to-with anthropogenic primary OA (POA) sources: hydrocarbon-like OA (HOA, 5.2 % of 38 OA mass in winter and 6.8 % in summer), biomass burning OA (BBOA, 10.6 %, and 6.1 %) and coal combustion OA (CCOA, 39 23 %, and 8.7 %). Another two are secondary/processed oxygenated OA (OOA) sources: less-oxidized OOA (LO-OOA, 28.4 40 %, and 36.7 %) and more-oxidized OOA (MO-OOA, 32.8 %, and 41.8 %). Since equivalent black carbon (eBCeBC) was 41 clearly associated with the identified POA factors (sum of HOA, BBOA, and CCOA, $R^2=0.87$), eBCeBC's contribution to 42 each of the POA factors was achieved using a multi-linear regression model. Consequently, CCOA represented the main 43 anthropogenic sources of carbonaceous aerosol (sum of OA and eBCeBC) not only during winter (56 % of POA in winter) but 44 also in summer (13 % of POA in summer), followed by BBOA (29 % and 69 % of POA in winter and summer, respectively) 45 and HOA (15 % and 18 % of POA in winter and summer, respectively). A seasonal air mass cluster analysis was used to understand the geographical origins of the different aerosol types and showed that during both winter and summer time, PM_1 46 47 (PM with an aerodynamic diameter smaller than 1µm) air masses with eastern influence was always associated with the highest 48 mass concentration and the highest coal combustion fraction. Since during winter time, CCOA is a combination of domestic 49 heating and power plants emissions, the summer contribution of CCOA emphasizes emphasizes the critical importance of coal 50 power plants emissions to rural-background aerosols and its impact on air quality, through long-range transportation.

51 1 Introduction

52 Human health effects of air pollution from particulate matter (PM) are well known, and efforts are being made across the world 53 (WHO, Expert Consultation, 2019) to minimize both long-term exposures to harmful levels and air pollution peaks. 54 Throughout all the PMs, the The submicronic particles known as PM_1 (particles with an aerodynamic diameter less than 1 55 µm), not only have a negative impact on human health (Pop and Dockery, 2016; Daellenbach et al., 2020) but also have a 56 significant effect on visibility (Shi et al., 2014) and climate (Shrivastava et al., 2017). It is ability to penetrate to respiratory 57 system make it more dangerous, therefore more relevant to mitigate adverse health impact-submicronic particles known as 58 PM_{+} (particles with an aerodynamic diameter less than 1 μ m), not only have a negative impact on human health (Daellenbach 59 et al., 2020) but also have a significant effect on visibility (Shi et al., 2014) and climate (Shrivastava et al., 2017). Since the 60 most numerous component of the atmospheric PM is the organic aerosol (OA) (Jimenez et al., 2009; Chen et al., 2022), 61 contributions to OA and explanations of its chemical and physical characteristics remain challenging, whereas the large variety 62 of OA can be attributed to primary emissions by various sources in different seasons, as well as different reactions to 63 atmospheric dynamics and complicated chemical mechanisms depending on meteorological parameters and geographical 64 locations.

In order to evaluate and recognize the sources of OA emission, aerosol mass spectrometers (AMS, Jayne et al., 2000) and 65 66 aerosol chemical speciation monitors (ACSM) (Ng et al., 2011; Fröhlich et al., 2013) are widely deployed worldwide (Chen 67 et al., 2022; Bressi et al., 2021; Fröhlich et al., 2015). AMS is commonly limited to short time periods due to the high 68 maintenance of the AMS measurements and their high operating costs. As a result, only a few studies run AMS continuously 69 (e.g., see Kumar et al., 2022 and O'Dowd et al., 2014). However, there was still a strong need for such a long-term analysis. 70 ACSM is designated for long-term monitoring purposes due to its robustness and much less labour-intense compared to AMS. 71 Therefore, the deployment of ACSM allows us to look at the long-term (more than one year) temporal changes and/or seasonal 72 variability of OA sources.

73 Regarding the identification of OA sources, source apportionment analysis using positive matrix factorisation algorithm (PMF, 74 Paatero & and Tappert, 1994) was has been intensively used over the past two decades on both AMS and ACSM measurements 75 (e.g. see Crippa et al., 2014; Poulain et al., 2020). However, this algorithm faced two main limitations when used during a long 76 time period: firstly, the factor profiles are static over the analyzing period (Paatero, 1997); and secondly, rotational ambiguity 77 which provides non-unique solutions. To solve these issues, a multilinear engine (ME-2, Paatero, 1999) has been implemented 78 in the PMF analysis, which allows to use of a priori knowledge to constrain the model to environmentally reasonable solutions 79 (e.g., Lanz et al., 2008; Canonaco et al., 2013 and; Crippa et al., 2014). To consider the temporal variation of the factor profiles, a rolling approach was suggested (Parworth et al., 2015; Canonaco et al., 2020). The rolling strategy involves advancing a 80 81 smaller PMF window (i.e., 14 days) and moving/rolling it over the whole dataset to catch the temporal changes of the source 82 profiles with a 1-day step.

83 Although several studies in Europe have already conducted source apportionment analyses of one year or more, most of them 84 were associated with urban or suburban environments (e.g., for urban studies: Stavroulas et al., 2019; Vlachou et al., 2019; 85 Huang et al., 2019; Qi et al., 2020; and for suburban studies: Katsanos et al., 2019; Y-Zhang et al., 2019), and only a few of 86 them were studied in rural-background sites (Schlag et al., 2016; Crippa et al., 2014; Vlachou et al., 2018; Paglione et al., 87 2020; Dudoitis et al., 2016; Heikkinen et al., 2020; Chen et al., 2021; and Chen et al., 2022), although the rural-background 88 sites represent the major advantage to be able to study the impact of long-range transport of anthropogenic emissions and their 89 changes over a long time period. The Leibniz Institute for Tropospheric Research (TROPOS) Central European observatory 90 Melpitz has been continuously measuring aerosol chemical compositions for 30 years. The station is a unique place in Europe, 91 sitting at the border between marine-influenced Western Europe and continental Eastern Europe. A direct consequence is that 92 the aerosol chemical compositions and mass concentrations strongly depend on the air mass origins, showing less polluted air 93 masses coming from the West and more polluted air masses from the East (Birmili et al., 2010). 94 However, only a few studies were done on the source identification of the aerosol reaching the station by covering short time 95 periods mostly during winter (van Pinxteren et al., 2016, 2020, 20202).

96 The current study comprehensively investigates the PM₁ aerosol particle chemical compositions and the various OA sources

- 97 for Melpitz as a rural-background station, based on ACSM and multi-angle absorption photometer (MAAP) measurements
- 98 from September 2016 to August 2017, using the most advanced rolling PMF with ME-2 implemented in the SoFi Pro package

99 (Datalystica Ltd., Villigen, Switzerland) (Parworth et al., 2015; Canonaco et al., 2013; Canonaco et al., 2020). Although 100 previous papers already considered this dataset, they were focused on quality assurance (Poulain et al., 2020) to depict the 101 European aerosol chemical composition (Bressi et al., 2021 and Chen et al., 2022) or the relationship between the CCN 102 properties (Wang et al., 2022, Schmale et al., 2017), none of these papers were focused on carbonaceous source identification 103 (OA and eBC) nor discussed the strong dependency of the aerosol chemical composition to the air mass origin. Therefore, a 104 multi-linear regression model was used to estimate the contribution of equivalent black carbon (eBC) to the various primary 105 organic PMF factors such as hydrocarbon-like organic aerosol, biomass burning organic aerosol, and coal combustion organic 106 aerosol. Meanwhile, to better understand the emission area of PM₁ chemical composition and PMF factors, the influence of 107 air mass origin was investigated based on self-developed back-trajectory cluster methods (BCLM). The current study 108 comprehensively investigates the PM1 aerosol particle chemical compositions and the various OA sources for Melpitz 109 based on ACSM and multi-angle absorption photometer (MAAP) measurements from September 2016 to August 2017, 110 using the most advanced rolling PMF with ME-2 implemented in the SoFi Pro package (Datalystica Ltd., Villigen, 111 Switzerland) (Parworth et al., 2015; Canonaco et al., 2013; Canonaco et al., 2020). Moreover, a multi-linear regression 112 model was used to estimate the contribution of equivalent black carbon (eBC) to the various PMF factors. Meanwhile,

113 the influence of air mass origin was investigated to identify the emission area of the different PM1 sources.

114 2 Methodology

115 2.1 Sampling site

116 The atmospheric aerosol measurements were carried out at the TROPOS research station Melpitz (51.54° N, 12.93° E, 117 86 m a.s.l.), located approximately 50 km northeast of Leipzig, Germany. The station itself is mainly encircled by agronomical 118 pastures and forests within a rural area, which that is why the station is recognized as a rural-background station (Spindler et 119 al., 2013). Since 1992, the station has been monitoring the influence of atmospheric long-range transport on the background 120 air quality of Central European (e.g. Spindler et al., 2012 2013). The Melpitz station is part of EMEP (European Monitoring 121 and Evaluation Programme; Level 3 station, Aas et al., 2012), ACTRIS (Aerosol, Clouds and Trace gases Research 122 Infrastructure), GAW (Global Atmosphere Watch of the World Meteorological Organization), and GUAN (German Ultrafine 123 Aerosol Network, Birmili et al., 2009, 2015, 2016). For a general description of the chemical and physical aerosol 124 characterization analysis techniques, check e.g. Spindler et al., (2004, 2010, 2012, 2013); and Poulain et al., (2011, 2014, 125 2020).

126 2.2 ACSM

127 The chemical compositions and mass loadings of non-refractory PM_1 (NR-PM₁: organic, sulphate, nitrate, ammonium, and 128 chloride) with a 30-minute time resolution were measured by an Aerodyne quadrupole ACSM. The ACSM sampling technique 129 and operational information were previously detailed by Ng et al., (2011). 130 Briefly, after PM₁ transmits across a 100 µm critical orifice, the aerosols are centralized into a slender-narrow beam in an 131 aerodynamic lens (Liu et al., 2007). Non-refractory particulate material that evaporates at the oven temperature (generally 132 600°C) is recorded and chemically determined using electron impact quadrupole mass spectrometry at 70 eV (Ng et al., 2011). 133 The ions are then detected using a quadrupole residual gas analyser (RGA, Pfeiffer Vacuum Prisma Plus). The ACSM takes 134 30-second samples of both ambient and particle-free air. The difference in these measurements identifies the aerosol mass 135 spectrum. To change the signal spectra into organic or inorganic species concentrations, the fragmentation table (Allan et al., 136 2004), the ion transmission correction, and the Response Factor (RF) are applied. To improve the particle loss as a result of 137 bouncing off the vaporizer, the ACSM data were processed according to manufacturer guidelines with using a composition 138 dependent collection efficiency (CDCE) correction relying on the algorithms suggested by Middlebrook et al. (2012). 139 Calibrations of Ionization Efficiency (IE) and Relative Ion Efficiency (RIE) were performed using a 350 nm monodispersed 140 ammonium nitrate and ammonium sulphate (Ng et al., 2011). The final mean value for IE was $4.93(\pm 1.45) \times 10^{-11}$ and the 141 mean values for RIEs for ammonium and sulphate respectively were 6.48 \pm 1.26, and 0.68 \pm 0.13. The quality assurance of 142 the ACSM measurements was performed by comparing them with collocated measurements including MPSS, and high-volume 143 filter samples (PM_1 and $PM_{2.5}$) for the total particle mass concentration, water-soluble ions (nitrate, sulphate, and ammonium), 144 as well as OC/EC. Details on the OA/OC and instrumental uncertainties can be found in Poulain et al., (2020). Details on the 145 QA/QC for this dataset can be found in Poulain, et al., (2020). 146 The ACSM ammonium mass concentration mainly corresponds to ammonium nitrate and ammonium sulphate salts. Previously 147 by Poulain et al. (2020), the neutralization state of the particles was estimated for datasets assuming complete neutralization 148 by nitrate, sulphate, and chloride. Therefore, the particles are neutralized when considering nitrate, ammonium, and sulphate 149 in this study. Furthermore, the significant role of organo-nitrate and organo-sulphate on signals of nitrate and sulphate is not 150 negligible (Kiendler- Scharr et al. (2016). Since the O-ACSM is working at a unit mass resolution (UMR), it is not possible to

- 151 distinguish nitrate and sulphate from organic. Therefore, estimating the organo-nitrate would only introduce uncertainties to
- 152 measurements, therefore, we did not consider to conduct this analysis in this study.

