1 Long-range transported pollution from the Middle East and its 2 impact on carbonaceous aerosol sources over Cyprus. Ambient 3 carbonaceous aerosol levels in Cyprus and the role of pollution transport from the Middle East. 4

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Abstract. The geographical origin and source apportionment of submicron carbonaceous aerosols (organic aerosols, OA, and 23 black carbon, BC) have been investigated here for the first time by means of deploying high-time resolution measurements at 24 25 an urban background site of Nicosia, the capital city of Cyprus, in the Eastern Mediterranean. This study covers a half-year period, encompassing both the cold and warm periods with continuous observations of the physical and chemical properties 26 27 of PM1 performed with an Aerosol Chemical Speciation monitor (ACSM), an Aethalometer, accompanied by a suite of various ancillary off and on-line measurements. Carbonaceous aerosols were dominant during both seasons (cold and warm periods), 28 29 with a respective contribution of 57% and 48% to PM₁, respectively, and exhibited recurrent intense night-time peaks (>20-30 30 µg m⁻³) during the cold period associated with local domestic heating. The Ffindings of this study show that high 31 concentrations of sulfate (close to 3 µg m⁻³) were continuously recorded, standing among the highest ever reported for Europe 32 and originating from the Middle East region. 33 Source apportionment of the OA and BC fractions was performed using the Positive Matrix Factorization (PMF) approach and 34 the combination of two models (aethalometer model and multilinear regression), respectively. Our study revealed elevated hydrocarbon-like organic aerosol (HOA) concentrations in Nicosia (among the highest reported for a European urban 35

background site), originating from a mixture of local and regional fossil-fuel combustion sources. Although air masses from 36 37

the Middle East had a low occurrence and were observed mostly during the cold period, they were shown to strongly affect 38 the mean concentrations levels of BC and OA in Nicosia during both seasons. Overall, the present study brings to our attention

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the need to further characterize primary and secondary carbonaceous aerosols in the Middle East; an undersampled region 40 characterized by continuously increasing fossil fuel (oil and gas) emissions and extreme environmental conditions, which can

contribute to photochemical aging. 41

42 1. Introduction

43 At the crossroads of three continents (Europe, Africa, Asia), the Eastern Mediterranean and Middle East (EMME) region faces 44 many challenges, such as rapid population growth - with its currently 400 million inhabitants - as well as political and socioeconomic instabilities. Environmental conditions in the region are exceptional, with the two largest deserts worldwide (Sahara 45 46 and, Arabian) being among the most water scarce ecosystems on the planet (Terink et al., 2013). Climate change in this region 47 is extraordinarily rapid; summer temperatures, in particular, are increasing by more than twice the global mean rate (Lelieveld 48 et al., 2014), with significant impacts, especially in urban areas (Mouzourides et al., 2015). While aerosol mass loadings over 49 the EMME are dominated by desert dust, concentrations of fine particles due to anthropogenic emissions are also high (Basart 50 et al., 2009) and will likely increase with continued population growth (Pozzer et al., 2012), making anthropogenic pollution in the area a leading health risk and an important climate forcer (Osipov et al., 2022). 51 52 Based on modelling studies, it has been also concluded that the EMME is characterized by highly favourable conditions for photochemical smog and ozone (O₃) formation leading to air quality standards being drastically exceeded (Lelieveld et al., 53 54 2014; Zanis et al., 2014). These enhanced concentrations of fine particulates and ozone have major human health implications. contributing to premature mortality (Giannadaki- et al., 2014; Lelieveld et al., 2015), which may be further exacerbated by the 55 56 effects of heatwaves occurring during summer within the EMME region (Zittis et al., 2022). 57 Although data derived from satellite observations of NO₂ and SO₂ has revealed strong air pollution trends in the Middle East 58 since 2010 (Lelieveld et al., 2015a), many pollution sources are still missing in emission inventories (Mclinden et al., 2016). 59 Thus, there is a current lack of a regional approach to characterize air pollution, with in-situ observation being insufficient, 60 unavailable, or of low quality (Kadygrov et al., 2015; Ricaud et al., 2018; Paris et al., 2021), limiting the possibility to reduce 61 uncertainties in regional emission inventories and implement efficient abatement strategies. 62 Significant efforts have been put forward in recent years to characterize the atmospheric composition in-situ over Cyprus, a 63 central location of the EMME region (e.g., Kleanthous et al., 2014; Debevec et al., 2017 and 2018; Pikridas et al., 2018; Dada et al., 2020; Baalbaki et al., 2021; Vrekoussis et al., 2022). In-situ ground-based PM observations have clearly shown that 64 65 contributions of dust to PM_{10} over Cyprus are among the highest for the entire Mediterranean basin (Querol et al., 2009; Pey 66 et al., 2013; Kleanthous et al., 2014; Pikridas et al., 2018; Achilleos et al., 2020), during dust storm events, leading to increased 67 hospitalization, particularly attributed to cardiovascular-related diseases (Middleton et al., 2008; Tsangari et al., 2016) and 68 short-term effects associated with daily mortality (Neophytou et al., 2013). These high levels of regional particulate matter are responsible for exceedances in PM10 EU limits in major Cypriot cities (Querol et al., 2009). Past studies on PM trends and 69 70 sources highlighted the important contribution of local (urban) emissions to PM₁₀ (Achilleos et al., 2014; Pikridas et al., 2018) 71 but also showed a predominant regional pattern for PM2.5 with a major contribution of sulfphur-rich sources (Achilleos et al., 72 2016). Based on 17 years of continuous observations of reactive gases in Cyprus, Vrekoussis et al., (2022) further confirmed

the major contribution of long-range transport (incl. Middle East) in the observed concentration levels of carbon monoxide
 (CO) and sulfphur dioxide (SO₂), two tracers of combustion sources.

75 Those studies have highlighted the unique location of Cyprus as a receptor site of major regional pollution hotspots, making 76 the island one of the most polluted EU member states in terms of PM and O3 concentrations; the only one impacted by long-77 range transport of poorly-regulated air pollutants originating from Middle East countries. However, still few studies are 78 currently available to assess the contribution of regional anthropogenic emissions to PM levels in Cyprus. The filter-based 79 chemical speciation study reported by Achilleos et al., (2016) is currently the most exhaustive one and was based on 24-h 80 integrated ($PM_{2.5}$ and PM_{10}) filter samples collected every 3 days for a period of one year (2012) in four cities in Cyprus. This 81 study concluded that Cypriot cities, like many others in Europe, are characterized by a major contribution of regional sulfphate 82 and local (urban) emissions from traffic and domestic heating biomass burning.

 $83 \quad Here with, a \ detailed \ description \ of \ submicron \ (<1 \mu m, PM_1) \ chemical \ composition \ and \ the \ further \ source \ apportionment \ of \ BC$

and OA is presented for the first time in Cyprus. State-of-the-art on-line instrumentation (e.g., Q-ACSM, Aethalometer) were

was deployed for the investigation to investigate of the temporal variability of aerosol composition at a location representative 85 of the urban background pollution in the capital city of Nicosia. Source apportionment of submicron organic aerosols was 86 87 performed using the organic fragments of the ACSM and Positive Matrix Factorization (PMF). The consistency of these results was assessed against the chemical analysis of parallel filter samples and on-line measurements of external tracers. This study 88 89 was extended to a 6-month duration in order to cover the two main seasons of the semi-arid Eastern Mediterranean climate 90 (short, mild and wet winter vs. long, hot and dry summer), offering a comprehensive understanding of the daily and monthly 91 variability of local and regional sources of carbonaceous aerosols. Cold and warm periods were compared to highlight the 92 complexity of local (combustion) sources and the importance of regional ones. These results were further processed to apportion Black Carbon sources in Nicosia with emphasis on local versus regional contribution. 93

94 2. Material and Methods

95 2.1 Sampling site

96 *Cyprus*: Cyprus is the third largest island in the Mediterranean Sea, extending approximately 240km long from east-to-west
97 and 100km wide. The closest countries and their distance from the capital city of Nicosia are respectively Turkey (110km),
98 Syria (250km), Lebanon (250km), Israel (300km), Egypt (400km), Jordan (430 km), and Greece (900 km from the Greek
99 mainland), (Fig. 1a).

100 The population of Cyprus (approximatelyca, 1 million inhabitants) is rather small compared to its neighbouring countries and 101 the rapidly growing (overall 400 million) population of the region (Lelieveld et al., 2013). The main urban areas of the island 102 shown in Fig. 1b, are those of Nicosia (c.a. 245,000 inhabitants), Limassol (c.a. 150,000 inhabitants), Larnaca (c.a. 50,000 103 inhabitants) and Paphos (c.a. 35,000 inhabitants). Cyprus has a Mediterranean and semi-arid climate with two main seasons: 104 a mild cold season (from December to March) and a hot warm season lasting about eight months (from April till-to November). 105 Rain occurs mainly in the cold season, with the warm one being extremely dry (i.e., almost no rain between May and 106 September)(Michaelides et al., 2018). 107 Nicosia: Nicosia is the largest city onf the island and the southeasternmost of the European Union Member States' capitals.

Nicosia is currently partitioned in two, with a buffer zone in-between under the control of the United Nations; the southern part being the capital of the Republic of Cyprus. The northern part of Nicosia (and <u>the</u>_northern part of the island) is not controlled by the government of the republic of Cyprus (Resolution 550, UN security council, 1984) (Fig. 1c). Geographically, Nicosia is located in the centre of the island, within the Mesaoria plain, 150 m above sea level (asl), which is delimited on its northern and southern edges by two mountain ranges; the Kyrenia Range culminating at 1,024 m asl, and the Troodos Mountains culminating at 1,952 m asl, respectively. This topography channels winds within a more or less west-east corridor (Fig. S5), feeding the city of Nicosia with long-range transported air masses from Europe, Africa, or the Middle East.

115 Measurements were performed at the Cyprus Atmospheric Observatory's Nicosia station (CAO-NIC) located at the Cyprus

116 Institute premises (Athalassa Campus; 174 m asl; 35.14N, 33.38E; Fig. 1c). The measurement site is considered as an urban

background site, located within a low population density residential area with no significant local pollution hotspots in its vicinity (i.e., no dense road traffic, industry, commercial centers, restaurants, etc.) and next to the Athalassa Forestry Park.

The period and duration of measurements presented here (07 December 2018 - 31 May 2019) were chosen to i) capture weather

120 conditions, atmospheric dynamics, and long-range pattern of the two main seasons, ii) investigate the contribution of domestic

121 heating emissions in winter, and iii) assess the potential increasing contribution of photochemical produced secondary aerosols

122 during the start of the dry and warm season. Local time (LT) in Cyprus is given as Eastern European Standard Time (EET)

123 (UTC+02:00 in winter and UTC+03:00 during the summer).



(a)



(b)

Figure 1: (a) Geographic location of the island of Cyprus and its closest Northern African and Middle Eastern neighbouring countries. (b) Location of the main cities of the Republic of Cyprus. Maps a,b were created by QGIS software v.3.26.3 utilizing the Natural Earth data (https://dgis.org). (c) Satellite view of the Nicosia agglomeration (grey area). The buffer zone dividing the island and the city is marked with red stripes; the location of the measurement site (CAO-NIC; The Cyprus Institute, Athalassa campus) is noted in red. (© OpenStreetMap contributors 2022. Distributed under the Open Data Commons Open Database License (ODbL)

(c)

130 2.2 On-line Aerosol Instrumentation

On-line aerosol instrumentation has been operated following the Standard Operating Procedures defined by ACTRIS
 (<u>https://www.actris.eu</u>), the European Research Infrastructure on Aerosols, Clouds, and Trace Gases_ACTRIS

133 (https://www.actris.eu), and Cost COLOSSAL (CA16109, 2021).

134 Non-refractory submicron (NR-PM1) aerosol chemical composition, i.e. organics, sulfphate, nitrate, ammonium and chloride, was continuously monitored using a Quadrupole ACSM (Aerosol Chemical Speciation Monitor; Aerodyne Research Inc.) at 135 136 a 30-min time resolution (Ng et al. 2011a). The instrument, along with a scanning mobility particle sizer (SMPS, described below), sampled through a sharp cut cyclone operated at 4 L min⁻¹ (SCC 1.197, BGI Inc., USA), and was equipped with a PM₁ 137 138 aerodynamic lens, yielding an aerosol cut-off diameter of approximately 1.3µm. A nafion dryer was installed upstream, keeping sample RH below 30%. Data were retrieved using ACSM local v.1.6.0.3, implemented within Igor Pro (v. 6.37, 139 140 Wavemetrics Inc., USA). The ACSM is designed and built around similar technology as the aerosol mass spectrometer (Jayne et al., 2000), where an aerodynamic particle focusing lens is combined with particle flash vaporization in high vacuum on the 141 142 surface of a standard tungsten vaporizer heated at 600 °C, followed by electron impact ionization, separation and final detection 143 of the resulting ions using a quadrupole mass spectrometerat 600, °C, electron impact ionization, separation and final detection 144 of the resulting ions using a quadrupole mass spectrometer. Mass concentrations are ealculated corrected for incomplete 145 detection due to particle bounce, using thea chemical composition-dependent collection efficiency (CDCE) (Middlebrook et 146 al., 2012). The determined parameters, response factor (RF) and relative ionization efficiency (RIE) are reported in table S2. 147

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Black carbon (BC) measurements were conducted using an AE-33 7-wavelength aethalometer (AE-33 Magee Scientific, US) 148 149 at a 1-min time resolution. The aethalometer sampled ambient aerosol through a PM2.5 aerosol inlet (SCC 1.829, BGI Inc., 150 USA) at a flow rate of 5 L min⁻¹, after passing through a nation dryer. The instrument internally corrected the filter loading 151 effect in real-time, while a fixed value (C₀=1.39) was applied to compensate for the multi-scattering effect (Drinovec et al., 152 2015). BC was apportioned to source specific components, namely BC_{ff} related to fossil fuel combustion and BCwb related to 153 wood burning, by applying the "aethalometer model" (Sandradewi et al., 2008) on the 470 - 950 nm wavelength pair. They 154 using the instrument's default values for fossil fuel combustion and wood-burning aerosol Absorption Ångström Exponent, 155 AAE_{if}=1 and AAE_{wb}=2, respectively were selected after performing a sensitivity analysis on the AAE values (Supplement

156 Section 3).-

157 2.3 Ancillary measurements

158 SMPS: Particle number size distributions were monitored using a scanning mobility particle sizer (SMPS) consisting of an 159 electrostatic classifier (model 3080, TSI Inc., USA) coupled with a condensation particle counter (CPC; model 3070, TSI Inc. 160 USA) operating at a 5-min time resolution and at a 1 L min⁻¹ sample flow rate, measuring particles with <u>a</u> diameter ranging 161 from 9 to 700 nm. Ambient aerosols were drawn through a nafion dryer, and placed upstream, keeping sample RH below 30 162 %. Volume concentrations of assumed spherical particles derived by the SMPS were converted into mass concentrations using a variable density calculated by the methodology described in Bougiatioti et al. (2014). The respective mass fractions time 163 164 series of chemical species were calculated based on the ACSM measurements. A density value of 1.77 g cm⁻³ was used for 165 ammonium sulfphate, and 1.35 g cm⁻³ for organics (Florou et al., 2017; Lee et al., 2010), the two dominant compounds of PM₁ 166 in Nicosia as detailed further below.

Filter sampling: Co-located 24h PM_{2.5} samples were collected on quartz fiber filters (Tissuquartz, 47mm diameter, Pall) using
 a low volume sampler (Leckel SEQ47/50) operating at a flowrate of 2.3 m³ h⁻¹. The filter samples were analysed for i) organic
 and elemental carbon using an OC/EC Lab Instrument (Sunset Laboratory Inc., OR, USA) implementing the EUSAAR II
 protocol (Cavalli and Putaud, 2008), ii) carbohydrates, including levoglucosan, mannosan, galactosan, using an Ion
 Chromatography Pulsed Amperometric Detection method (Thermo - Model ICS-3000) and iii) anions (Cl⁻, NO₃⁻, SO₄²⁻, MSA,
 Oxalate) and cations (K⁺, Na⁺, Mg²⁺, Ca²⁺) using ion chromatography (Thermo - Model ICS-5000).
 Proton Transfer Reaction - Mass Spectrometry (PTR-MS): Air was sampled through a 20m long, 3/8" o.d. (1/4" i.d.)

174 sheathed Teflon line that ran from the roof of the building to the instrument. A Teflon filter (0.2µm diameter porosity) was 175 installed at the inlet to prevent large aerosol particles and insects from entering the sampling line. The resulting residence time 176 of air in the line was estimated to be ea. approximately 0.5 min. The Ttemporal resolution of Volatile Organic Compounds 177 (VOCs) measured by the PTR-MS (Ionicon Analytik, Austria) was approximately two minutes (the time required to measure 178 55 different ions at 2 seconds per ion). The basic operation principles of the PTR-MS instrument have been described in detail 179 by Lindinger et al. (2011). Briefly, a stable flow of air and high concentrations of H_3O^+ ions are continuously sampled into a 180 drift tube held at 2.2 mbar pressure. There, compounds with a proton affinity greater than water, including a large selection of Oxygenated Volatile Organic Compounds (OVOCs), undergo efficient proton-transfer reactions with the H₃O⁺ ions to produce 181 182 protonated organic product ions, which can be detected by a mass spectrometer. 183 Meteorological Parameters: Standard meteorological parameters (temperature, relative humidity, wind speed and direction)

184 were obtained <u>174 asl</u>0 m above ground at the Athalassa Forestry Parkfrom the <u>m</u>Meteorological station of the Cyprus

185 Department of Meteorology, located installed 10 m above ground, located at the Athalassa Forestry Park (164 m asl) lying

186 <u>approximatelyat e.a.</u> 1.3 km east from of the CAO-NIC station. Wind speed and direction data were further used in this study

for component-specific non-parametric wind regression analysis (NWR) performed using the ZeFir toolbox (Petit et al., 2017)
 developed within the Igor Pro software (Wavemetrics Inc.). A co-located automatic CIMEL CE370 micro-LIDAR was

189 operated continuously to retrieve the Planetary Boundary Layer Height (PBLH) and better assess the influence of atmospheric

190 dynamics on in-situ ground-based observations.