153 2.3 Additional measurements

- 154 In parallel to the ACSM, a MAAP was used to measure the mass concentrations of equivalent black carbon (eBC) (model
- 155 5012 Thermo Scientific; Petzold & and Schönlinner, 2004). Conversion of the The eBC mass concentration from the PM₁₀
- 156 data was multiplied by a constant factor of 0.9 following Poulain et al (2011) to estimate the eBC mass concentration in the
- 157 PM₁ fraction. Consequently, all the eBC mass concentrations reported and discussed here correspond to the eBC in the PM₁
- 158 fraction and are referred to as eBC-PM₁. eBC mass concentration from the PM₁₀ inlet to the ACSM PM₁-cut off was made by
- 159 applying a correction factor of 0.9 following Poulain et al (2011). Furthermore, a dual mobility particle size spectrometer
- 160 (TROPOS-type T-MPSS; Birmili et al., 1999) was used to measure the particle number size distribution (PNSD) from 3 to
- 161 800 nm (mobility diameter, d mob) at ambient and 300°C temperatures (Wehner et al., 2002). The MAAP was situated in the

- 162 same laboratory container as the ACSM and these instruments sampled the same PM_{10} inlet after a dryer, and the sampled air
- 163 distribution among the instruments was equally assured by an isokinetic splitter (Poulain et al., 2020).
- 164 In addition to the online measurements, high-volume samplers (DIGITEL DHA-80, Digitel Elektronik AG, Hegnau,
- 165 Switzerland) were utilized to capture daily PM_{2.5} samples on a quartz filter (for 24 hours from midnight to midnight). For more
- 166 details on the sample preparation and evaluation methods, see Spindler et al., (2013). Levoglucosan as a tracer for wood
- 167 burning combustion was measured following linuma et al., (2009) using high performance anion exchange chromatography
- 168 coupled with an electrochemical detector (HPAEC-PAD) Levoglucosan as a tracer for wood burning combustion was measured
- 169 following Jinuma et al., (2009) using high performance anion exchange chromatography coupled with an electrochemical
- 170 detector (HPAEC PAD). that was used for the analysis of anhydro-monosaccharides (linuma et al., 2009).
- 171 Trace gas measurements were also carried out. Ozone was determined by a U.V. Photometric gas analyser mode 49C (Thermo
- 172 Scientific, UK), SO₂ by an APSA-360A (Horiba, Kyoto, Japan), and NO and NO₂ using a customized Trace Level NOx
- 173 Analysis Model 42i-TL (Thermo Scientific) equipped with a blue light converter. Standard meteorological parameters
- 174 (temperature, relative humidity, solar radiation, precipitation, wind direction, and wind speed) were regularly measured.

175 2.4 Rolling PMF (ME-2) source apportionment of OA

176 This work conducted the most advanced source apportionment analysis following a standardized protocol developed by Chen 177 et al., (2022). The PMF method was used to allocate the source of the OA (Paatero & and Tappert, 1994) through the Source 178 Finder professional (SoFi Pro, version 8.0.3.1, Canonaco et al., 2021) software package (Datalystica Ltd., Villigen, 179 Switzerland), within the Igor Pro software environment (Igor Pro, version 8.04, Wavemetrics, Inc., Lake Oswego, OR, USA). 180 Two matrices of factor profiles F and factor contributions G, defined the dataset X, and the matrix E named the residual matrix 181 is the fraction which cannot be described by the model. Time series and the chemical fingerprint of sources respectively have 182 been represented by F_{kj} and G_{ik} , respectively. The dimension of F_{kj} and G_{ik} are based on the order p, which is the number of 183 factors selected to represent the data which is defined by the user:

184

185
$$X_{ij} = \sum_{k=1}^{p} G_{ik} \times F_{kj} + E_{ij}$$
 (1)

186

In this study, since the measurement covers a period of 12 months (full four seasons), four separate PMF inputs were prepared. Unconstrained PMF was applied with 4 to 6 factors runs for all the seasons; throughout the pre-result and while referring to previous studies (Crippa et al., 2014 and van Pinxteren et al., 2016) primary factors were separated as hydrocarbon-like OA (HOA), biomass burning OA (BBOA) and coal combustion OA (CCOA). However, unconstrained PMF did not result to separate the primary factor profiles. Introducing constraints based on prior knowledge is an efficient strategy for avoiding the mixing of primary factors (Canonaco et al., 2013; Crippa et al., 2014). For this reason, the multilinear engine (ME-2) algorithm (Paatero, 1999) enables the incorporation of time series and factor profiles constraints in form of the *a*-value approach. In dealing with a profile constraint, the *a-value* specifies the variety of a factor that can deviate from the anchor profile during the PMF iteration:

196
$$f_{i,solution} = f_i \pm a \cdot f_i$$
 (2)

The constraints applied through ME-2 for HOA and BBOA sources used the anchor profile of Crippa et al., (2014), and Ng et al., (2010), respectively. The anchor profile used for CCOA was generated from our own winter data during this work (SI, 1.1). For each of the four seasons, primary profiles were subject to a sensitivity analysis with *a-values* ranging from 0-0.4 for HOA and BBOA, 0-0.5 for CCOA, and steps of 0.1 to choose the best *a-value* combination for these three factors.

In the PMF approach, there is the intrinsic property of static factor profiles during the period of PMF analysis. Even though for short-term measurements (like one/two season/s) this might be a sensible estimation, long-term observations as are typical for current ACSM study (one year and more), are expected to be subject to evolving factor profiles based on seasonality. To consider the temporal changes, the rolling PMF window method was developed (Canonaco et al., 2021b; Parworth et al., 2015). This technique is applied to a small window, which is slowly extended throughout the whole dataset. Based on the dataset, the user determines the width of the PMF window, the shift parameter, and the number of PMF repeats per window; for the current work, we set 14-day windows, 1-day shifts, and 100 repeats per window.

208 In addition, this rolling PMF analysis was coupled with the bootstrap re-sampling approach (Bootstrap Methods: Another Look 209 at the Jackknife on JSTOR, 1979), which can randomly select a part of the original matrix and repeat a part of the rows to 210 generate a new same-sized matrix to test the stability of solutions and to estimate the statistical error. Overall, we have 211 combined rolling PMF with ME-2 and bootstrap to conduct the source apportionment investigation, and more information on 212 this new approach was described in Canonaco et al., (2020). This approach for a yearlong dataset generates an enormous 213 amount of PMF runs (N= 35800) and not all of the solutions are environmentally reasonable. Since it is practically impossible 214 to manually inspect all PMF runs, the criterial-base selection was introduced in SoFi Pro to automatically and objectively 215 select environmentally reasonable PMF solutions (Canonaco et al., 2020). Finally, the resulting factors were interpreted as 216 HOA, BBOA, CCOA, and two oxidized OA (OOA) factors named less-oxygenated OOA (LO-OOA) and more oxygenated 217 OOA (MO-OOA). The steps and setups utilized in the evaluation of this dataset are detailed in the supplement (Sect. 1).

218 <u>2.5 eBC-PM₁ source apportionment</u>

The eBC-PM₁ correlated with each of the three identified primary organic factors (HOA, BBOA, and CCOA) during the source apportionment analysis (Table. 1, which will be discussed later on in the result Sect.). The total amount of these primary factors (known as POA) was highly correlated with eBC-PM₁ (R^2 = 0.87; Fig. 8a). As a result, the different sources of eBC-PM₁ were evaluated for each factor utilizing a multilinear regression model, as suggested by Laborde et al., (2013); Zhu et al., (2018) and Poulain et al., (2021), for instance. The following assumes that the eBC-PM₁ mass is associated with the separate contribution from each OA factor (i.e., eBC-PM_{1HOA}, eBC-PM_{1BBOA}, and eBC-PM_{1CCOA}) at any time:

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227	
228	The eBC-PM ₁ emission from each source is expected to be proportionate to the separate source mass concentration generated
229	in each season (m_{HOA} , m_{BBOA} , and m_{CCOA} , respectively). As a result, the multilinear regression model can be described as
230	<u>follows:</u>
231	
232	$eBC(t) = am_{HOA} + bm_{BBOA} + cm_{CCOA} $ (4)
233	
234	where a, b, and c are the linear regression coefficients for m_{HOA} , m_{BBOA} , and m_{CCOA} , respectively, that will be applied to evaluate
235	the contribution of eBC-PM ₁ per each POA factor for each season (Table, S4).
236	2.65 Air mass trajectory analysis
230	
237	Non-parametric wind regressions (NWR) were used to approximate the OA source concentrations at a given wind direction
238	and speed (Henry et al., 2009) in order to investigate not only the local but also the prevalent wind sector associated with
239	transported emission sources (Marin, et al., 2019). Non-parametric wind regressions (NWR) were used to approximate the OA
240	source concentrations at a given wind direction and speed (Henry et al., 2009). The NOAA HYbrid Single-Particle Lagrangian
241	Integrated Trajectory (HYSPLIT-4) model was used to analyse 96 h backward trajectories at 500 m above the model ground
242	of the sampling place (Draxler & and Hess, 19972004). The trajectory results were used for two independent but
243	complementary analyses to better depict the emission areas of the aerosol: by identifying the potential areas of aerosol sources
244	area-and by clustering the trajectories.
245	A cluster analysis of the different trajectories was performed. The synoptic-scale air mass condition, together with geographical
246	locations and paths, is a crucial driver of local pollutant concentrations (e.g. Sun et al., 2020; Ma et al., 2014). Local particle
247	mass concentrations and meteorological conditions can play a significant role and be associated with specific air mass
248	trajectories. In addition, the trajectories of the air mass can influence aerosol compositions. For example, the stability of the
249	atmosphere is also meaningful since it influences both the vertical dilution of pollutants and the overall particle mass
250	concentrations. Therefore, the effects of inter-annual variations in air mass conditions and the stability of atmosphere on
251	observed patterns were inspected using a self-developed back-trajectory cluster method (BCLM), concerning air mass
252	backward trajectories, pseudo-potential temperature profiles, PM_{10} mass concentration profiles over Melpitz, and seasons
253	(Birmili et al., 2010; Ma et al., 2014). Descriptive analysis, cluster processing, and data processes and products are all described
254	in detail by Sun et al., (2020) and Ma et al., (2014).
255	In this method, the different clusters can be divided according to the different seasons (CS: cold season; TS: transition season;

- and WS: warm season), and meteorological synoptic patterns (ST: stagnant; A1: anticyclonic with air mass coming from 256
- Eastern Europe; A2: anticyclonic with air mass coming from the west; C1: cyclonic with air mass coming from relatively 257

- 258 south; C2: cyclonic with air mass coming from the west and south west). However, the clustering approach did not consider
- 259 spring and fall separately, and therefore the transition clusters correspond to both spring and fall. Finally, a total of fifteen
- 260 clusters were identified, corresponding to different meteorological conditions over the course of the year. Descriptive analysis,
- 261 cluster processing, and data processes and products are all described in detail by Sun et al., (2020) and Ma et al., (2014).

262 3 Results and discussion

263 3.1 PM₁ chemical composition

264 In this work, we investigate one-year-long measurements of PM₁ for Melpitz, Germany. All the data is-are presented in UTC, during the winter and summer, the time zone is one and two hours behind local time, respectively. Yearly time series, seasonal 265 266 variation, and diurnal cycles of aerosol particle chemical compositions including mass concentrations and mass fractions, as measured by ACSM and MAAP, are shown in figures Figures 1, 2, and 3, respectively. Over the entire period, the chemical 267 composition of PM_1 was basically made up of organic aerosol (46 % of the total mass; Fig. 1c), sulphate (16 %), nitrate (21 268 269 %), ammonium (11 %), eBC-PM₁eBC (6 %), and chloride (close to 0 %). However, a mean mass concentration of 10.497 μ g/m³ (Fig. 1) was obtained with an obvious seasonal trend which detected the highest total mass concentrations (15.95 μ g/m³) 270 271 during the winter time and lowest mass concentration during the summer time; 6.24 µg/m³ (Fig. 1a and Fig. 2a). Compared to 272 previous AMS measurements of Poulain et al., (2011) at the same station, a similar seasonal trend was observed in the period 273 2008/2009, while the absolute masses differed (Table. S1), which is at least partially related to the inter-annual changes of the 274 meteorological conditions. Compared to previous ACSM long-term measurements of Poulain et al., (2021) at the same station, a similar mean mass concentration of PM₁ was observed in the period from June 2012 to November 2017 (Poulain et al., 2021: 275 $10.23 \,\mu\text{g/m}^3$ and this study: $10.49 \,\mu\text{g/m}^3$; respectively), and presented same seasonal trends for all the chemical species (Table. 276 277 S2) with a highest mass concentration in the winter and lowest mass concentration in the summer time $(13.15 \text{ µg/m}^3 \text{ and } 7.64 \text{ m}^3)$ 278 μ g/m³, respectively; Table. S2). Consequently, the results obtained from the current study can be considered as a representative 279 ACSM study for Melpitz station. Fig. S2-S3 presents the coming high polluted air masses for total PM₁ to the measurement 280 site in the current study; the-polluted Eastern Europe flow with high mass concentration and south-west with lower mass 281 concentration was more clearly found in winter time rather than in other seasons, which will be comprehensively discussed in 282 the Sect. 3.4. 283 In comparison with other ACSM/AMS rural-background stations in Europe which can be divided into three parts Northern

- Europe (NE), Southern Europe (SE), and Mid-latitude Europe (ME) (Bressi et al., 2021), the annual PM₁ mean mass
- 285 concentration measured at Melpitz is similar to the value obtained at other ME stations, such as Magadino $10.1 \,\mu$ g/m³, Kosetice
- $286 \quad 8.5 \ \mu g/m^3$ (Chen et al., 2022), $9.1 \ \mu g/m^3$ on average of PM_1 mean mass concentration of 6 stations (Ispra, Melpitz, Magadino,
- 287 Cabauw, Sirta, and Hohenpeissenberg, Bressi et al., 2021).