191 Air masses back trajectory analysis: Five-day air mass back trajectories arriving at 1000m altitude above the sampling site

192 were computed every 6 hours, using the Hybrid Single-Particle Lagrangian Integrated Trajectory model (HYPLIT4; Stein et

193 al., 2015) using the Global Data Assimilation System (GDAS 1) meteorological data fields (with 1° spatial resolution). Back

trajectories were coupled to measured concentrations, assessing origins and source contributions to specific chemical components, by applying the Potential Source Contribution Function (PSCF) technique as implemented in the ZeFir toolbox

196 described above.

197 2.4. Source Apportoinment Apportionment analysis

Positive Matrix Factorization (PMF) is an advanced multivariate factor analysis tool that attempts to identify the contributing factors, or sources, of atmospheric pollutants at a sampling site. For this study, source apportionment was performed on the organic mass spectra dataset collected by the ACSM. The (PMF) method (Paatero and Tapper, 1994) using the multilinear engine (ME-2) model developed by Paatero (1999) was implemented using the SoFi (Source Finder) toolkit (SoFi 6D; Canonaco et al., 2013). PMF allows the decomposition of the OA mass spectra matrix X into two matrices, G and F and a remaining residual matrix, E:

204 X = G * F + E (1)

205 Where X is the input dataset matrix (measured quantity), F is the resulting source profile matrix, G is the source contribution 206 matrix (temporal variability of each source), and E represents the model residual matrix. Based on a number of criteria, the optimal solution is selected, aiming at being physically meaningful that can be supported by external indicators (ancillary 207 208 measurements), and trying to minimize values in the residual matrix E. Model input data and error matrices (in µg m⁻³), were 209 exported using the ACSM software. Data points with a signal-to-noise (S/N) ratio smaller than 0.2 were removed. and pPoints 210 with S/N between 0.2 and 2 were down_weighted by increasing their estimated error values (Ulbrich et al., 2009; Paatero and 211 Hopke, 2003). Mm/z (mass-to-charge ratio) values ranging from 10 to 120 were used in the analysis. CO2-related variables 212 were excluded from the PMF and finally reinserted into the solution.

213 Source apportionment of OA was performed following the general steps described by Crippa et al. (2014) and the recently 214 updated harmonised standard operating procedures for seasonal OA PMF (Chen et al., 2022). As a first step, unconstrained 215 PMF analyses were performed with a number of factors ranging from 2 to 8 in order to identify the most relevant number of 216 factors and potential sources. If primary organic aerosol factor profiles such as Hydrocarbon-like OA (HOA); or biomass 217 burning-like OA (BBOA) were found, then the corresponding site-specific primary OA (POA) mass spectra (see discussion 218 below) or spectra found in the literature (e.g., Ng et al., (2011) and Crippa et al., (2014)) were set as constraints in the PMF, 219 using the "a-value" approach (Paatero and Hopke, 2009; Canonaco et al., 2013). A sensitivity analysis was then performed 220 with different a-values to assess the level of constrain introduced in each factor with-i) a constrained HOA using, as an anchor 221 the HOA spectrum found in Ng et al. (2011) with the a-values ranging between 0.05 and 2.0, ii) a constrained BBOA factor 222 with the a-values from 0.2 to 0.5 from Ng et al. (2011), and iii) a constrained cooking OA (COA) factor from Crippa et al. 223 (2014) Mohr et al. (2012) with a-values from 0.2 to 0.5. Once this sensitivity analysis was completed, the evaluation of the 224 PMF results showed that the BBOA factor could not account for the entire m/z 60 mass fragment, which fragment was 225 distributed within 2 factors. Additionally, the correlation of BBOA with BCwb showed to be unsatisfactory (section S1). On 226 the other hand, given the BBOA factor's sensitivity to the type of solid fuel used, different biomass-burning factor profiles 227 have been reported in various regions around the world (Mohr et al., 2012)(Xu et al., 2020; Trubetskaya et al., 2021).(Xu et 228 al., 2020; Trubetskaya et al., 2021)(Xu et al., 2022); Trubetskaya et al., 2021)(Xu et al., 2020; Trubetskaya et al., 2021)(Xu et 229 al., 2020; Trubetskaya et al., 2021)(Xu et al., 2020; Trubetskaya et al., 2021)(Xu et al., 2020; Trubetskaya et al., 2021). 230 Consequently, a site-specific BBOA factor profile (BBOAcy) was selected. The BBOAcy spectrum was calculated as an average 231 of 20 PMF runs from the initial unconstrained PMF for the cold period, validated by it's time_series correlation to BCwb. Since

aged OA (i.e. Oxygen-like OA, OOA) factors show more variability between measurement sites in terms of their mass spectra, 232

- no constrain was introduced for these factors (Canonaco et al., 2015). 233
- 234 In this study, the BBOA factor - a major contributor of OA during winter - could not be properly resolved when performing

the PMF analysis on the entire period dataset. A seasonal approach was followed- instead, separating the OA dataset into two 235

236 periods that were then used to describe both the two periods (cold and warm, respectively). The criteria used to delineate those 237 two periods are presented and discussed in the below section 3.2.

238 One factor was consequentially constrained with the resulting BBOAcy spectrum (with an a-value in the 0-0.5 range, using

239 steps of 0.02), obtaining the optimal solution using an a-value equal to 0.46. A widely referred-to standard mass spectrum (Sun

240 et al., 2016; Duan et al., 2020) derived from Ng et al. (2011) was used to constrain the HOA factor, with an a-value of 0.2,

241 thus obtaining the best correlation with BCff, a tracer for traffic-traffic-related emissions. A detailed description of the OA

242 source apportionment analysis can be found in section S1 in the supplementary material.

3. Results and Discussion 243

244 3.1. On-line aerosol data quality check

245 A chemical mass closure exercise for PM₁ was performed at a temporal resolution of 1h to check the quality of the on-line aerosol measurements. Chemically reconstructed PM1 was calculated as the sum of the mass concentration of all non-refractory 246 247 species measured by the ACSM (OA, NO3, SO42, NH4+, Cl) plus the BC concentrations measured by the Aethalometer AE-248 33 (Putaud et al., 2004). The contribution of other chemical constituents to submicron aerosols, such as sea salt and dust (measured by co-located filter sampling), was found to be low and therefore neglected here. A scatter plot of the ACSM + AE-249 250 33 measurements vs. the SMPS-derived PM1 concentrations is shown in Figure S4ab. The results obtained indicate a very 251 good correlation ($r^2 = 0.88$; N=1823) and a slope of 1.2 (Fig. S4ab). This 20% discrepancy lies within the uncertainty of the 252 on-line instruments-and. It could be attributed to the cut-off size of the SMPS at 700nm, that-which is slightly lower compared 253 to than the ACSM. In addition, ACSM individual chemical species were compared with co-located off-line analyses performed 254 on daily PM2.5 filters. As shown in Fig. S4be-ef, very good agreement was obtained between on-line and off-line measurements 255 with $r^2 \ge 0.80$ (N=165-175) for all species. The discrepancy between ACSM and filter measurements for nitrate (slope of 1.3) 256 could potentially be attributed to the volatilization of HNO3 from the filter surface due to the presence of semi-volatile 257 ammonium nitrate. The obtained slopes for ammonium and sulfate below 1:1 (0.81 and 0.85, respectively) is are consistent 258 with the fact that fine (NH4)₂SO₄ aerosols, mainly originating from secondary processes and long-range transport (Sciare et 259 al., 2010; Freutel et al., 2013), can be found at a large size mode possibly exceeding 1 µm, consequently not being sampled by the ACSM. 260

261 The study investigated the aerosol ion balance using both online and offline inorganic measurements. The ratio of the measured 262 concentration of NH_4^{\pm} Measured and the estimated concentration of NH_4^{\pm} predicted, as calculated in (Jiang et al., (2019), was used for 263 this purpose. The results showed a slope of 0.80 for online measurements and 0.96 for offline measurements. These findings, 264 suggest that the atmospheric aerosol observed during the study period was mostlyfairlypredominantly neutral, taking into 265 account the uncertainties of ammonium concentrations reported in Q-ACSM intercomparison studies (Crenn et al., 2015), as 266

well as the species' relatively high detection limit (Ng et al., 2011).

267 An very strikinginteresting result obtained from the comparison of OA (ACSM) with OC (from filters) is an OM-to-OC ratio 268 of 1.42 which is at the lower end of ratios reported for urban environments, which usually exhibit typical values of 1.6 ± 0.2 (Petit et al., 2015; Theodosi et al., 2011; Brown et al., 2013). Without neglecting the fact that two different size fractions were 269 270

compared (PM₁ for the ACSM and PM_{2.5} for the filter sampling), this low ratio probably point to long-chain hydrocarbon OA 271

that often are related to primary combustion (poorly oxidized) OA This low ratio clearly denotes a major contribution of long-

272 chain hydrocarbon OA that often refer to primary combustion (poorly oxidized) OA(Aiken et al., 2008). As such, this ratio

273	could represent an independent means of verification of the consistency of our source apportionment between primary and
274	secondary OA.
275	As such, this ratio will represent a valuable and independent means of verification of the consistency of our source
276	apportionment between primary and secondary OA.
277	Finally, black carbon concentrations derived from light absorption measurements (Aethalometer AE-33) were compared
278	against filter-based EC measurements (see Fig. S4 \underline{fa}). Data from the two techniques correlate very well (r ² =0.83) ₁ with a
279	BC/EC ratio of 1.67 being similar to studies in other urban areas (Rigler et al., 2020; Liu et al., 2022), highlighting the existence
280	of a BC absorption enhancement (E_{abs}) attributable to a lensing effect induced by other chemical species, among which

281 secondary OA may play an important role (Zhang et al., 2018).

282

283 3.2 Meteorological conditions

284 Delineation of cold vs. warm seasons: The ACSM organic mass at m/z (mass to charge ratio)-60 is characteristic of the 285 fragmentation of levoglucosan, a product of cellulose pyrolysis and well-established biomass burning marker (Alfarra et al., 286 2007). Its respective contribution to total OA ($f_{\delta\theta}$) was used in this study as an indicator of biomass burning for domestic 287 heating to delineate cold vs. warm seasons, comparing with the 0.3% threshold proposed by Cubison et al., (2011) for air 288 masses influenced by biomass burning. Except for a single small peak in early May, corresponding to open fires for the 289 celebration of the Greek Orthodox Easter, the last instance when f_{60} was above the threshold was recorded during the first week of April (Fig. 2). From then onwards, daily air temperature started rising constantly, from aboutea. 15°C in-at_the 290 291 beginning of April up to to 30°C at the end of May. These two features dictated the division of the dataset into two periods: a cold period of four months (07/12/2018-08/04/2019), with an average temperature of 12 \pm 4°C, and a warm period of two 292 293 months (09/04/2019 – 31/05/2019), with an average temperature of $20 \pm 7^{\circ}$ C.



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Figure 2: Time series of air temperature (blue), m/z 60 organic concentration (org60, brown) and f_{60} fragment (green) for the cold and warm periods. The vertical line is used to delineate the measurements within the two seasons.

297 Wind sectors: During these two periods, a distinct pattern in the wind sectors and the air masses arriving at the sampling site

298 was observed. As seen in Fig. S5, the dominant wind direction for the cold period was the NW-SW [225 ° - 315 °] sector

299 encompassing 48% of the total wind directions, while the NE-SE [45 ° - 135 °] sector covered 26%. During the warm period,

300 the weight of this proportion is shifting even more towards the NW-SW [225 ° - 315 °] sector, having a 62% of total air masses

301 while only 17% are arriving from the NE-SE [45 $^{\circ}$ - 135 $^{\circ}$] sector.

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302 Air mass origin: A cluster analysis was performed (Fig. S6a,b) for both periods in order to better assess the main upwind 303 regions responsible for long-range transported air pollution over Cyprus and their change relative to the period of the year. 304 The number of clusters used in each season was determined by considering the percentage change in Total Spatial Variance 305 (TSV) as a function of the number of clusters of merged trajectories (Fig. S6c,d) and the mean trajectory paths of each cluster 306 (Fig. S6e,f). The first large drops observed in TSV from the two - to - three and the three - to - four cluster transition could 307 not represent all the recorded trajectories and especially the ones describing air masses arriving in Nicosia from the east. The 308 next remarkable decrease in TSV was recorded when moving to seven clusters. Thus, for both periods, seven clusters were 309 chosen to better represent all air masses arriving in Nicosia. 310 A significant part of most of the calculated mean trajectory path representing clusters arriving in Cyprus was found to be 311 related to the wider western sector, with many of them though, passing over Turkey before reaching Cyprus. Interestingly, this 312 analysis showed one cluster (Cluster 1) arriving from the Middle East (close to Lebanon and Syria) and another four (Cluster 313 1, 2, 5, 6) passing over the western part of Turkey for the cold period. For the warm period, the only clusters arriving from the 314 Middle East were the ones related to Turkey (Clusters 1, 5, 6). Most of the air masses arriving in Cyprus were found to 315 originate from Europe; many of them passing over Turkey before reaching Cyprus. Interestingly, this analysis showed one 316 cluster arriving from Middle East for the cold period, whereas there were not enough trajectories passingPlotting all individual

<u>72h back trajectories (Fig S6e,f) showed that a clear portion (almost 25% of the calculated trajectories) are being influenced</u>
 <u>by the Middle East, especially for the cold period (Fig S6e).</u> -over the Middle East to calculate a cluster for this area during the

319 warm period.

320 3.3. Chemical composition of PM1

321 Seasonal perspective of PM₁: <u>The Tt</u>ime series of PM₁ chemical composition derived from the ACSM (OA, SO₄²⁻, NH₄⁺,

 NO_3 , Cl⁻) and the Aethalometer (BC_{ff}, BC_{wb}) are depicted for the entire measuring period in Figure 3. Averaged data (6h

averaging period) are shown here for clarity. Furthermore, the relative average contribution of each chemical constituent to
 total PM₁ concentrations; is depicted in- the respective <u>inserted-inner</u> pie charts for both periods.



9

326	Figure 3: Stacked area plots of the chemical composition Fime series of the chemical composition of for PM44; in Nicosia derived	
327	from 6-hour averages of ACSM and AE 33 measurements. The vertical dashed red line separates the cold from the warm season.	1
328	The average relative contribution of each species is shown in the respective pie charts (inner panels) for each season	1
329	Although intense and shortduration peaks are observed for carbonaceous aerosols (OA, BC _{ff} , BC _{wb}), background NR-PM1	1
330	concentration levels (between peak values) remain well below 10 μ g m ⁻³ for the 6-h average in both seasons. In other words,	
331	no PM_1 pollution episodes (with e.g., concentrations above 10 μ g m ⁻³) lasting for consecutive days were observed. Such lack	
332	of intense and persistent PM_1 pollution episodes differs from what is reported in central and northern Europe, where stagnant	
333	(anticyclonic) conditions occur together with continental (polluted) air masses, mainly in winter and springtime (e.g., Petit et	
334	al., 2015). This suggests that the relatively low emissions from Cyprus (compared to the neighboring countries) together	
335	withand- its remote marine location (i.e., far from densely populated areas) may prevent the build-up of high PM1 pollution	
336	$events \ over \ Nicosia. \ On \ the \ other \ hand, \ clear \ differences \ can \ be \ observed \ between \ both \ periods, \ with \ significantly \ higher \ PM_1$	
337	concentrations during the cold period, associated with repeated, intense peaks of OA and BC - not observed during the warm	
338	$season-and\ suggesting\ local\ combustion\ emissions.\ The\ highest\ PM_1\ concentrations\ were\ observed\ between\ December\ 28^{th}$	
339	2018 and January 13th 2019 (Fig. 3) and were associated with low temperatures and Christmas holidays, both likely to promote	
340	the use of domestic heatingdomestic heating use. During the warm period, the higher contribution of sulfate, and lower	
341	contribution of OA, are clearly noticeable. The contribution of nitrate during the warm period, most probably in the form of	
342	semi-volatile NH4NO3, remains marginal, possibly due to non-favourable thermodynamic conditions preventing its formation	
343	and accumulation.	
344	PM₁ chemical composition : For the cold period, the average calculated mass concentration of PM ₁ (calculated as the sum of	

345 chemical components measured by AE_33 and ACSM) was $12.32-35 \pm 9.77 \text{ µg m}^3$, with $10.340 \pm 7.92 \text{ µg m}^3$ being the 346 average concentration of the non-refractory species (Table 1). OA constitutes the larger fraction of PM1 mass, with an average 347 concentration of 5.03 \pm 5.48 µg m³ (41 %), followed by sulfate (23 %), black carbon (16 %), nitrate (10 %), ammonium (9 348 %), and chloride (13%). These concentrations and the overall distribution of chemical components in NR-PM1 are quite similar 349 to those measured by ACSM in other European cities (Bressi et al., 2021). Concentrations appear to decline during the warm 350 period, with an average calculated PM₁ concentration of $8.18 \pm 4.65 \ \mu g \ m^3$, including $7.15 \ 18 \pm 3.80 \ 81 \ \mu g \ m^3$ from the non-351 refractory components. The dominant species during the warm period were sulfate and OA, each representing 35 % of PM1, 352 followed by black carbon (12 %), ammonium (11 %) and nitrate (6%). During that period, chloride concentrations were

353 negligible, contributing less than 1 % (Table 1).

354Table 1: Species mean, standard deviation, median concentrations and respective contribution to PM1 during cold and warm periods355in Nicosia.