288 3.1.1 Inorganic

289 The seasonality of the inorganic species can be associated with their variations in emissions and/or the changes in their 290 chemical atmospheric processes. Throughout the year, the mass concentration and their respective contribution to the total 291 PM₁ mass of nitrate, ammonium, and chloride increased from a minimum value in summer (11 %, 7 %, and 0 %, respectively; 292 Fig. 2b) and reached a maximum value in winter (24 %, 12 %, and 1 %, respectively; Fig. 2b). Moreover, the comparison 293 between Bressi et al., (2021) and current study (Fig. S4 from Bressi et al 2021, Fig. 3 from the current study) for Melpitz 294 station with different time coverage shows that the daily variation of ACSM sulphate, nitrate, and ammonium are similar in 295 both winter and summer seasons. In comparison with other ACSM/AMS rural-background stations in Europe (Fig. S4, Bressi 296 et al., 2021), the mean daily cycle of the PM₁ chemical components (sulphate, nitrate, and ammonium) does not show a similar 297 pattern to the other stations (Bressi et al., 2021) due to the different geographical location and meteorological conditions.

298 Sulphate showed a slightly different behavior. Although the contribution of sulphate to the total PM₁ decreased slightly from 299 summer (20 %) to winter (15 %), its mass concentration remained higher in winter compared to summer (2.38 μ g/m³ and 1.23 300 μ g/m³, respectively; Table. 1). The increment enhancement-is not as drastic as other inorganic species since sulphate is least 301 volatile, therefore, more fraction of sulphate stayed in particle phase even in summer. Moreover, the sulphate contribution to 302 the total PM₁ was higher during the summer than winter time, since with enhanced irradiations in summer, sulphate formation 303 from photochemistry could be enhanced as well. This sulphate higher contribution in summer over winter is consistent with 304 the mean PM_1 mass concentration measured by AMS for the three periods during fall (16. September.2008 to 03. 305 November.2008), winter (24. February.2009 to 25. March.2009), and summer (23. May.2009 to 09. June.2009) campaigns 306 reported by Poulain et al., (2011). In comparison with previous ACSM long-term measurements of Poulain et al., (2021) at 307 Melpitz station, a similar mean mass concentration of sulphate was observed in the period from June 2012 to November 2017 308 (Poulain et al., 2021: 1.54 μ g/m³ and this study: 1.67 μ g/m³; respectively; Table. S2). This comparison indicates the current 309 study as a case study of ACSM for Melpitz station within 5-year ACSM data, with the best data coverage of time in a year.-Moreover, with enhanced irradiations in summer, sulphate formation from photochemistry could be enhanced as well. This 310 311 result is consistent with the mean PM₁ mass concentration measured by AMS for the three periods during fall (16. September.2008 to 03. November.2008), winter (24. February.2009 to 25. March.2009), and summer (23. May.2009 to 09. 312 313 June 2009) campaigns reported by Poulain et al., (2011). The diurnal cycles of sulphate (Fig. 3) showed a different daily pattern 314 in warm and cold seasons. In summer, sulphate mass concentration increased during the day and reached its maximum level 315 at 12:00 UTC (Fig. 3) due to sulphur dioxide photochemical oxidation processes in the atmosphere, which also presented the 316 highest mass concentration during the day, along with maximum temperature and sun radiation in summer time (Fig. S43). 317 Furthermore, the NWR plots (Fig. S3) show that during the winter time, sulphate mostly comes from the north and east sectors 318 with wind speeds above 5 m/s which can be associated with dominant transported sulphate sources. Although the eastern wind 319 sector remains visible for the sulphate in the summer time, the high concentrations of sulphate can be observed during periods 320 with low wind speed and without a specific wind sector; which corresponds to local sulphate formation. Sect. 3.4 will go into

321 detail about the long-range transported emissions later on)... Furthermore, the wind rose analysis showed a high mass 322 concentration of sulphate at low wind speed (Fig S3). Although this locally formed emissions of sulphate (Fig. S32 and Fig. 323 9) can explain this peak during the day in summer, this photochemical process is not the only source of sulphate. It especially 324 cannot explain the highest mass concentrations during the winter time with almost no diurnal variation (Fig. 3). For winter, 325 the emission of domestic heating processes, which could be enhanced in the atmospheric boundary layer (Stieger et al., 2018), 326 along with the long-range transported emissions, which came from north-east toward the measurement site (Fig. S32 and Fig. 327 9), and also high ammonium nitrate due to partitioning according to temperature, explain the high mass concentration but the 328 low relative contribution of sulphate.

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330 Nitrate is mostly found in the form of ammonium-nitrate (NH_4NO_3), which is reliant on the gas phase precursor concentrations, 331 temperature, humidity, and aerosol chemical composition (Poulain et al. 2011; Stieger et al., 2018). Both nitrate and 332 ammonium showed a minimum mass fraction and mass concentration in summer (11 %, 0.68 µg/m³, 7 %, 0.43 µg/m³, respectively; Fig. 2), an increasing trend toward the cold months and reached their maximum mass fraction and mass 333 334 concentration in winter time (nitrate 24 %, 3.87 µg/m³, ammonium 12 %, 2 µg/m³, respectively; Fig. 2). The diurnal cycles of 335 nitrate and ammonium (Fig. 3) showed a relatively similar daily pattern in all seasons, which means the highest values were 336 reached in the morning, due to the beginning of vertical mixing and a reduction in the afternoon followed by an increase during 337 the night, reflecting their night time production during every season. The volatile behaviour of ammonium-nitrate strongly 338 affects its temporal variation during warm days leading to the formation of the gaseous nitric acid and ammonia compounds 339 at higher temperatures and low humidity (Fig. S43, and S3S8). Nitrate profiles from NWR plots (Fig. S3) present two different 340 wind directions for the whole period which might be associated with transported nitrate from Leipzig and Torgau (50 km in 341 the south-west and 7 km in the north-east of Melpitz, respectively) with higher wind speed. Since the reaction pathway of OH 342 and NO_2 can result in nitrate formation (Yang et al., 2022), this mechanism can be linked to traffic emissions in residential 343 areas. These long-range transported sources together with locally formed emissions could describe higher mass concentrations 344 for nitrate and ammonium due to e.g., meteorological conditions and abundant precursors in winter time. However, in winter, 345 ammonium-nitrate remains mainly in the particle phase (Seinfeld and Pandis, 2006) since it can totally be changed from gas 346 to particle phase at lower temperature (Spindler et al., 2010). High values of nitrate and ammonium in spring time are linked 347 to agronomical fertilization (Stieger et al., 2018)). In winter, ammonium nitrate remains mainly in the particle phase (Seinfeld 348 and Pandis, 2006) and, like sulphate, arrived at the measurement site due to the long range transported emissions which not 349 only came from the north-eastern but also south-western flow, describing higher mass concentrations for nitrate and 350 ammonium (Fig. S2). High values of nitrate and ammonium in spring time are linked to agronomical fertilization (Stieger et 351 al., 2018). These seasonal contribution results for both, nitrate and ammonium, are consistent with the previous AMS study 352 (Poulain et al., 2011), with minimum fraction to the total AMS-PM₁ during summer (nitrate 5 % and ammonium 8 %; Table. 353 S1), and maximum fraction during winter time (nitrate 34 % and ammonium 17 %; Table. S1). However, it is known that a 354 fraction of the nitrate signal can be attributed to nitrogen containing organic species (Kiendler-Scharr et al., 2016), which can

355 affect the overall nitrate mass concentration (Poulain et al., 2020).

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357 Although chloride had the lowest annual mass concentration $(0.05 \,\mu g/m^3)$ compared to all other PM₁ chemical components 358 (Table, 1), it showed the highest mass concentration and mass fraction in winter (0.11 µg/m³, 1 %, respectively; Fig. 2a&b; 359 Table. 1) compared to the other seasons; as seen in the previous AMS study of Poulain et al., (2011) (2 %, Table. S1). It could 360 be related to the surrounding and transported emissions which mass concentrations were high for air masses from north-easterly and south-westerly directions (Fig. S_{32}). In a multi-year analysis of the hourly PM₁₀ chloride mass concentration 361 measurements using a MARGA, Stieger et al., (2018) attributed the chloride sources of Melpitz during winter to the 362 resuspension of road salt used for the de-icing of streets, mainly coming from the cities of Torgau and Leipzig. These sites are 363 364 also located in the wind directions along with the coal and wood combustion emission region, which could explain the highest 365 mass concentration of chloride during the winter. Furthermore, the existence of chloride might be due to low mass 366 concentration marine influences consisting of sea-salt aerosols during all the seasons in the south-westerly direction (Fig. S32) 367 which was-were previously studied by Stieger et al., (2018). However, it is known that the AMS -technology cannot properly detect sea salt (S-Huang et al., 2018; Ovadnevaite et al., 2014) because the majority of chloride is in the refractory part which 368 369 cannot be flash vaporized at 600 °C. Consequently, the chloride detected by the ACSM is mostly related to combustion 370 processes (wood, coal combustion as well as trash burning; Li et al., 2012).

371 **3.1.2** <u>eBC-PM₁eBC</u> and organics

372 The eBC-PM₁eBC showed its maximum mass concentration and mass fraction to PM mass during winter time at $1.38 \,\mu g/m^3$ 373 and 9 %, respectively (Fig. 2), and only $0.25 \,\mu g/m^3$ and 4 %, respectively, during summer time (Fig. 2). This is consistent with 374 the expected highest anthropogenic emissions from fossil fuel consumption (house heating and energy productions) in winter 375 compared to summer (Spindler et al., 2010). Furthermore, considering measured eBC-PM₁ in regard to wind speed and wind 376 direction from NWR plots (Fig. S3), eBC-PM₁ presented transported and local emissions. The highest mass concentrations for 377 fall, winter, and spring seasons could be linked to north-easterly and south-westerly winds with higher wind speed (above 10 378 m/s). While in summer time it is mostly linked to the surrounding emissions regardless of wind direction with lower wind 379 speed (Fig. S3). Furthermore, considering measured eBC in regard to wind speed and wind direction (Fig. S2), the highest 380 mass concentrations could be linked to north easterly and south westerly winds for fall, winter, and spring seasons, while in 381 summer time it is mostly linked to the surrounding emissions (Fig. S2). Significant changes in the diurnal profiles of eBC-382 PM₁eBC for the different seasons can be found with the highest mass concentrations throughout the cold months compared to 383 warm months owing to house heating (Fig. 3). It also showed morning and evening peaks during all seasons (Fig. 3). This is 384 consistent with those observed for the nitrogen oxides (Fig. S43), which might be attributed to liquid fuel emissions and 385 possibly the impact of the traffic rush hours on the main street, B 87, located approximately 1 or 1.5 km north of the station, 386 (Yuan et al., 2021). In the following chapter, diurnal patterns showed lower mass concentrations at noon, and increased in the late afternoon to become nearly constant from 8 p.m. until midnight (Fig. 3). This ambient particulate pollution resulting from very surrounding sources in the village was reported by van Pinxteren et al., (20202). Diurnal increments of <u>eBC-PM1eBC</u> were smaller in fall and spring compared to winter; the increment in summer is also correspondingly low due to the absence of house heating emissions, and the diurnal variation in the increment is determined by surrounding motor vehicle emissions in combination with the mixing layer height (van Pinxteren et al., 20202). Further discussions on the seasonal trend of the <u>eBC-PM1eBC</u> can be found in Sect. 3.3.

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Organic aerosol (OA) was the predominant species throughout the whole year, with a mean mass concentration of $4.84 \,\mu g/m^3$ 394 395 and a mass fraction of 46 % (Fig. 1c; Table. 1). The OA mass fraction decreased from the maximum value in summer and 396 attained a minimum mass fraction in winter (58%, 39%, respectively; Fig. 2b). Similar to the comparison of previous inorganic 397 AMS measurements performed at Melpitz (Poulain et al., 2011), AMS-OA contribution to total PM₁ showed maximum 398 contribution during summer (59%, Table, S1), and minimum contribution during winter (23%) as well. However, the mass 399 concentration of OA increased from its lowest value in summer and reached its highest value in winter time $(3.67 \,\mu\text{g/m}^3, 6.21 \,\mu\text{g/m}^3)$ 400µg/m³ respectively; Fig. 2, Table. 1). Similar to eBC-PM₁eBC, by analyzing NWR plots, OA measured in according to wind 401 direction and wind speed showed the highest average mass concentrations for north-easterly and south-westerly winds in 402 winter (Fig. S32). In fall, polluted air masses came from the north-easterly direction, and in spring and summer OA, 403 surrounding emissions closer to Melpitz were identified (Fig. S32). The diurnal cycle of the organic had an identical pattern 404 across all seasons (Fig. 3), showing the highest mass concentration in night time, a small peak in the early hours of the morning 405 related to rush hours, and the lowest mass concentrations around the early afternoon. The peak observed around 12:00 UTC in 406 summer time (Fig.3) can be due to the local photochemical production that leads to the formation of secondary organic aerosol 407 mass during the day, similar to the diurnal behavior of sulphate (previously discussed in Sect. 3.1.1). However, the reduction 408 in total OA mass concentration throughout the day (Fig. 3), which was mostly observed during the warm seasons (spring and 409 summer), could be clearly related to the dilution effect of increasing mixed layer height. During warm days, evaporation of 410 semi-volatile organics from the particle phase cannot be completely excluded (Schaap et al., 2004; Keck and Wittmaack, 411 2005). In comparison between Bressi et al., (2021) and the current study for Melpitz station, the daily variation of organic are 412 similar in both winter and summer seasons, while there are differences between Melpitz with other rural-background stations 413 due to the different geographical location and meteorological conditions (Bressi et al., 2021),-

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416 Overall, <u>eBC-PM1eBC</u> and OA can be composed of various sources with strong seasonal dependencies, as well as be
417 influenced by different responses to atmospheric dynamics depending on meteorological parameters, geographical locations,
418 and chemical processes. Therefore, a comprehensive analysis of the OA and <u>eBC-PM1eBC</u> sources was performed using source
419 apportionment techniques.