	Cold Period				Warm Period				
µg m⁻³	Mean	Std	Median	Contribution (%)	Mean	Std	Median	Contribution (%)	
OA	5.03	5.48	3.35	40.81<u>41</u>	2.83	1.91	2.51	34.57<u>35</u>	
SO 4 ²⁻	2.8 <mark>41</mark>	1.89	2.60	22.83 23	2.87	1.50	2.61	35 .08	
NO₃⁻	1.22	1.25	0.75	9.87<u>10</u>	0.53	0.56	0.34	6 .46	
NH4 ⁺	1.14	0.77	1.01	<u>9.259</u>	0.92	0.55	0.84	11 .30	
CI-		0. 25 2				0. 11 0			
CI	0. 31<u>14</u>	<u>1</u>	0. 23<u>07</u>	2.53<u>1</u>	0. 05<u>03</u>	<u>8</u>	0.0 <u>1</u> 2	0.62<u><1</u>	
BC	2.01	2.31	1.26	16.33<u>16</u>	1.01	1.46	0.66	12 .36	
PM1	12. 32 3 <u>5</u>	9.77	10.01	100 .00	8.18	4.65	7.53	100 .00	

356

Interestingly, sulfate concentrations recorded in Nicosia are-much higher compared to what is commonly observed in other European countries and Mediterranean cities (Table 2) and likely reflect a regional pattern of sulfur-rich emissions compared to Europe, where SO₂ emissions have strongly decreased during the last decades (Smith et al., 2011; Chin et al., 2014) thanks

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to the implementation of specific abatement measures on reducing sulfur emissions (European NEC Directive (EU, 2016) and United Nation Gothenburg (1999) protocol). More specifically, the importance of sulfur emissions in Turkey (2 455 Gg, EEA 2021)₂ which were 50% higher compared to the total SO_x emissions of the EU 28 in 2019, together with the fact that half of the air masses reaching Cyprus are passing over Turkey (see Fig. S6) are key contributors to the high concentrations of sulfate in our study.

365 Shipping emissions appear to have a relatively minor impact on the concentration of sulfate. To more accurately determine the 366 contribution of shipping emissions to SO₄⁻², SO₂, and total PM_{2.5} a supplementary analysis was conducted using the WRF-367 Chem model, which simulates both physical and chemical processes occurring in the atmosphere. This model has been 368 extensively evaluated in several studies for the Eastern Mediterranean (Kushta et al., 2018) and Europe (Berger et al., 2016; Tuccella et al., 2012). Following the set-up used in (Giannakis et al., (2019) and driven by the EDGAR v.5 anthropogenic 369 370 emission inventories (Crippa et al., 2019), two annual-long simulations were performed: firstly, including all sectoral emissions 371 in the model (baseline simulation So) and a second simulation where shipping emissions have been omitted (scenario 372 simulation, S1) to identify the impact of shipping on gaseous and aerosol sulfur-related species concentrations (SO2 and SO4-373 2) and total PM_{2.5} over the Central and Eastern Mediterranean. The figures (S final) describe the contribution of shipping in 374 absolute terms (Fig. S7 a,c,e) and as a percentage (Fig. S7 b,d,f) for the SO₄⁻², SO₂ and total PM_{2.5} calculated for each species. 375 According to these results, the highest impact of shipping on near-ground modelled concentrations of the three species (SO4-376 ², SO₂ and PM_{2.5}) was estimated along the central Mediterranean region (yellow grids, west of the Balkans and Greece), as

377 well as a small section south of Greece. The Levantine basin, where Cyprus is located, experiences significantly lower

378 influence under the no-shipping emissions sensitivity test. More specifically, over the East Mediterranean, SO4⁻² concentrations

379 represent a relative change of only about 6-8% when including shipping emissions

380

Table 2: Comparison of concentration, and percentage contribution to PM₁, between the main submicron chemical species derived by ACSM.

	PM_1	OA	SO_4^{2-}	$\mathrm{NH_4^+}$	NO ₃ -	Cl	Poforonco
	(µg m ⁻³)	Kelelence					
Nicosia Cold (DJFM)	12.32	5.03	2.81	1.14	1.22	0.31	This study
Nicosia Warm (AM)	8.18	2.83	2.87	0.92	0.53	0.05	This study
Cyprus RB* (Annual)	7.6	3.26	2.66	0.98	0.23	-	Chen et al. (2022)
European UB** (Annual)	10.6	5.3	2.0	=	1.9	-	Bressi et al. (2021)
S. Europe RB*** (Annual)	6.3	3.5	1.3	=	0.8	-	Bressi et al. (2021)
Athens Winter	18.7	13.13	2.4	=	1.8	0.14	Stavroulas et al. (2019)
Athens Spring	6.42	3.3	2.1	0.6	0.4	0.02	Stavroulas et al. (2019)
Marseille Winter	11.9	6.17	1.12	0.86	1.58	0.09	Chazeau et al. (2021)
Marseille Spring	8.09	3.86	1.06	0.70	1.13	0.04	Chazeau et al. (2021)
Barcelona (Annual)	9.85	4.10	1.70	1.05	1.35	0.06	Via et al. (2021)

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383 * Cyprus Regional background

384 ** European urban background = Barcelona (Spain) + London (UK) + Prague (Czech) + Tartu (Estonia) + Zurich (Switzerland)

385 ***Southern European regional background = Ersa (Corsica, France) + Finokalia (Crete, Greece)

386 The main difference between the cold and warm periods lies in the decrease of <u>in the</u> concentration of carbonaceous aerosols

(OA, BC) and NO_3^- by almost a factor of two. Several phenomena can explain this significant seasonal variation: the absence of a domestic heating source (mainly biomass burning as explained in Fig. 2): the absence of Middle East air masses during

388 of a domestic heating source (mainly biomass burning as explained in Fig. 2); the absence of Middle East air masses during 389 the warm period (see discussion later on); the increase in the Planetary Boundary Laver Height (PBLH) above Nicosia (Fig.

the warm period (see discussion later on); the increase in the Planetary Boundary Layer Height (PBLH) above Nicosia (Fig.
 S<u>8</u>7) enhancing vertical dilution of local emissions during the warm period and therefore lowering ground-based

391 concentrations; less favourable thermodynamic conditions, with warmer and dryer air, also preventing the condensation of

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392 semi-volatile species (e.g., ammonium nitrate). Sulfate concentrations do not exhibit a similar seasonal pattern and therefore 393 seem to be less affected by the above factors. On the contrary, the increase in photochemistry enhances the formation of sulfate 394 aerosols, and the decrease in precipitation enhances aerosol lifetime, strengthening_strengthening_the impact of long-range 395 transport.

3.4. Diurnal variability of PM1 chemical constituents 396

397 Figure 4 shows the diurnal variability of the PM1 species derived from the ACSM and AE-33 for both the cold (Fig. 4a) and warm (Fig. 4b) periods. The diurnal variability of the apportioned BC related to fossil fuel combustion (BC_{ff}) and wood-398

399 burning (BCwb) are also depicted here.



400 Figure 4: Median Ddiurnal variability trends of the main submicron chemical constituents (OA, SO4², NO5[,], NH4⁺, Cl⁻ and BC) during the 401a) cold and b) warm periods. The Diurnal profiles of BCff and BCwb are embedded shaded area represents the 25th and 75th percentiles of 402 the diurnals

403 Organic aerosols: Organic aerosols clearly dominate the cold period PM1 concentration levels, exhibiting a night-time 404 maximum above 12 μ g m³; and a second smaller maximum at 4 μ g m³; coinciding with local traffic rush hour (06:00-09:00

405 LT). Elevated OA concentrations in the cold period during the night (max at 22:00 LT) are a common, well-documented 406 feature in many urban environments across Europe and the Mediterranean (e.g., Florou et al., 2017; Stavroulas et al., 2019;

407 Chazeau et al., 2021). They can be attributed to higher emissions from domestic heating, evening traffic peak and cooking

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408	activities. The strong correlation between OA and BCwb (R ² =0.81; N=2934; Fig. S28) suggests that residential wood burning
409	is an important contributor to this nighttime peak. Interestingly, this peak is not significantly amplified by a lower PBLH
410	during night-time, which seems to remain relatively stable with no significant diurnal variability during the cold period (Fig
411	S_{27}^{87}). It is also worth noting that background OA concentrations observed both at the end of the night and middle of the day
412	when local emissions are minimal, remain relatively high at aroundea. $3 \ \mu g \ m^{-3}$. The diurnal variability of OA is much less
413	pronounced during the warm period, suggesting a more important contribution of regional sources to OA compared to the
414	strong dynamic of local emissions. The assumption of a more important contribution from regional OA during the warm period
415	is further supported by a mean OA concentration of 2.83 μ g m ⁻³ (Table 2) that is close to the averaged OA concentrations of
416	$3.2\mu g\ m^{-3}$ reported for a 2-year period continuous observations with Q-ACSM (2015-2016) at the rural background site of the
417	Cyprus Atmospheric Observatory at Agia Marian Xyliatou (CAO-AMX), at aca. 40km distance roughly 40 km from Nicosia
418	(Chen et al., 2022). During the warm period, a small OA peakremains visible in the morning, with a similar amplitude to
419	the cold season, likely to be related to traffic emissions. A second peak can be observed at 21:00 LT (not observed in BC)
420	which may possibly originatepotentially originating from cooking activities. Heavy oil combustion from shipping could
421	possibly contribute to OA. Further to the poor contribution of shipping emission on OA, a model study of sources of organic
422	aerosols in Europe using CAMx (Jiang et al., 2019) showed that the contribution of "other anthropogenic sources" (gathering
423	shipping, industry, and energy production) on OA (POA+SOA) was, typically, of the order of 10% during summer and winter
424	in the Eastern Mediterranean region close to Cyprus. Based on a simple receptor model, PM25 source apportionment performed
425	in Nicosia, Achilleos et al. (2016) showed that the contribution from shipping is approximately 8% to PM25. Most of the
426	transported mass is attributed to $SO_4^{2^2}$ with a minor contribution from carbonaceous aerosols. In conclusion, shipping emissions
427	are likely to play a minor role in OA concentrations.
428	Black carbon: During the cold season, BC follows a bimodal diurnal pattern, which can be further apportioned by focusing
429	on its source-specific components BC_{ff} and BC_{wb} . The fossil fuel component exhibits two maxima, one in the early morning
430	coinciding with traffic rush hour, and one in the late afternoon, most probably related to both traffic and an increase in energy
431	demand due to domestic heating (see discussion later on). On the other hand, BCwb diurnal variability is dominated by a night-
420	

432 time maximum (20:00 - 01:00 LT), peaking one hour after BC_{ff} and linked to wintertime residential wood-burning emissions, 433 contributing up to 33 % of total BC. During the warm season, the BC diurnal pattern is characterised by the absence of a night-434 time maximum, while still exhibiting a significant peak in the morning, dominated by BCfr. The very low contribution of 435 biomass-related combustion particles during the warm period, as previously noted from m/z 60 in Fig. 2, is further supported 436 here, with BCwb exhibiting a nearly flat diurnal variability with close-to-zero mass concentrations. The contribution of shipping 437 in the Mediterranean on Black Carbon (BC) concentrations was investigated from model estimates by (Marmer et al., (2009) 438 based on three (3) most commonly used ship emissions inventories: 1) EDGAR FT by (Olivier et al., (2005), 2) (Eyring et al., 439 (2005), and 3) EMEP by Vestreng et al. (2007). Results showed that shipping emissions were contributing to typically 15-25% 440 of BC in the E. Mediterranean, far from the shipping routes (which is the case for Cyprus). A similar result was found from a 441 more detailed (Positive Matrix Factorization) PM2.5 source apportionment analysis performed in Nicosia in 2018, with heavy 442 oil combustion contributing 7% to PM2.5 (Bimenyimana et al., 2023 under review), and the relevant factor containing less than

443 0.1μg m⁻³ of EC.

Secondary inorganic aerosols: During the cold season, non-refractory nitrate and chloride detected by the Q-ACSM are mostly present in the form of semi-volatile NH₄NO₃ and NH₄Cl (Guo et al., 2017; Theodosi et al., 2018). They show a nighttime maximum (Fig. 4-a), reflecting the presence of gas precursors (NH₃, HNO₃, HCl) and the more favourable thermodynamic conditions with lower temperatures, higher relative humidity, and condensation sink due to high PM concentrations of combustion aerosols (traffic, domestic heating). Additionally, there is a smaller morning NO₃⁻ peak_± most probably linked to traffic (Foret et al., 2022). This is not observed for chloride, suggesting that HCl may not be as abundant in the morning Formatted: Subscript

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450 compared to the evening. The less favourable thermodynamic conditions during the warm period leads to very small 451 concentrations of semi-volatile NO3⁻ and Cl⁻ (Fig. 4b). As expected, sulfate does not show a pronounced diurnal pattern, 452 irrespective of the period, and pointing to regionally-processed aerosols (Fig. 4a,b).

453 3.5. OA Source Apportionment

454 3.5.1 OA source apportionment during the cold period

455 For the cold period, the optimal PMF result has been found using a 5-factor solution following the approach detailed in section 2.4. The identification of OA sources related to these 5 factors was then performed following the typical combination of 456 457 information from i) OA mass spectra (Fig. \$59a), ii) the correlation of each factor with source-specific tracers (see Fig. 5b), 458 iii) their diurnal variability (Fig. 6a), and iv) their daily (week days vs. week-end) pattern (also Fig. 6b). The five factors were 459 then assigned to the following sources: A primary BBOA (Biomass Burning Organic Aerosol), two primary HOA 460 (Hydrocarbon-like Organic Aerosol; HOA-1 and HOA-2) and two secondary OA sources, namely low-volatile MO-OOA 461 (Low volatile More-Oxidized Oxygenated Organic Aerosol) and semi-volatile LO-OOA (Semi-volatileLess-Oxidized 462 Oxygenated Organic Aerosol). This source apportionment is presented and justified below for each factor:

463 HOA-1 (Hydrocarbon-Like OA Type 1): The mass spectrum of HOA-1 (Fig. 59a5a) is consistent with a fossil fuel (traffic) 464 combustion source that can be identified by the prevailing contributions of the ion series representing C_nH_{2n-1} (m/z = 27, 41, 465 55, 69, 83, 97, typical fragments of cycloalkanes or unsaturated hydrocarbon chains) and C_nH_{2n+1} (m/z = 29, 43, 57, 71, 85, 466 99, typical fragments of alkane chains). Hence, this factor mass spectrum is well correlated to eight selected HOA factors 467 related to vehicular traffic found in the literature (Fig. S10a) and relevant to European and Mediterranean environments. The 468 traffic-related origin of the HOA-1 factor can be further confirmed by the good correlation with BC_{ff} (R^2 =0.65; N=2934; Fig. 469 S11a), benzene (R²=0.72; N=1165; Fig. S11b). The diurnal variability of HOA-1 shows a bimodal cycle with a sharp maximum 470 during the morning rush hour with an amplitude similar to BCrf (Fig. 6a), and a broader maximum in the evening possibly 471 encompassing emissions from traffic and diesel-fired residential heating systems. In the weekly cycle, as depicted in Fig. 6b, 472 the morning peak decreases in on Saturday and . It is nearly absent on Sunday mornings, aligned with the de-escalation of 473 traffic emissions usually observed during weekend mornings.

474

475 BBOA (Biomass Burning OA): The mass spectrum of the site-specific BBOA factor (reported as BBOA_{cy} in section 2.4) 476 exhibits characteristic peaks at m/z 29, 60, and 73 (Fig. 59a5a), which are indicative of biomass burning (Crippa et al., 2014). 477 The mass spectrum is quite similar to other BBOA spectra found in the Mediterranean and Europe (Fig. S10c), with a key 478 difference here being the rather low contribution of <u>a</u> signal at m/z=43.- The biomass burning-related origin of the factor is further confirmed by the strong correlation with BC_{wb} (R²=0.81; N=2934; Fig. S11c), benzene (R²=0.61; N=1162; Fig. S11d) 479 480 and levoglucosan (R²=0.94; N=125; Fig, S11e) a typical tracer of biomass burning (Fourtziou et al., 2017). The BBOA diurnal 481 pattern exhibits an expected well-marked night-time maximum around 22:00 LT, consistent with residential wood_-burning activities. This night-time maximum is observed throughout the week (Fig. 6a), confirming the important role of wood burning 482 483 for heating in the city. Interestingly, the higher concentrations of BBOA as well as BCwb (Fig. 6b) were observed on Sunday 484 evenings, pointing to a-the recreational use of fireplaces, leading to enhanced residential wood-burning emissions during the 485 weekend, a feature also reported in other sites in Europe and the US (Bressi et al., 2016; Rattanavaraha et al., 2017; Zhang et 486 al., 2019).

HOA-2 (Hydrocarbon-Like OA Type 2): The mass spectrum obtained for this factor (Fig. <u>S9a5a</u>) is similar to the HOA-1 factor, with high signals for the ion series $C_nH_{2n+1}^+$ and $C_nH_{2n-1}^+$. The main differences between these two factors occur in the relative contribution of m/z 41 compared to m/z 43 and the relative contribution of m/z 55 compared to m/z 57, which are both much higher for HOA-2, than for HOA-1. Furthermore, the contribution of signal to m/z 44 is more significant in HOA-2, which can imply a mix of various sources and/or -a possible possibly higher degree of atmospheric processing. Other discrepancies with HOA-1 concern its diurnal variability, with an intense maximum at night (Fig. 6a), and even more its average concentration levels, which are almost three times higher than HOA-1.