420 3.2 Source apportionment of OA

421 The chosen solution for the organic aerosol source apportionment contained five different factors based on their time series 422 and mass spectra (Fig. 4). The source apportionment solution is based on a partly constrained rolling approach with three 423 primary organic factors (POA), namely HOA (on average $0.30 \,\mu g/m^3$ and 6 % of the total OA; Table.1 and Fig. 4), BBOA (on average 0.39 μ g/m³ and 7.9 % of the total OA) and CCOA (on average 0.77 μ g/m³ and 15.4 % of the total OA). In addition to 424 425 these POA factors, two oxygenated organic aerosols (OOAs) were identified as LO-OOA (on average 1.62 µg/m³ and 32.4 % of the total OA), and MO-OOA (on average $1.92 \,\mu \text{g/m}^3$ and $38.4 \,\%$ of the OA). The seasonal average mass concentrations and 426 427 relative mass fractions of each OA factor to the total OA mass and their seasonal diurnal variation are presented in Figures 5 428 and 6; respectively. They will be discussed separately in the following sections.

429 3.2.1 POA factors

430 The HOA mass spectrum (Fig. 4b) is recognized by mass fragments at unsaturated and saturated hydrocarbon chain pairs m/z. 431 41 (C_3H_5), 43 (C_3H_7), m/z 55 (C_4H_7) and 57 (C_4H_9) (Zhang et al., 2005; Canagaratna et al., 2004), which are representative of 432 liquid fuel combustion emissions and are associated with either traffic emissions or domestic heating fuel (Wang et al., 2020). 433 This result designates HOA as a minimal source of OA at the monitoring site, which is consistent with previous studies in the 434 PM_1 range made in the same place: a) total average was 7 % of the organic mass concentration in a study by Crippa et al., 435 (2014) total average was 3 % of PM size range between 0.05-1.2 µm mass concentration in a study by van Pinxteren et al., 436 (2016) (Table, S₃²). However, in comparison with other ACSM/AMS stations in Europe (22 stations; Chen et al., 2022), 437 Kosetice with 9.7 % as a rural-background site, and Bucharest with 13.7 % as an urban-background site showed the minimum 438 annual HOA mean contribution of total OA, which is similar to the contribution at Melpitz.

439 Mass concentration of HOA followed a slightly increasing seasonal pattern towards the cold months, from 0.23 μ g/m³ in 440 summer to $0.36 \,\mu\text{g/m}^3$ in the winter (Fig. 5a; Table. 1). HOA presented a low correlation with nitrogen oxides over the entire 441 period ($R^2 = 0.17$, Table. 1), but it correlated well with eBC-PM₁eBC in winter ($R^2 = 0.52$; Table. 1) and showeds a weaker 442 correlation in summer (R^2 = 0.28; Table. 1). Possibly HOA is also associated with household heating (35 % by oil and 11 % 443 by liquid petroleum gas, van Pinxteren et al., 20202) rather than traffic emissions, especially during the cold months. Analyzing 444 the NWR plots demonstrates the highest HOA mass concentration was observed at low wind speed during the warm period 445 (Fig. 7) indicating, rather local emission sources. While during the cold period a clear increase of the mass concentration can be associated with the highest wind speed (> 10 m/s) mostly coming from the North to East sector. During periods with wind 446 447 speeds below 10 m/s, the two dominant wind sectors (NE and SW) can be observed. The first one might be associated with 448 emission plumes coming from either the surrounding traffic emissions (the federal street B 87), as well as the domestic 449 emissions are associated not with house heating in summer but with hot water production (van Pinxteren et al., 2020), as well 450 as the city of Torgau (with approx. 20,000 inhabitants, distance from 7 km). Although the SW sector shows a lower HOA mass 451 concentration in comparison to the NE one, it corresponds to the direction of the city of Leipzig (above 600 000 inhabitants,

- 452 approx. 50 km). Therefore, it might be associated with the influence of the pollution plume of the city of Leipzig. Analyzing 453 the pollution wind rose, the highest winter HOA mass concentrations are associated with the north easterly wind direction 454 regardless of the wind speed suggesting the influence of long range transported emissions (Fig. 7). During the warm months, 455 the emissions were more from the surrounding area, still associated with a north easterly wind direction but only at low wind 456 speed (Fig. 7), which might be associated with either the surrounding traffic emissions, as well as the domestic emissions 457 associated not with house heating in summer but with hot water production (van Pinxteren et al., 2020).
- 458 The diurnal patterns of HOA reproduced two peaks in the morning and evening for all seasons (Fig. 6), which is related to 459 traffic rush hours and linked to surrounding emissions from the main street (B 87, approx, 1.5 km north of the station), Melpitz 460 village itself, and emissions coming from Leipzig and Torgau residential areas The diurnal patterns of HOA reproduced two 461 peaks in the morning and evening for all seasons (Fig. 6). The small time shift for the start of the evening increase corresponds 462 to the time shift of the sunrise between winter and summer. The diurnal cycles reached a systematic minimum during the day 463 time probably not only owing to emission decrease but also emphasizing the effect of dynamic atmospheric processes (e.g. 464 mixing layer height (MLH) and planetary boundary layer (PBL)) (Fig. 6, and S4). Oppositely to what can be seen during the 465 day time, night time mass concentrations appeared to be unaffected by the seasons, showing similar mass concentrations all 466 year round, i.e. their mass concentration rose continuously in the early evening and remained at a very similar mass 467 concentration over the night, which supports the hypothesis of yearlong continuous rather surrounding emissions. 468 Nevertheless, the differences between HOA mass concentration during the night time from summer to winter season (Fig. 6) 469 are small and can be covered by the uncertainties of PMF result (± 32.5 %, Fig. S2), however, it can be explained by different 470 emission sources, condensation of POA (Chen et al., 2022), evaporation, oxidation processes (Saha, et al., 2018), and potential 471 night time aging process by high ozone concentration (Kodros, et al., 2020).
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- 473 The mass spectra of BBOA are identified by ions at m/z 29, 43, 60, and 73 (Fig. 4b), known as fragments tracers of anhydro-474 sugars like levoglucosan (Alfarra et al., 2007), which have been identified recognized as indicators of wood combustion 475 processes (Simoneit et al., 1999; Simoneit & and Elias, 2001). This is confirmed by the correlation between BBOA and 476 levoglucosan over the whole period (R^2 = 0.65; Table. 1). On average, BBOA mass concentration and contribution were 0.39 477 µg/m³ and 7.9 %, respectively (Table. 1 and Fig. 4a). However, its contribution is highest during winter time (10.6 %; Fig. 5), 478 which is similar to previous studies in different PM ranges for the Melpitz station during the cold months: a) in PM₁ range, 14 479 % of OA mass concentration in fall (Crippa et al., 2014); b) in 0.05-1.2 µm range, highest contribution with 10 % of PM mass 480 concentration in winter (van Pinxteren et al., 2016); and c) in PM₁₀ range, highest contribution with 16 % of PM mass 481 concentration in winter (van Pinxteren et al., 20202).
- 482 By analyzing the NWR model, the high mass concentration of BBOA in cold months, regardless of wind speed can be observed
- 483 with two wind sectors coming from north-east and south-west directions. These BBOA emissions are mainly attributed to
- 484 residential heating in Melpitz village and also indicate the effect of transported biomass burning emissions to the sampling site
- 485 with higher wind speed (> 10 m/s, Fig. 7). The high value of BBOA is mainly attributed to residential heating, and indicates

- the effect of transported biomass burning emissions to the sampling site in cold months (Fig. 7), Wwhile in summer time, it is still observable as surrounding emissions during periods of low wind speed (Fig. 7 and Fig. S4) with a mass concentration of 0.21 μ g/m³ and a contribution of 6.1 % to total OA (Fig. 5). The presence of BBOA in the summer can be linked to water heating systems using wood briquettes and logs (estimated at 32 % of total central heating in this area, van Pinxteren et al., 202<u>0</u>9). Moreover, it can also be related to recreational open fires and/or barbecue activities (van Pinxteren et al., 202<u>0</u>2). This result is similar to other ACSM/AMS rural-background stations in Europe (22 stations; Chen et al., 2022); both Magadino and Kosetice showed the highest contribution of BBOA during winter time (27.4 % and 15.5 % respectively).
- The diurnal cycles, peaking from early evening to early morning in winter (Fig. 6), match the expectations for a factor related to domestic heating activities, along with a better <u>eBC-PM1eBC</u> correlation during winter than during summer time (R^2 = 0.81, and R^2 = 0.42, respectively; Table. 1). Finally, in opposition to HOA, the night time BBOA mass concentration showed a strong seasonal variation, having its highest mass concentration during winter nights and lowest during summer time, the influence of the impact of house heating emissions on the BBOA emissions. However, the day time behavior reflects the influence of enhanced vertical mixing during day time (higher temperature, Fig. S43) combined with high wind speeds (Fig. 11) can readily cause dilution and thus low pollutant concentrations near the ground (Chen et al., 2021; Via et al., 2020; Paglione et al., 2020).
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501 The mass spectrum of CCOA is characterized by fragments at m/z 77, 91, and 115 (Fig. 4b) as previously reported by Dall'Osto 502 et al., (2013); Xu et al., (2020); Tobler et al., (2021) and Chen et al., (2022). These specific fragments can be associated with 503 unsaturated hydrocarbons, particularly ion peaks related to polycyclic aromatic hydrocarbon (PAH). The CCOA time series 504 showed the strongest correlation with eBC-PM₁eBC (R^2 = 0.9; Table. 1). In addition, several studies reported that coal 505 combustion emissions are often accompanied by high chloride mass concentrations (e.g. Iapalucci et al., 1969; Yudovich & 506 and Ketris, 2006 and Tobler et al., 2021). Here, the correlation between CCOA and chloride was higher during winter than 507 during summer time ($R^2 = 0.41, 0.15$ respectively; Table. 1), as the gas-particle phase equilibrium dramatically changes with 508 rising temperatures (Tobler et al., 2021). Although chloride is almost observable in the particle phase as ammonium chloride 509 (NH₄Cl) at lower temperatures, chloride is typically observable in the gas phase as hydrogen chloride (HCl) at higher 510 temperatures (Tobler et al., 2021).

CCOA represented on average 15.4 % of the total OA (0.77 µg m⁻³), (Table. 1; Fig. 4a) and is the most important POA over 511 512 the entire period. No CCOA factor was identified in the previous AMS measurements made at Melpitz (Crippa et al., 2014). 513 Most likely this factor was not properly resolved and/or it was not possible to properly separate it from the other factors since 514 no reference mass spectra for CCOA was-were reported in the literature at that time. CCOA showed the highest mass 515 concentration and mass fraction during the winter (1.58 μ g/m³, 23 %, respectively; Fig. 5a; Table. 1). By analyzing the NWR 516 plots, this high mass concentration during winter time can be related to the surrounding emissions and long-range transported 517 air masses coming from two different directions, north-easterly and south-westerly (Fig. 7). CCOA showed the highest mass 518 concentration and mass fraction during the winter (1.58 µg/m³, 23 %, respectively; Fig. 5a; Table. 1), which is related to the 519 surrounding emissions and long range transported air masses coming from two different directions, north easterly and south520 westerly (Fig. 7). Not surprisingly, the lowest mass concentration and contribution were observed during the summer time 521 $(0.30 \text{ µg/m}^3, 8.7 \%, \text{ respectively; Fig. 5a; Table. 1.)}$ which most probably correspond to only long-range transport as later 522 discussed in Sect. 3.4 (Fig. 9). Moreover, this result is consistent with previous measurements made in the same place. For the 523 size range 0.05-1.2 µm van Pinxteren et al., (2016) reported a contribution of 29 % and 21 % of the PM in winter and summer 524 respectively, and a contribution of 7 % and 0 % for winter and summer respectively for the PM_{10} range was found (van 525 Pinxteren et al., 20202). From all ASCMACSM/AMS stations (22 stations; Chen et al., 2022) only Melpitz as a rural-526 background site and Krakow as an urban-background site showed the coal combustion emissions with the maximum 527 contribution during winter for both sites (Krakow: 18.2 % and Melpitz: 23 %) compared to summer (Krakow: 4.5 % and 528 Melpitz: 8.7%). The drastic seasonal changes in Krakow are attributed to the common use of coal burning for residential 529 heating reasons during the winter time (Casotto et al., 2022; Tobler et al., 2021), while in Melpitz, as discussed above, coal 530 combustion is affected by both surrounding and transported emissions from other sites.

531 Mass concentrations of CCOA during night time were much higher than during day time throughout all seasons (Fig. 6), further

532 verifying the increased coal combustion emissions from coal heat generation at <u>night in</u> winter time night and the potential

533 decrease in emissions during the day due to a strong influence of atmospheric dynamics.