494 Influence of cooking activities: The HOA-2 diurnal profile has a small peak at 13:00 LT and a significantly higher one at 21:00 495 LT, effectively coinciding with typical meal times in Cyprus as well as those reported in the literature for Greece (Siouti et al., 496 2021), therefore indicating the influence of cooking activities to this factor. When plotting f_{55} vs f_{57} (Mohr et al., 2012) and 497 colouring the data points by the corresponding time of day, a distinct pattern appears with data of higher f_{55} over f_{57} being 498 clustered to the top left of the triangle, close to the fitted lines representing cooking (Fig. S12) and coinciding with midday and 499 evening hours. The night-time maxima pattern is consistent throughout the week, with the higher concentrations being recorded on Friday and Saturday evenings (Fig. 6b), in line with an expected food service sector activity increase as part of Nicosia 500 inhabitants' leisure in the weekend. The mass spectrum of HOA-2, even though left unconstrained, is highly correlated to COA 501 502 found in other studies (Fig. S10b) in both Mediterranean and continental European urban environments, Additionally, the nonnegligible signal at m/z=60 points to the widely spread habit of meat charbroiling (Kaltsonoudis et al., 2017). 503

Influence of power plant emissions: A closer look at the diurnal variability of the HOA-2 factor shows a certain persistence of 504 505 this factor throughout the day, even when cooking activities are more or less absent (Fig. 6a). Such pattern could imply the 506 influence of other combustion sources, not necessarily of local origin. The influence of other combustion sources would also 507 help to explain why HOA-2 average concentrations are roughly 3 times higher than OA related to traffic (HOA-1), as it is very 508 unlikely that cooking activities can contribute solely to the observed HOA-2 concentrations. A possible contributing source 509 could be related to the energy production sector in on the island, which relies exclusively on heavy fuel oil. In a recent study, 510 Vrekoussis et al. (2022), utilizing satellite observations, have identified that power plants located to the North (Teknecik 511 powerplant, PP4, 362MW), North-East (Kalecik powerplant, PP5, 153MW) and South-East (Dhekelia power station, PP3, 512 460MW) of Nicosia at 22 km, 60 km and 38 km, respectively, are significantly contributing to columnar NO₂ concentrations 513 over the island. The importance of these emission hotspots, along with their location on the island, during both the cold and 514 warm periods is illustrated in Fig. S13 and shows, in particular particularly for the Northern power plants (PP4, PP5), emissions 515 as high as the traffic-related NO2 over Nicosia. Interestingly, -in a source apportioning study on VOCs performed at the Cyprus 516 Atmospheric Observatory - Agia Marina Xyliatou (CAO-AMX), a rural remote site 32 km southwest of Nicosia, Debevec et 517 al. (2017) have resolved a factor related to industrial activity/power generation, exhibiting a connection with winds arriving 518 from the wider eastern sector. 519 In order to assess the possible influence of Cypriot power plant emissions, the coupling of wind velocity, and wind direction

520 with the HOA-2 time_series was performed through NWR analysis (Fig S14b). This analysis highlights the association of 521 stagnant conditions (low wind speed / low dispersion) with high HOA-2 concentrations (i.e., night-time peaks), pointing to a 522 more local origin for this OA source. On the other hand, different features appear when wind velocities are higher, showing 523 emissions originating from the NW and the E-NE sectors; i.e. downwind of power plants PP47 and PP5, although long-range 524 transport influence cannot be ruled out. This is illustrated by the NWR of sulfate (Fig. S14f), which shows a dominant E sector 525 likely to originate from regional emissions. Given the positioning of the sampling site, close to the edge of Nicosia'sNicosia 526 urban fabric, with the Athalassa park lying to the east, such an observation can suggest the transport of plumes from the 527 operating powerplants, namely PP5 and PP3 to the city. Interestingly, a similar, yet even clearer image stands for SO_2 528 concentrations - only half -of which are considered to be of urban origin (Vrekoussis et al., 2022) - measured at a suburban 529 background site (NicRes) and a traffic site (NicTra) in the city (Fig. S14g-h), with elevated SO₂ concentrations being related 530 to eastern winds of higher velocity, further corroborating that power generation related polluted plumes, traveling through the 531 Mesaoria plain arriving to Nicosia can contribute to the HOA-2 factor.

532 Other combustion sources: Interestingly, chloride shows a good correlation with HOA-2 (r²=0.61; N=2945; see Fig. S11f, Fig.
533 5b). Chloride detected by the ACSM is in the form of NH₄Cl (a secondary highly-volatile species). The source of this chloride

534 is still widely debated and may originate from industrial activity or municipal (plastic-containing) waste burning (Gunthe et

al., 2021). Another possible explanation of the good agreement between HOA-2 and chloride would be the use of Cl-rich coal
 as a means for outdoor cooking in Nicosia could therefore reflect the influence of cooking activities that comprises a fraction
 of the HOA-2 factor

Less-Oxidized Oxygenated OA (LO-OOA): With elevated contribution of m/z 44, the mass spectrum of this factor is 538 539 consistent with a secondary OOA source. A higher m/z 43, and a lower m/z 44 (Fig. 59a5a) compared to MO-OOA, implies a 540 less oxygenated (less-processed) component (Mohr et al., 2012). Finally, the time series of this factor is quite similar to NO3-, with an overall good correlation value ($R^2 = 0.67$, N=2943; Fig. S11h), highlighting its semi-volatile character. This is further 541 542 corroborated by the very good correlation of LO-OOA with chloride (R² = 0.73, N=2943; Fig S11i), another semi-volatile 543 compound measured by the O-ACSM. The diurnal variation of LO-OOA displays 1.5 times higher concentrations during the 544 night compared to daytime (maximum of 1.84 ± 0.31 µg m⁻³ at 22:00 LT; Fig. 6a); a pattern that is much more pronounced 545 than the variability observed for MO-OOA. This feature highlights the fact that the presence of LO-OOA, is not exclusively 546 controlled by photochemical processes. Instead, changes in thermodynamic equilibrium (due to lower T and increased RH), 547 favouring the condensation of gas-phase semi-volatile material on the one hand, and intense night-time chemistry (gas phase 548 or heterogenous) on the other hand, are among the processes that may account for the rapid night-time formation of LO-OOA. 549 Atmospheric processing of biomass burning OA during periods of low photochemical activity (such as in winter or at night). 550 known alsoalso known as "dark" aging, have has been reported recently (Kodros et al., 2020; Jorga et al., 2021) and could 551 have contributed to the observed night-time formation of LO-OOA. Notably, the weekly cycle of LO-OOA, and its night-time 552 maxima, appears to have the same pattern and intensity as those observed for BBOA (e.g., low peaks on Tuesday/Thursday, 553 maximum on Sunday) (Fig. 6b). On the other hand, the factor is correlated with both BBOA (R²=0.81; Fig. S11k) and BC_{wb} 554 (R²=0.66; Fig. S11j). This observation could indicate a biomass_-burning contribution to LO-OOA; through fast oxidation of 555 primary emissions, supported in-by several studies showing biomass burning linked to OOA sources at night (Stavroulas et 556 al., 2019: Kodros et al., 2020: Chen et al., 2021). 557 More-Oxidized Oxygenated OA (MO-OOA): The MO-OOA factor typically accounts for secondary organic aerosol formed 558 in the atmosphere from gas-to-particle conversion processes of VOCs and their products, as well as atmospheric ageing of 559 primary OA (Petit et al., 2015; Stavroulas et al., 2019). Numerous VOC sources can contribute to OOA but lose their mass 560 spectrum fingerprint owing to extended oxidation due to photochemical aging, which leads to enhanced signal at the m/z 44 561 fragment (CO2⁺), a dominant tracer for OOA (Ng et al., 2011). The predominance of m/z 44 and the near absence of m/z 43 in

562 the mass spectrum of the resolved MO-OOA factor (Fig. 59a5a) points to highly oxidized/aged secondary OA (i.e., originating 563 from long-range transport). This is further supported by the relatively good agreement (R²=0.55; N=2943; Fig. S111) between 564 concentrations of MO-OOA and sulfphate (Fig. 5b), a species of regional origin (Sciare et al., 2003). Nevertheless, the diurnal 565 variability of MO-OOA does not closely follow sulfate showing a small increase of 20-30% every evening (Fig. 6a,b), which furthermore cannot be explained by atmospheric dynamics (c.f. the negligible PBLH diurnal variability for the cold period 566 567 shown in Fig. S§7). Alternatively, this would suggest that a fraction of MO-OOA is produced locally through night-time 568 oxidation mechanisms as previously observed for LO-OOA. Similar nighttime increases of high oxygenated OA factors, 569 related to local sources, have been reported in both northern European urban sites (Zhang et al., 2019; Lin et al., 2022) as well 570 as in the Eastern Mediterranean urban environment (Athens, Greece), where a link to oxidized primary residential wood

571 <u>burning emissions as a potential driver of the low volatility OOA factor diurnal variability, was also suggested (Stavroulas et</u> 572 <u>al., 2019).</u>



577 Figure 6: Diurnal variability (left) and weekly cycles (right) of the five OA factors averaged over the cold period..

578 3.5.2. OA source apportionment during the warm period

For the warm period, the optimal PMF solution was obtained using a 4-factor solution (HOA-1, HOA-2, MO-OOA, LO-OOA). As expected, the BBOA factor could not be resolved, as previously highlighted by the low concentrations at m/z 60 reported during this period (Fig.2). Again, the identification of OA sources related to the 4 OA factors was performed following the typical combination of information from i) OA mass spectra (Fig. S9b7a), ii) the correlation of each factor with external sourcespecific tracers (Fig. 7b and Fig. S15), iii) their diurnal variability (Fig. 8a), and iv) their daily (week-days vs. week-end) pattern (also Fig. 8b). The mass spectra profiles for the 4-factor PMF solution during the warm period (Fig. S9b7a) were quite similar to the ones from the cold period (Fig. S9a5a).