534 **3.2.2 OOA factors**

- 535 The two OOAs (Fig. 4) referred to as LO-OOA and MO-OOA are known to be characterized by the different ratios of their 536 m/z 43 and m/z 44 fragments (Fig. 4b), that-which represent the oxidation level (Canagaratna et al., 2015). While m/z 43 could 537 be derived from $C_2H_3O^+$ (a signature of the semi-volatile) and/or C_3H7^+ (a signature of the primary emissions of the 538 hydrocarbon-like), m/z 44 is mainly derived from the fragment of CO_2^+ (a signature of oxygenated, particularly acids) 539 (Canonaco et al., 2015; Ng et al., 2010). As presented in Fig. 4b, MO-OOA mass spectra showed a notable peak at m/z 44. 540 This spectrum has been extensively recognized as low volatility OOA (LV-OOA) and described to be made--up of aged 541 secondary OA (SOA) and highly oxidized OA (Lanz et al., 2007; Ulbrich et al., 2009; O-Zhang et al., 2011; Ng et al., 2011b); 542 while the mass spectra of LO-OOA in this study presented a higher m/z 43 (Figs. 4b) compared to MO-OOA, which is similar 543 to the mass spectral pattern of the previously reported freshly formed semi-volatile OOA (SV-OOA) (Jimenez et al., 2009; 544 Ng et al., 2010). To differentiate the variations of the OOAs factor, the f44 vs f43 space was used which is a typical diagnostic 545 tool based on atmospheric aging (Ng et al., 2010).
- The seasonal *f*44/*f*43 for OOAs measured points and the *f*44/*f*43 for modelled factor profiles (LO-OOA and MO-OOA) are presented in Fig. S<u>5</u>4. The data points in Fig. S<u>5</u>4 are distributed differently according to the season (Chen et al., 2021; Canonaco et al., 2015; Crippa et al., 2014; Chazeau et al., 2022). Furthermore, the modelled factor profile points represent a high variability in space, especially for LO-OOA. This assumes how an annual or seasonal PMF solution, unless a larger number of factors are used, would perform poorly in capturing all of the variations of SOA. In order to capture time-dependent changes, in particular for LO-OOA, it is, therefore, advantageous to perform rolling PMF analysis. The triangle plot defined by Ng et al. (2010) is also shown in Fig. S<u>5</u>4. As assumed the LO-OOA points were concentrated in the lower part of the space,

whereas more aged MO-OOA points relocated to the upper part of the space during the aging process. The fall, spring, and summer data points were all located on the right side of the triangle (Fig. S54), however, the winter data points were located near the top and inside the triangle. The data points on the right side of the triangle correspond to the time exposed to higher temperatures more than those that are within the triangle. This could be attributed to an increase in biogenic SOA emissions if the temperature increased, as biogenic OOA appears to be dispersed all along the right the side of the triangle. Furthermore, as the temperature is reduced, the increased biomass emissions cause the OOA points to lie vertically inside the triangle, as seen in the winter data.

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561 The two OOAs were the two most significant contributors to the total OA fraction (Fig. 4) over the entire period. The seasonal 562 mean mass concentrations of MO-OOA varied from higher mass concentrations during winter $(2.25 \,\mu g/m^3)$ and lower during 563 summer time (1.44 µg/m³, Table. 1). However, the highest MO-OOA mass concentrations found during the cold periods are 564 similar to the seasonal patterns in POA. This high mass concentration in cold seasons can be seen from the NWR plot (Fig. 7) 565 presenting local emissions with low wind speed (> 5 m/s) and transported emissions from east, north-east, and south-west 566 directions with high wind speed (< 5 m/s). -Furthermore, high mass concentrations of MO-OOA are generally found at high 567 relative humidity (RH > 80 %) and low temperature (< 0 $_^{\circ}C$), i.e., conditions during winter time (Fig. S₀5). This low air temperature condition can be linked to a possible scenario for an increase in the MO-OOA precursor emissions from biomass 568 569 burning and coal combustion as a result of residential heating activities during winter time. Therefore, significant enhancement 570 appears to be an effect of RH during winter, proposing that the aqueous-phase heterogeneous mechanisms could also play a 571 crucial way in the regional MO-OOA formation through winter as suggested by Gilardoni et al., (2016). In contrast, no RH-572 temperature-dependent trends for the MO-OOA were found in the other seasons (Fig. S_{5}), indicating more complex formation 573 processes during other seasons. Meanwhile, MO-OOA diurnal cycles presented a seasonal variation as well, with a remarkable 574 enhancement in the evening and night time during winter (Fig. 6), indicating a potential regional formation mechanism 575 containing night time chemistry (Tiitta et al., 2016), and descending pattern from night time to day time due to planetary 576 boundary layer effectMeanwhile, MO OOA diurnal cycles presented a seasonal variation as well, with a remarkable 577 enhancement in the evening and night time during winter (Fig. 6), indicating a potential regional formation mechanism 578 containing night time chemistry (Tiitta et al., 2016).- While in fall, spring, and summer, MO-OOA displayed a considerable 579 increase during the day (Fig. 6), indicating that higher temperatures result in considerable regional photochemical production 580 of SOA particles (Fig. S43) and enhanced solar radiation (Petit et al., 2015). Furthermore, regarding the correlation of mass 581 concentration of MO-OOA with sulphate, the latter is regarded as a local secondary production indicator (Petit et al., 2015, and Table. 1). Consequently, alongside almost stable mass spectra throughout the year, MO-OOA seems to be derived from a 582 583 variety of seasonal-dependent formation mechanisms and sources (such as aged background, biomass burning, coal 584 combustion, and biogenic sources).

585

586 The seasonal mean mass concentrations of LO-OOA varied from higher mass concentrations during fall $(2.13 \, \mu g/m^3)$ and 587 lower mass concentrations during spring time (1.24 μ g/m³, Table. 1). Temperature had a significant effect on LO-OOA, and 588 showed a distinguishable seasonal variation pattern. The temperature-RH dependence of the LO-OOA was not quite similar 589 depending on the season (Fig. S_{65}). The highest winter time LO-OOA mass concentrations were found mostly at low 590 temperatures and high RH environments, indicating that gas-particle partitioning might have a key role in LO-OOA formation 591 throughout this season. The freshly formed SOA deriving from primary biomass burning and coal combustion emissions, as 592 found in previous studies (Crippa et al., 2013; Zhang et al., 2015; Y. Sun et al., 2018; Stavroulas et al., 2019) can also affect 593 the LO-OOA during the cold months. Furthermore, since nitrate could be originated locally or arrived from a long distance to 594 Melpitz (Sect. 3.1.1), with a good correlation between LO-OOA and nitrate ($R^2 = 0.59$) during winter, the long-range 595 transported LO-OOA from different directions reaching to measuring site could be explained (Fig. 7). Furthermore, during 596 winter time the correlations between LO-OOA and nitrate (R^2 =0.59) were found. Different LO-OOA daily cycles were also 597 found in different seasons (Fig. 6). The daily changes in LO-OOA displayed higher mass concentrations in night time compared 598 to day time in fall, spring, and summer (Fig. 6), highlighting the significant roles of night time chemistry and/or gas-particle 599 partitioning in the LO-OOA formation; while the decrease during the day is partly linked to the atmospheric dilution effect 600 (Fig. S43), evaporation and photochemical aging into MO-OOA (Fig. 6). For winter night increments, lower temperature in 601 favor of condensation; and more abundant precursors present considering increased BBOA emission, therefore enhanced night-602 chemistry activities, leads to higher LO-OOA; moreover, shallow boundary layer in winter and night time inversion caused 603 pollutants to accumulate.

604 3.3 eBC sSource apportionment of eBC-PM1

By applying a multilinear regression model, The during the source apportionment analysis, eBC-PM₁ correlated with every one of the three identified primary organic factors (HOA, BBOA, and CCOA, eBC correlated with each of the three identified primary organic factors (HOA, BBOA, and CCOA) during the source apportionment analysis (Table. 1)). The total amount of these primary factors (known as POA) was highly correlated with eBC (R^2 = 0.87; Fig. 8a). As a result, the different sources of eBC were evaluated for each factor utilizing a multilinear regression model, as suggested by Laborde et al., (2013); Zhu et al., (2018) and Poulain et al., (2021), for instance. The following assumes that the eBC mass is associated with the separate contribution from each OA factor (i.e., eBC_{HOA}, eBC_{EBDA}, and eBC_{CCOA}) at any time:

612

613	$\rho RC(t) = \rho RC$ (t)	$+ eBC_{BBOA}(t) + eBC_{e}$	(t)	(3)
015	$-c_{BC}(t) - c_{BC}(t)$	$BBOA(c) + c D c_{BBOA}(c) + c D c_{f}$	CCOA (V)	-(3)

614

615 The eBC emission from each source is expected to be proportionate to the separate source mass concentration generated in 616 each season (m_{HOA} , m_{BBOA} , and m_{CCOA} , respectively). As a result, the multilinear regression model can be described as follows: 617

618 $eBC(t) = am_{HOA} + bm_{BBOA} + cm_{ccOA}$

(4)

- 619
- 620 where a, b, and c are the linear regression coefficients for m_{HOA} , m_{BBOA} , and m_{CCOA} , respectively, that will be applied to evaluate
- 621 the contribution of eBC per each POA factor for each season (Table. S3)..
- 622 CCOA appeared to <u>have be</u> the largest source of <u>eBC-PM_1eBC</u>, contributing half of it (<u>eBC-PM_1eBC</u>-CCOA 55 %, Table. 1), 623 followed by <u>eBC-PM_1eBC</u> associated with BBOA 37 % (<u>eBC-PM_1eBC</u>-BBOA), while the lowest contribution was found for 624 <u>eBC-PM_1eBC</u>-HOA (8 %). However, the contribution of sources to the total <u>eBC-PM_1eBC</u> strongly depends on the season.
- 625 Looking at each individual source, the hydrocarbon-like emissions contributed most to the eBC-PM₁eBC fraction in the fall 626 (eBC-PM₁eBC-HOA with 22 %, Table. 1; Fig. 8b), while biomass burning emissions dominated the eBC-PM₁eBC in summer 627 and coal combustion emission dominated in winter (eBC-PM₁eBC-BBOA and eBC-PM₁eBC-CCOA with 69 % and 56 %, Table. 1). In the diurnal cycle, contribution to the total eBC-PM₁eBC of eBC-PM₁eBC-HOA showed two peaks in the morning 628 629 and evening for fall, spring, and summer (Fig. S76), reflecting the impact of the traffic rush hours as mentioned in Sect. 3.2.1, 630 and the minimum contributions during the day time due to the effect of lowest emissions and PBL effect (Fig. S43). However, 631 winter time did not show a strong variation in the diurnal cycle (Fig. S76). This indicates the potential influence of continuous 632 emissions at the measurement site. Biomass burning combustion with its maximum contribution during the day in summer 633 (Fig. S76) can be related to a variety of different eBC-PM₁eBC-POA mass concentrations (Fig. S76b), while the BBOA mass 634 concentration was almost constant, the other POA mass concentration decreased during the day. Coal combustion showed an 635 increasing contribution during night time in all the seasons (Fig. S756), especially during the winter time, which further 636 confirms the enhanced coal combustion emission in winter nights (Fig. S76b).
- 637

638 3.4 Impact of airSeasonal air mass clustering mass origin and trajectory analysis

639 As mentioned before, the geographical origin of the PM_1 chemical species and also PMF components are not only emitted 640 from the surrounding area but also transported. Therefore, to better identify the origin of their sources, trajectory analysis, and 641 their clustering analysis were applied using the self-developed back-trajectory cluster method (BCLM) (Sun et al., 2020; Ma 642 et al., 2014; Hussein et al., 2006). A total of fifteen clusters were identified, corresponding to different meteorological 643 conditions over the course of the year at Melpitz (Fig. 9a). The different clusters can be divided according to the different 644 seasons (CS: cold season; TS: transition season; and WS: warm season), and meteorological synoptic patterns (ST: stagnant; 645 A1: anticyclonic with air mass coming from Eastern Europe; A2: anticyclonic with air mass coming from the west; C1: 646 eyclonic with air mass coming from relatively south; C2: cyclonic with air mass coming from the west and south west). 647 However, the clustering approach did not consider spring and fall separately, and therefore the transition clusters correspond 648 to both spring and fall. Regarding this cluster approach, six air masses were identified at Melpitz station for the winter season, 649 four air masses for the transition seasons, and five air masses for the summer season (Fig. 9a). The number of clusters with 650 their corresponding mean mass concentration of PM₁ chemical species and PMF factors of organics are summarized in Table. 651 2 and with more details in Tables S3-S4 and S4S5.

652 3.4.1 Winter

653 Fig. 9b and 9c illustrate the mass concentration and contribution of PM₁ chemical species and PMF factors of organic for each 654 air mass type at Melpitz based on the type of air masses. For the winter season, the cluster CS-ST corresponds to more 655 surrounding emission origin with a PM mean value of $21.95 \,\mu g/m^3$, which occurred during 14 % of the total measurement 656 period. These surrounding emissions refer to the emissions from Melpitz station directly, Melpitz village, and short distance 657 transported particles like particles from Leipzig and Torgau. Thi. This cluster presented the highest mass concentration of LO-OOA to the PM mass (2.73 µg/m³). In fact, SOA is considered to be formed by biomass burning as well as coal combustion, 658 659 particularly during the winter when biogenic emissions and UV radiation are low (Lanz et al., 2010; Kodros, et al., 2020). In 660 this condition and in the presence of NO_2 and O_3 , the biomass burning emissions could age rapidly and produce SOA. In 661 conclusion, this cluster could confirm the role of freshly formed SOA which originated from the primary biomass burning and 662 coal combustion emission (mass concentrations of 0.97 μ g/m³ and 1.89 μ g/m³, respectively). s cluster with the highest mass concentration of LO OOA to the PM mass (2.73 µg/m3) could confirm the role of freshly formed SOA originating around the 663 station from primary biomass burning and coal combustion emissions (mass concentration of 0.97 µg/m3 and 1.89 µg/m3, 664 665 respectively). Furthermore, nitrate showed a high mass concentration and contribution in this air mass (5.38 μ g/m³ and 25 %, respectively) due to e.g., meteorological conditions and abundant precursors. 666

- 667 The cluster CS-A1 with the highest mass concentration of PM (29.14 µg/m³) represented Eastern European continental air masses (passing Poland and the Czech Republic) during anticyclonic flow which occurred during 18 % of the total 668 measurement period, meaning that Melpitz was under their influence during winter. This air mass, with the highest POA mass 669 670 concentration (5.56 µg/m³), especially coal combustion emissions (CCOA and eBC-PM₁eBC-CCOA with an average mass concentration of 4.01 μ g/m³ and 1.93 μ g/m³, respectively), highlight the importance of long-range transported emissions. This 671 672 cluster also contained the highest mass concentration of sulphate (5.39 μ g/m³) and can-could support the importance of coal 673 combustion on sulphate formation, which is known to be strongly emitted by coal power plants (Wierońska-Wiśniewska et al., 674 2022).
- The air mass CS-A2 identified as marine-influenced air <u>mass</u> with a mean value of 13.39 μ g/m³ of PM came from the United Kingdom with the anticyclonic flow, which occurred during 8 % of the total measurement period. This cluster presented a low mass concentration of POA and for two OOAs almost the same mass concentration and contribution (Table. <u>S3-S5</u> and Table. <u>S4S6</u>). Since Melpitz is placed away from the coast, therefore the sampling location is affected by aged maritime air masses (Poulain et al., 2011). Inorganics are dominated by nitrate in this cluster with the high mass concentration (3. 86 μ g/m³) and which represents the highest mass fraction (50 % of the total inorganic species).
- 681 The CS-C1 air mass with a mean value of $15.99 \,\mu g/m^3$ characteristic of Southern European air mass, came from an industrial
- and polluted area starting from Spain and partly crossing Italy with the cyclonic flow, which occurred during 10 % of the total
- 683 measurement period. POA mass concentration and contribution were low in this cluster, while SOA, especially MO-OOA,
- 684 showed the highest mass concentration of PM over the entire period ($3.77 \,\mu g/m^3$) and the highest contribution during the winter

season (24 %). This can be linked to the high sulphate in this air mass (2.99 μ g/m³), which showed that the regional influence by contribution from aged BBOA and CCOA might be manifested in MO-OOA (as discussed in Sect. 3.2.2).

687 Finally, CS-C2a and CS-C2b were both associated with cyclonic and marine influence conditions which only occurred for a 688 short time (3 % and 2 % of the total measurements, respectively), showing the lowest PM mean value (4.09 μ g/m³ and 2.60 689 μ g/m³, respectively). Both of them showed almost the same mass concentration and contribution of POA (Fig. 8a and b; and 690 Table. \$3-\$5 and \$4\$6). However, similarly to CS-A2, cluster CS-C2a contained a marine component at the beginning point 691 of the air masses, and in the following time it was dominated by continental areas (France and southern Germany), where due 692 to the longer time transferring over continent and aging process, it showed more nitrate mass concentration and contribution than CS-C2b (1.35 µg/m³, 16 µg/m³; and 28 % 14 %, respectively). Whereas CS-C2b started near Iceland with same history 693 694 of the air mass over the continent, and in comparison, with CS-C2a, it presented a higher contribution of sulphate (29 % and 695 19 %, respectively), which could be associated with aged marine air mass due to the higher contribution of MO-OOA (21 % 696 and 18 %, respectively).

697 3.4.2 Transition seasons

698 For transition seasons (fall and spring), whereas the four clusters showed a quite similar PM mass concentrations (Fig. 9) 699 which might be linked to the overall weather situation during these two times of the year, their chemical composition strongly 700 depended on their origins. TS-A1 and TS-A2 corresponded to two different types of anticyclonic air masses with respective 701 mean PM mass concentrations of 6.06 μ g/m³ and 5.86 μ g/m³. Cluster TS-A1 which occurred during 4 % of the total 702 measurements period, started from Finland, crossing the Estonian, Latvian, Lithuanian, and Polish coasts before arriving at 703 Melpitz. Although it might contain a certain marine component, this cluster mostly followed coastal areas, which means that 704 in this cluster OA mass concentration dominated PM (2.95 μ g/m³). Furthermore, this cluster showed continental and polluted 705 aspects with the highest LO-OOA mass concentration and contribution during transition seasons (1.03 μ g/m³ and 17 % 706 respectively), which is linked to originating from freshly formed SOA from primary biomass burning and coal combustion 707 emissions around coastal areas. On the other hand, cluster TS-A2 (4 % of the measurements period) is characterized as a 708 marine cluster and started from the south of Iceland/Greenland. This cluster showed inorganics as the dominant components 709 in PM with a high mass concentration and a mass fraction (3.35 μ g/m³ and 58 % respectively). Since Melpitz is influenced by 710 aged marine air masses, this cluster showed a maximum nitrate mass concentration during the transition seasons $(1.54 \,\mu g/m^3)$ 711 and a contribution of 26 %, respectively).

- Finally, two other clusters TS-C1 and TS-C2 were two different types of cyclonic air masses in fall and spring time, with mean PM mass concentrations of $4.69 \,\mu$ g/m³ and $4.94 \,\mu$ g/m³ respectively. These trajectories with different types of marine influenced air masses occurred for a very short period of time (3 % and 4 % of the total measurements period, respectively). The first one, TS-C1, started from the Atlantic Ocean near Spain and is associated with a more continental influence, which is why organic mass concentration and contribution were higher than inorganics. However, The LO-OOA contribution of this cluster was the highest during this time period (26 %) due to the aging processes of primary organic aerosols especially CCOA, which had a
 - 22

718 maximum mass concentration (0.31 µg/m³ and mass fraction of 7 %, respectively). While the second one, TS-C2, was almost

719 a pure marine cluster, coming from the Norwegian Sea. In opposition to TS-C1, PM was dominated by inorganics in TS-C2,

720 with a high mass concentration of nitrate $(1.35 \ \mu g/m^3)$ representing the aging effect due to the long-time transfer over the

721 continents.

722 **3.4.3 Summer**

During the summer season, the different clusters showed strong changes in both chemical compositions and total mass concentrations. Cluster WS-ST was identified as the local air mass with a mean value of 8.97 μ g/m³, which occurred for a short period, 6 % of the measurement. However, this cluster contained a low POA mass concentration but a maximum contribution of MO-OOA (32 %), assuming important regional photochemical roles of SOA particles with higher temperatures (Fig. <u>44S4</u>) and enhanced solar radiation (Petit et al., 2015).

728 Air masses WS-A1 and WS-A2 were two different types of anticyclonic air masses with different directions and different 729 mean PM mass concentrations. Cluster WS-A1, known as the highest mass concentration during summer time, $(16.95 \,\mu g/m^3)$ 730 and contribution of 11 % of the measurement period) was the continental air mass which was coming from Eastern Europe 731 during the anticyclonic flow (starting from Belarus, crossing Poland and the Czech Republic). This air mass included maximum 732 inorganics and organics especially CCOA mass concentration $(1.28 \,\mu g/m^3)$ during summer time, which can explain the existing 733 higher CCOA during summer, and showed the role of long-range transported emissions in the summer season. However, WS-734 A2 air mass, with a mean value of 9.48 µg/m³ was a marine-influenced air masse and was coming from the North Sea, which 735 only occurred for a short period (6 % of the total measurement period).

Moreover, two cyclonic air masses, WS-C1 and WS-C2, were also identified as two different marine clusters. These trajectories did not occur very often, only 5 % and 3 % of the total measurement period, respectively. The starting point of WS-C1 with a mean value of 8.41 μ g/m³ was the Celtic Sea, but in the following time, it predominantly passed over continental areas (France and southern Germany), which means it could be aged and the result can be shown in the high mass concentration of nitrate and sulphate in this cluster (1.63 μ g/m³ and 1.86 μ g/m³, respectively). Finally, the starting point of WS-C2 with a mean value of 4.46 μ g/m3, was near Iceland, with the lowest PM mass concentration during summer. However, it showed the highest sulphate contribution (27 %) at this time which could be associated with aged marine air mass like other marine air masses.

743 **3.4.4 Cluster seasonality**

A parallel comparison can be made between the winter and summer clusters. Clusters CS-A1 and WS-A1 both show the highest POA contribution dominated by coal combustion, which emphasizes that the origin of this source could be associated with the transport of the coal power plants emissions from Eastern Europe (e.g. Eastern part of Germany, Poland, Czech Republic and further countries located in the East). <u>They-These clusters</u> were not only affected by the winter air quality but also the summer air quality.

- 749 Clusters CS-ST and WS-ST, which were known as local air masses, showed the seasonal effect on the chemical component.
- 750 First, the volatility of ammonium nitrate at higher summer temperatures could explain their lower value in summer. Then,
- 751 atmospheric photochemical oxidation processes affected the sulphate locally formed emission in summer, which its highest
- value over inorganic components during summer can be confirmed. Not surprisingly, due to the residential heating effect, POA
- 753 mass concentration was very high during winter; however, freshly formed SOA originating from biomass and coal emissions
- 754 can explain the higher LO-OOA mass concentration in winter.
- 755 During the whole period, some marine air masses with cyclonic and anticyclonic flow showed the important roles of aged
- 756 marine air masses over the measurement site: a) clusters CS-A2 and WS-A2 with anticyclonic pattern starting from the North
- 757 and/or Norwegian Sea, and b) CS-C2a, WS-C1, and TS-C1 starting from the Celtic Sea near Spain, and also CS-C2b and WS-
- C2 starting from Iceland, all with cyclonic patterns contain nitrate and sulphate during the transferring over the continental
 areas in different seasons.

760 4 Conclusion

- 761 The chemical compositions of non-refractory fine aerosol (NR-PM₁) at the German rural-background observatory Melpitz 762 were investigated in this study over a one-year period between September 2016 and August 2017. Overall, the averaged total PM_1 mass concentration is 10.49 μ g/m³ and follows a clear seasonal pattern, with the highest mass concentration during winter 763 764 (15.95 µg/m^3) and the lowest mass concentration during summer time (6.24 µg/m³). The organic aerosol was the most 765 significant component, accounting for 46% of total PM₁ and showing significant seasonal dependency (39% in winter to 58%) % in summer). It was followed by sulphate (15 % and 20 %) and nitrate (24 % and 11 %). For OA source apportionment, PMF 766 767 in a rolling fashion has been applied using the SoFi Pro, which provided the decomposition of time-dependent factor profiles 768 that were able to better capture the variability of OA sources across seasons in comparison with the conventional seasonal 769 PMF. The final solution enabled the identification of five factors throughout the one-year measurements of OA; HOA, BBOA, 770 CCOA, LO-OOA, and MO-OOA. Using the correlation between HOA, BBOA, and CCOA with eBC-PM₁, a multilinear 771 regression approach was applied to perform the source apportionment of eBC-PM₁.
- 772 Generally, in Melpitz, HOA as a minor source of OA (6 % of the contribution of total organic mass) and eBC-PM₁ (8 % of the 773 total eBC-PM₁) was associated with: a) low traffic emissions, b) household heating in winter, and c) the central heating for hot 774 water production for all the seasons which uses multiple fuel types in the Melpitz area. BBOA representing 7.9 % of the 775 contribution of total organic mass and 37 % of the total eBC-PM₁, showed a seasonal effect, emphasizing the impact of house 776 heating during winter. Similar to HOA, the presence of BBOA during summer was due to central heating which uses multiple 777 fuel types in the Melpitz area. The most dominant anthropogenic source was associated with CCOA with a 15.4 % contribution 778 of total organic mass and 55 % of the total eBC-PM₁ with the highest mass concentration and contribution of PM during winter 779 rather than summer. Although a certain fraction of CCOA could be linked to surrounding domestic heating (van Pinxteren et

780 <u>al., 2020</u>), it is rather associated with power plant emissions and long-range transport all year round which is supported by 781 <u>cluster and back-trajectory analysis</u>. LO-OOA and MO-OOA referred to oxidized oxygenated organic aerosol (32.4 % and 782 <u>38.4 % of the contribution of total organic mass</u>, respectively), were identified as a secondary organic aerosol with the highest 783 mass concentration during the cold months and the lowest mass concentration during the warm months. LO-OOA mass 784 concentration decreased during the day due to dilution, and the evaporation process resulted in aging into MO-OOA.

785 A combination of the NWR model and cluster analysis was used to better understand the origin of the aerosol reaching the 786 station. Overall, Melpitz is influenced by fifteen types of air masses, such as long-range continental, marine, and surrounding 787 emissions. During winter and summer time, easterly continental air masses, CS-A1 and WS-A1 with an anticyclonic pattern come from Eastern Europe and showed a significant particle mass concentration, especially high POA (and CCOA) mass 788 789 concentration at the measurement site. Marine clusters, mostly coming from the south/west/north side with aged marine air 790 masses including nitrate and sulphate, also have important roles in the PM mass concentration at the Melpitz site over the entire period (winter: CS-A2, CS-C2b, and CS-C2a, transition: TS-C, TS-A2 and TS-C2, and summer: WS-Ca, WS-C2, and 791 WS-A2). However, the surrounding emissions are recognized as another important source of emissions which include high 792 793 organic and inorganic components during winter and summer (CS-ST and WS-ST, respectively). Within this study, the change 794 in chemical compositions of non-refractory fine acrosol (NR-PM₁) at the German rural-background observatory Melpitz was 795 investigated during a one year period between September 2016 and August 2017, by applying PMF in a rolling fashion with 796 14 days window length and a 1 day shift using the SoFi Pro. This method provided the decomposition of time-dependent factor 797 profiles that were able to better capture the variability of OA sources across seasons, in particular for LO OOA. Overall, the 798 averaged total PM₁ mass concentration is 10.47 µg/m³ and follows a clear seasonal pattern, with the highest mass concentration 799 during winter (15.95 µg/m³) and lowest mass concentration during summer time (6.24 µg/m³). The organic aerosol was the 800 major component accounting for 46 % of total PM1 and showing a strong seasonal variability ranging from 39 % (in winter) 801 to 58 % (in summer). It was followed by sulphate (15 % and 20 %) and nitrate (24 % and 11 %). The final solution of the PMF 802 rolling approach for OA source apportionment enabled the identification of five factors throughout the one-year measurements 803 of OA; HOA, BBOA, CCOA, LO OOA, and MO OOA.

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806 Generally, in Melpitz, HOA as a minor source of OA (6 % of the contribution of total organic mass) was associated with: a) 807 low traffic emissions, b) household heating in winter, and c) the central heating for hot water production for all the seasons 808 which showed a small increasing mass concentration pattern toward cold months (winter and summer: $0.36 \ 0.23 \ \mu g/m^3$). The 809 HOA night time mass concentration was not affected by the seasons, which indicates the presence of a continuous emission 810 source. Biomass burning emissions (BBOA) representing 7.9 % of the contribution of total organic mass showed a seasonal 811 effect, emphasizing the impact of house heating during winter (winter and summer: 23 % and 8.7 %). This highest mass 812 concentration during the winter time showed the descending pattern from night time to day time due to domestic heating 813 activities and the planetary boundary layer effect; however similar to HOA, the presence of BBOA during summer was due to 814 central heating which uses multiple fuel types in the Melpitz area. The most dominant anthropogenic source was associated 815 with coal combustion (CCOA) with a 15.4 % contribution of total organic mass and 55 % of eBC with the highest mass 816 concentration and contribution of PM during winter rather than summer (1.58 0.30 µg/m³). Although a certain fraction of 817 CCOA could be linked to surrounding domestic heating (van Pinxteren et al., 2022), it is rather associated with power plant 818 emissions and long range transport all year round. Using the correlation between HOA, BBOA, and CCOA with eBC, a 819 multilinear regression approach was applied to perform the source apportionment of eBC. This analysis highlighted eBC 820 contribution related to the source of HOA (8 % of the total eBC), BBOA (37 % of the total eBC), and CCOA (55 % of the 821 total eBC), which showed the CCOA as the largest source of eBC during the measurement period. Moreover, from the seasonal 822 source apportionment, CCOA presented the largest fraction (56 % of the total eBC) during winter, while the highest fraction 823 is attributed to BBOA for summer time (69% of the total eBC). LO OOA and MO OOA referred to oxidized oxygenated 824 organic aerosol (32.4 % and 38.4 % of the contribution of total organic mass, respectively), were identified as a secondary 825 organic aerosol with the highest mass concentration during the cold months (fall: 2.13 µg/m³ and winter: 2.25 µg/m³. 826 respectively) and the lowest mass concentration during the warm months (spring: 1.24 µg/m³ and summer: 1.44 µg/m³, 827 respectively). LO OOA mass concentration decreased during the day due to dilution, and the evaporation process resulted in 828 aging into MO OOA.

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830 A combination of pollution wind rose and cluster analysis was used to better understand the origin of the aerosol reaching the 831 station. Overall, Melpitz is influenced by fifteen types of air masses, such as long range continental, marine, and surrounding 832 emissions. During winter and summer time, easterly continental air masses, CS A1 and WS A1 with an anticyclonic pattern 833 come from Eastern Europe and showed a significant particle mass concentration, especially high POA (and CCOA) mass 834 concentration at the measurement site. Marine clusters, mostly coming from the south/west/north side with aged marine air 835 masses including nitrate and sulphate, also have important roles in the PM mass concentration at the Melpitz site over the 836 entire period (winter: CS A2, CS C2b, and CS C2a, transition: TS C, TS A2 and TS C2, and summer: WS Ca, WS C2, and 837 WS A2). However, the surrounding emissions are recognized as another important source of emissions which include high 838 organic and inorganic components during winter and summer (CS-ST and WS-ST, respectively).

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Our results emphasize the importance of the long-range transported emissions of coal combustion related aerosol particles regardless of the season, which supports that the main CCOA source is related to coal power plants emissions. However, coal power plants emissions not only affect the surrounding air quality but can also be transported over long distances. It is important to note that the overall coal combustion mass concentration presented here can certainly be underestimated since the identified CCOA factor is associated with freshly emitted organic aerosol and no factor associated with potential aged coal combustion was identified. Because coal still is an important energy source in the European energy mix (68.4 % of all energy in the EU was produced from coal, crude oil, and natural gas, Energy Statistics - an Overview - Statistics Explained, 2022) as well as on 847 a global scale and also that it still will be in used for the coming decades (until 2040, Europe's Coal Exit - Europe Beyond

Coal : Europe Beyond Coal, 2022), further research<u>es</u> should be done on the identification of coal emissions across Europe in
 order to better understand its atmospheric aging processes.

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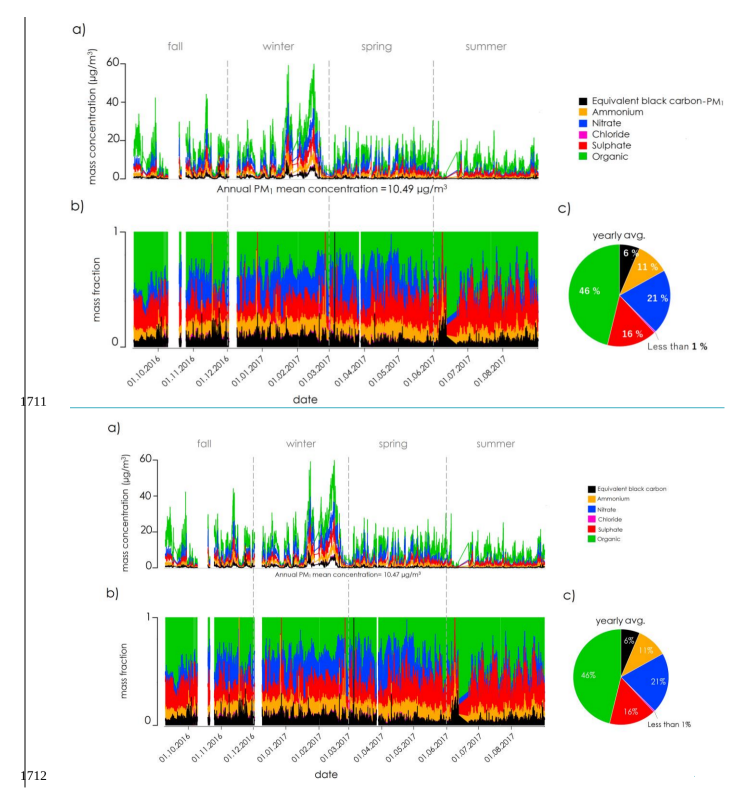
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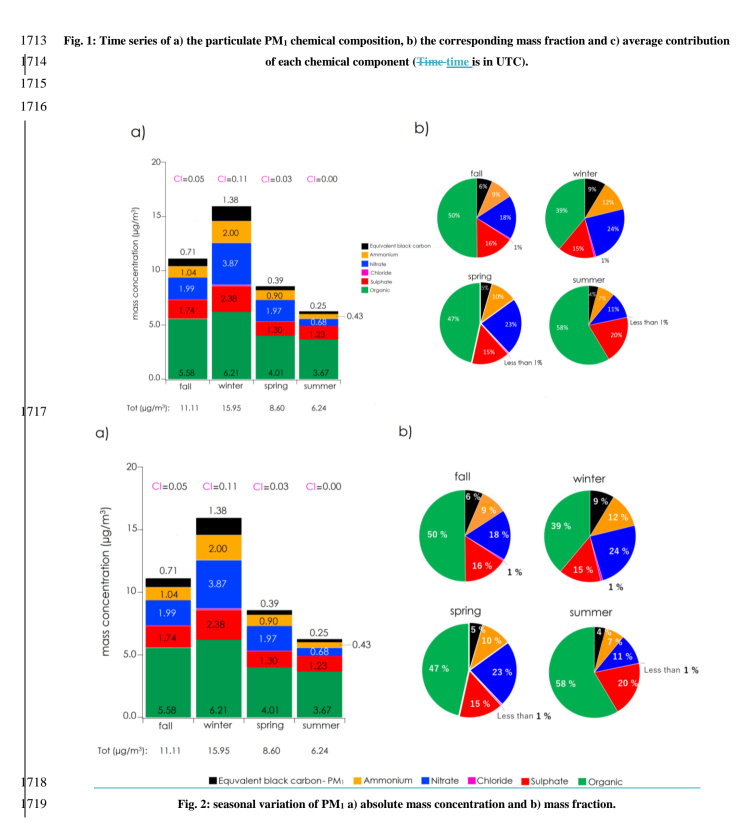
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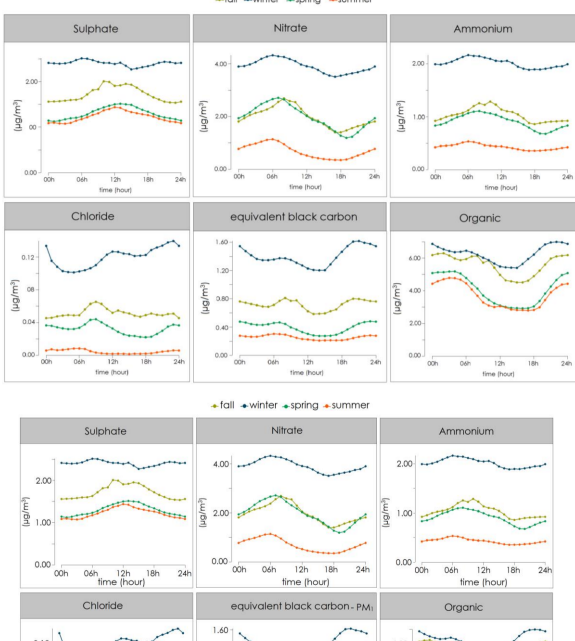
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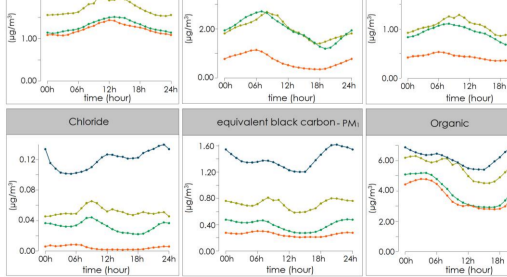






→ fall → winter → spring → summer





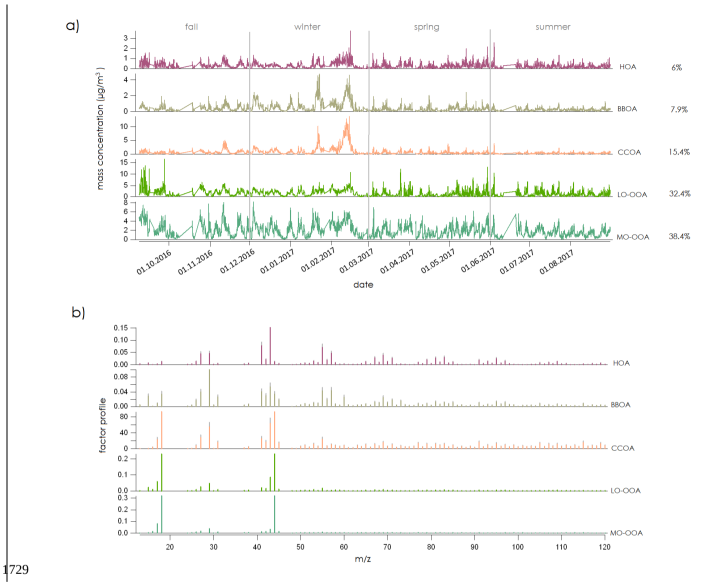
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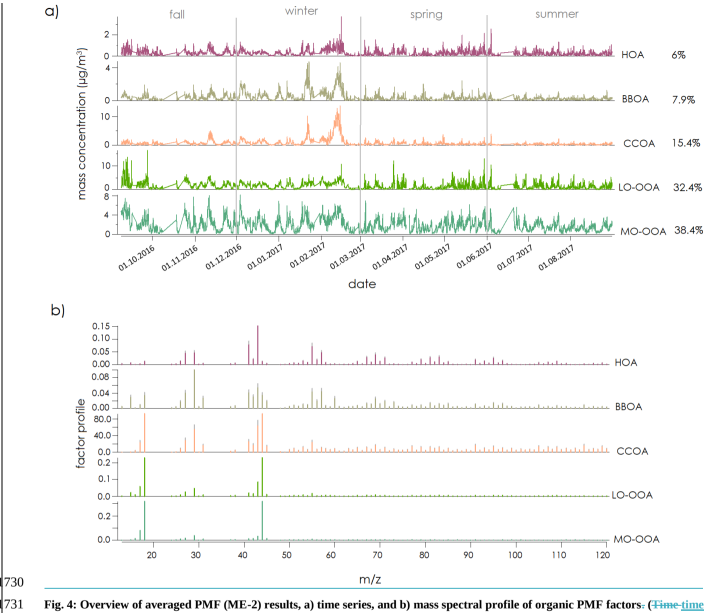
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Table. 1: Seasonal/yearly mass concentration of each ACSM species, each PMF factors, contribution of the different POA-PMF-

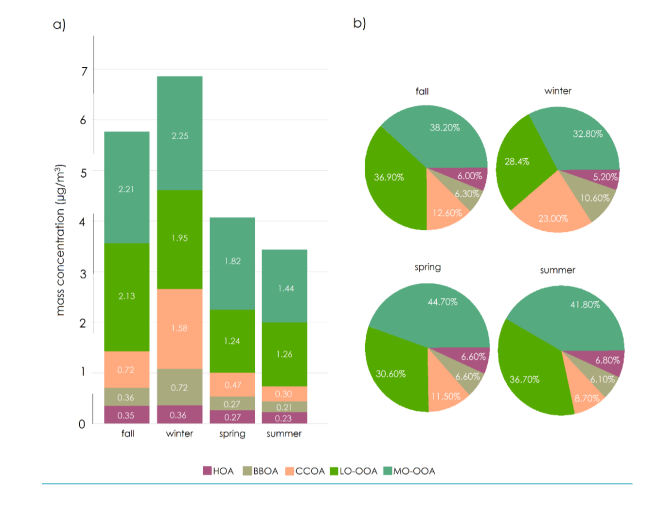
<u>eBC-PM₁eBC</u>	, and correlation	of each factors	with related	l species; PM1.
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Spe	ecies/ Factors	Fall	Winter	Spring	Summer	Yearly
	Org	5.58	6.21	4.01	3.67	4.84
ACSM	SO4 ²⁻	1.74	2.38	1.30	1.23	1.67
(µg/m ³)	NO ₃ -	1.99	3.87	1.97	0.68	2.16
(µg/m)	\mathbf{NH}_{4}^{+}	1.04	2.00	0.90	0.43	1.11
	Cl	0.05	0.11	0.03	0.00	0.05
MAAP						
(µg/m ³)	<u>eBC -PM₁eBC</u>	0.71	1.38	0.39	0.25	0.66
	НОА	0.35	0.36	0.27	0.23	0.30
	BBOA	0.36	0.72	0.27	0.23	0.30
PMF	CCOA	0.30	1.58	0.27	0.30	0.39
$(\mu g/m^3)$	LO-OOA	2.13	1.95	1.24	1.26	1.62
	MO-OOA	2.13	2.25	1.24	1.20	1.02
	eBC-PM ₁ eBC-HOA	0.16	0.19	0.03	0.04	0.05
<u>eBC-PM₁eBC</u>						
(µg/m ³)	<u>eBC-PM</u> _l eBC-BBOA	0.34	0.38	0.17	0.15	0.25
	eBC-PM ₁ eBC-CCOA	0.23	0.74	0.16	0.02	0.37
<u>eBC-PM₁eBC</u>	<u>eBC-PM₁eBC-HOA</u>	22	15	9	18	8
(%)	eBC-PM1eBC-BBOA	47	29	47	69	37
	eBC-PM ₁ eBC-CCOA	31	56	44	13	55
	HOA/ <u>eBC-PM</u> ₁ eBC	0.49	0.52	0.34	0.24	0.33
	HOA/NO _x	0.23	0.12	0.32	0.23	0.17
	BBOA/Levo.	0.19	0.59	0.09	0.07	0.54
Correlation	BBOA/ <u>eBC-PM1</u> eBC	0.62	0.81	0.48	0.42	0.77
(R ²)	CCOA/ <u>eBC-PM1</u> eBC	0.65	0.85	0.49	0.30	0.82
(K)	CCOA/Cl ⁻	0.40	0.41	0.18	0.15	0.46
	LO-OOA/ NO3	0.22	0.59	0.01	0.12	0.19
	LO-OOA/ SO4 ²	0.36	0.55	0.00	0.02	0.23
	MO-OOA / SO4 ²	0.58	0.47	0.34	0.42	0.44
	MO-OOA/ NO3 ⁻	0.24	0.47	0.16	0.24	0.31





is in UTC).



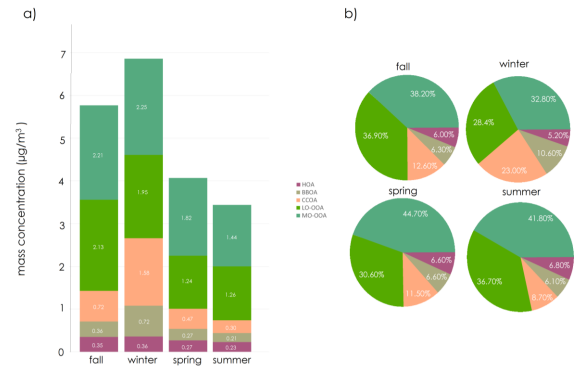
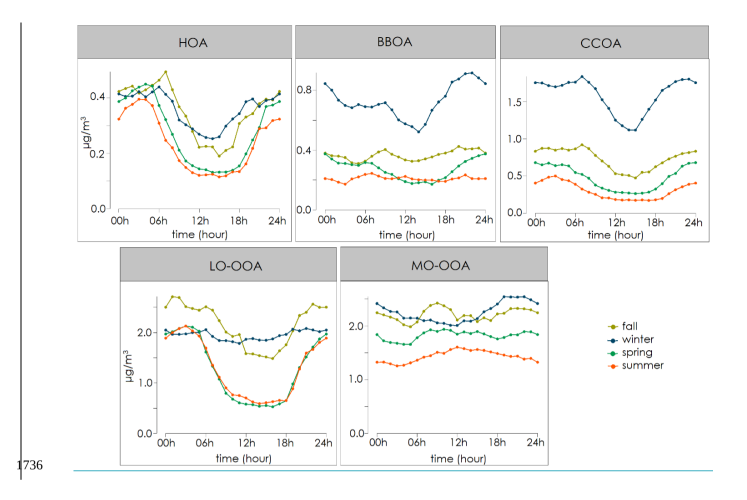
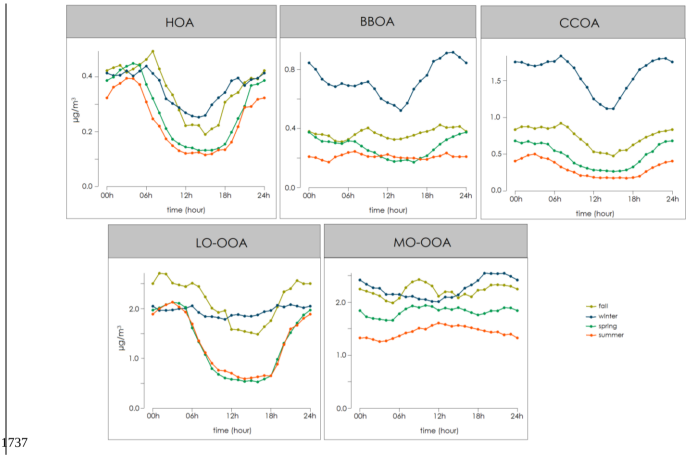


Fig. 5: Seasonal variation of a) mass concentration, b) mass fraction of PMF source factors.





1738 Fig. 6: Seasonal diurnal cycle (hourly averages) of the organic components HOA, BBOA, CCOA, LO-OOA and MO-OOA in UTC

time.

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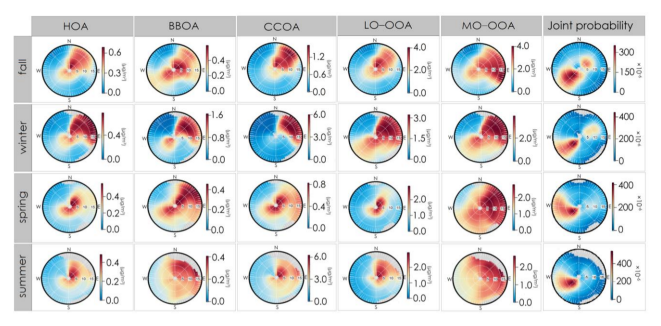
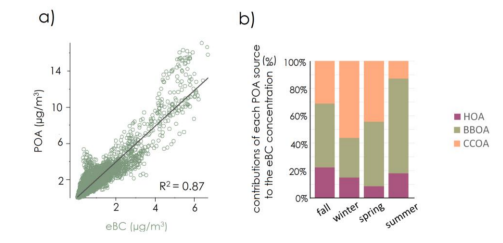
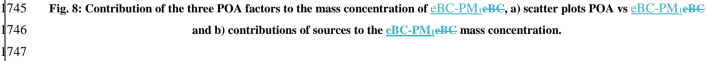
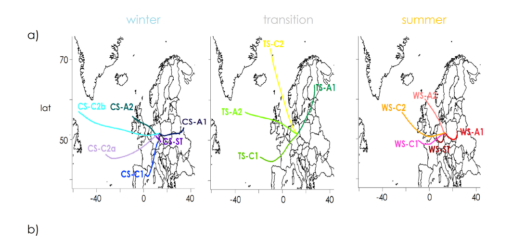
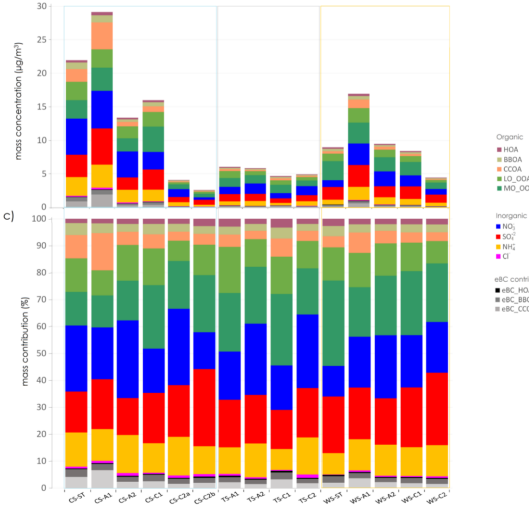


Fig. 7: Seasonal wind roses and NWR plots for the different PMF factors (in µg/m³).











NO₃
 SO₄⁻²
 NH₄⁺
 Cl⁻

eBC contribution ■ eBC_HOA ■ eBC_BBOA ■ eBC_CCOA

1749

air mass classification

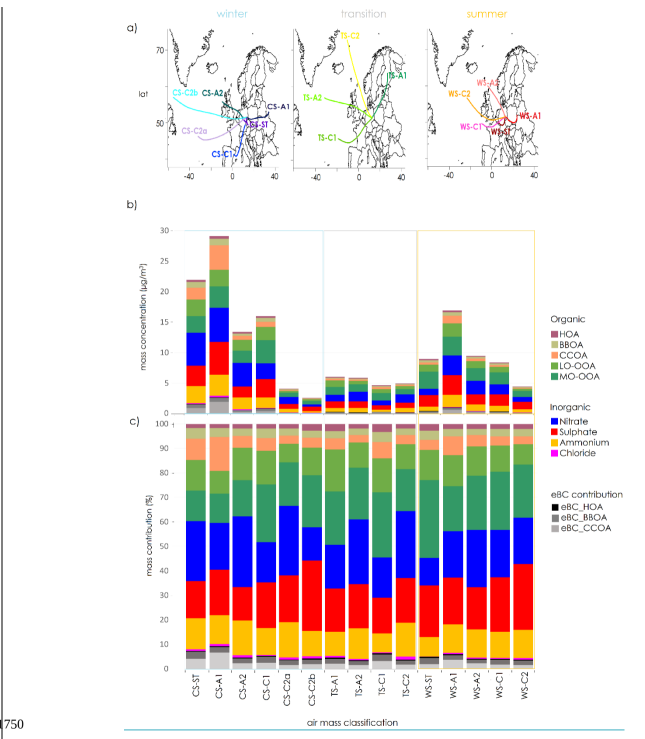




Table. 2: Main statistical details of the 15 air mass types for total PM₁

(CS=Cold Season, WS=Warm Season, ST=Stagnant, A=Anticyclonic, C=Cyclonic).

Main season	Airmass type	Wind direction	Vorticity	Frequenc y (%)	Total mean (µg/m³)
	CS-ST	Stagnating	Anticyclonic	14	21.95
	CS-A1	East	Anticyclonic	18	29.14
TT [*]	CS-A2	West	Anticyclonic	8	13.39
Winter	CS-C1	South	Cyclonic	10	15.99
	CS-C2a	South West	Cyclonic	3	04.09
	CS-C2b	West	Cyclonic	2	02.60
	TS-A1	North East	Anticyclonic	4	06.06
Transition	TS-A2	West	Anticyclonic	4	05.86
(Spring/ Fall)	TS-C1	South West	Cyclonic	3	04.69
	TS-C2	North West	Cyclonic	4	04.94
	WS-ST	Stagnating	Anticyclonic	6	08.97
	WS-A1	South East	Anticyclonic	11	16.95
Summer	WS-A2	North West	Anticyclonic	6	09.48
	WS-C1	West	Cyclonic	5	08.41
	WS-C2	West	Cyclonic	3	04.46