586 HOA-1: For the warm period, an a-value of 0.2 was selected for constraining the HOA-1 factor, again using the Ng et al. 587 (2011b) HOA profile as a reference. The resolved factor profile is nearly identical to the one obtained for the cold season (R^2 588 = 0.99, Fig. S10a), and It is also very well correlated to traffic-related HOA factor profiles found in other Mediterranean (Kostenidou et al., 2015; Gilardoni et al., 2016; Florou et al., 2017; Stavroulas et al., 2019) and European cities (Lanz et al., 589 2010; Crippa et al., 2014) as depicted in detail in Fig S10a. The HOA-1 time series follows the same pattern as the 590 corresponding traffic-related HOA-1 factor reported for the cold period, showing a good correlation with BC_{ff} , (R^2 =0.62, 591 592 N=1259; Fig. S15a). Its diurnal variability exhibits a bimodal pattern, with a typical sharp maximum in the morning (07:00 593 LT) and a smaller peak during the evening (Fig. 8a). On a weekly basis, this diurnal variability tends to be less pronounced on Saturdays and nearly absent on Sundays (Fig. 8b), reflecting reduced commuting during the weekend. 594

595 HOA-2: The HOA-2 factor still shows elevated concentrations during the warm period, close to 3 times higher compared 596 tothan HOA-1 (Table S_{32}^2). The profile remains quite unchanged between the cold and warm periods ($R^2 = 0.92$; Fig. S10b), 597 pointing to similar sources. No correlation was observed with chloride, which may be expected due to unfavourable 598 thermodynamic conditions hindering NH4Cl formation as well as the lack of significant chloride sources during this period. A 599 night-time maximum of HOA-2 is still observed when investigating the factor's diurnal variability (Fig. 8a). Furthermore, a somewhat broader, compared to the cold period, maximum in the middle of the day (Fig. 8a) can also be observed. When 600 going through the weekly variability, this midday maximum is particularly well defined on Sundays (Fig. 8b), while the 601 602 evening peaks of Sundays and Mondays are the lowest. The above observations remain consistent with the cold period 603 assessment,- that HOA-2 is on the one hand linked to cooking activities. For households activities are expected at noon and 604 evenings, while for restaurants, activity peaks on Sunday noon and is lower on Sunday evening and Monday, reflecting the 605 fact that such businesses remain closed on the first day of the week (Fig. 8b). On the other hand, the overall offset of HOA-2 606 observed against the HOA-1 diurnal profile persists, suggesting somewhat permanent background HOA-2 concentrations that 607 cannot be explained by cooking activities alone. A contribution to this source by continuous emissions from power plants (see 608 space-based (SP5-TROPOMI) vertical columns of NO2 during the warm period in Fig.S13d), should be sought. In addition, 609 the HOA-2 NWR plot for the warm period reveals an even more significant enhancement of concentrations when moderate 610 winds blow from the E-SE (Fig. S16b), a trend also observed for SO2 during the same period (Fig S16e,f).

611 The above observations remain consistent with our assessment for the cold period: the HOA-2 factor consists of a mixed OA 612 source that contains cooking activities (inc. coal combustion) and emissions from the powerplants located on the eastern part 613 of the island. Indeed, the HOA-2 midday maximum can be linked to an increase in electricity demand at that time of day during 614 the warm period due to an increase in air conditioning usage (Cyprus' NECP 2021-2030, 2019).

615 **LO-OOA:** The LO-OOA factor profile exhibits some differences with from the one resolved for the cold period ($R^2 = 0.66$)_a 616 as illustrated in the correlation matrix of comparison to selected factor profiles found in the literature (Fig. S10d) while being 617 very similar to those obtained in Athens/Piraeus during summer (Bougiatioti et al., 2014; Stavroulas et al., 2021). The LO-618 OOA time_series shows a low agreement with NO₃⁻ ($R^2 = 0.31$; N=1259; Fig. S15c) poorer than the observed correlation 619 during the cold period (Fig. S11h). The diurnal pattern of the factor (Fig. 8a) shows maximum concentrations persisting 620 throughout the night and early morning, while a secondary maximum during the midday can be observed. But overall, the

621 diurnal pattern of LO-OOA is rather flat compared to the cold period, suggesting that local production may not be so important 622 at that time compared to a less variable regional background. Interestingly a midday hump similar to the one observed for 623 HOA-2 is present, suggesting a common origin.

624 MO-OOA: The factor profile of MO-OOA resolved during the warm period is strikingly identical to the profile found in the cold period (their R² is almost 1; Fig. S10e), while being excellently correlated to other highly oxygenated OA factors resolved 625 626 in both the urban and regional background in the Eastern Mediterranean (Bougiatioti et al., 2014; Stavroulas et al., 2019, 2021) as well as in continental Europe (Crippa et al., 2014). The winter night-time peaks are not observed anymore (Fig. 8a), with 627 628 the factor's diurnal pattern exhibiting much less variability, highlighting its dominant regional character. The time series of 629 MO-OOA correlates good-well to SO4²⁻ (R²=0.53; N=1259; Fig.S15b), confirming this regional and highly processed origin. 630 The concentration levels of MO-OOA during the warm period are lower than in the cold (Table S32). However, its relative 631 contribution to total OA during the warm period remains similar (45 %).



634 spectra of the PMF (a) and the Ttime series of the four OA factors resolved along with corresponding tracer Figure 7: Mass 635 compounds (b) for the warm period.





637 3.6. Spatial and seasonal variability of OA sources

638 3.6.1. Seasonal variability of OA sources

Primary OA: The mass concentration of the three primary OA factors (HOA-1, HOA-2, BBOA) represents as much as 40 % 639 640 of total organic aerosols during the cold period (Fig. 9), with POA contribution significantly decreasing in the warm period (22% to total OA) due to the absence of the significant residential wood burning source which during the cold period accounted 641 642 for 12% of total OA. The important contribution of primary sources in Nicosia has been also been highlighted earlier, by the 643 rather low OA/OC ratio of 1.42 (Section 3.1). In a recent publication covering several European sites, Chen et al. (2022) 644 reported that in urban sites, solid fuel combustion-related OA components were 21.4 % of total OA during winter months, 645 higher than what is found for BBOA in Nicosia, owing to the rather milder winters in the city. 646 The traffic_related primary factor in Nicosia (HOA-1) was found to be rather stable in terms of contribution to total OA across

this study's two seasons, averaging 7% and 6%, respectively, for the cold and warm periods, being lower than the figure reported in other European Urban sites (12.7%, Chen et al., 2022). On the other hand, the HOA-2 factor represents ca 2/3 of the total HOA in Nicosia with little variation from winter (72 %) to summer (66 %) to total HOA (Fig. 9). Comparing it with COA in urban locations resolved by Chen et al. (2022), during both winter (14.4% compared to 21% in the cold season in Nicosia) and spring (15% versus 16% in Nicosia during the warm season), the higher values reported in Nicosia further support the assumption that the HOA-2 represents a mixed combustion source.

653 Secondary OA: A higher degree of oxidation is observed for the LO-OOA factor during the warm period, given the much 654 higher contribution of signal signal contribution at m/z 44 compared tothan the respective cold period factor. This discrepancy, 655 reported in several studies (Huang et al., 2019; Duan et al., 2020), is explained by higher photochemistry during the warm 656 period, which promotes the oxidation of OA, resulting in an LO-OOA profile with a higher m/z 44 fraction. This result is also 657 consistent with a less-oxidized LO-OOA formed during the cold period from night-time chemistry. The range of LO-OOA 658 concentration levels are is different between cold and warm periods (0.05-7.74 µg m⁻³ and 0.05-4.00 µg m⁻³, respectively), 659 while the mean concentrations for both periods are similar (0.86 and 0.95 μ g m⁻³ for cold and warm periods respectively). The 660 contribution of LO-OOA relative to total OA is double during the warm period compared to the cold, reflecting both the 661 absence of the biomass burning source as well as the prevailing conditions favoring atmospheric processing of primary OA 662 and SOA precursors. During the cold period, LO-OOA intense peaks suggest an influence from local emissions, while during 663 the warm period, the less-variable LO-OOA diurnal variability highlights the influence of more intense photochemical processing at medium-to-large geographical scale. MO-OOA is found to be the major contributor to total OA for both the cold 664

(44%) and warm (45%) periods, higher in both cases than the average MO-OOA contributions reported for other European
urban sites (Chen et al., 2022) underlining the importance of highly processed secondary OA over Nicosia (Fig. 9).



668 Figure 9: Relative contribution of PMF resolved OA sources to total OA for the cold period (left) and the warm period (right)₂ 669 respectively.

670 **3.6.2. Geographic origin of OA sources**

667

671 The geographic origin of OA sources (local vs regional) is further assessed here using both Non-parametric Wind Regression

672 (NWR) analyses as well as the regional scale coupling concentrations to air mass back trajectories through PSCF.

673 Cold period: During this period, primary OA factors, especially HOA-1 and BBOA, have an expected strong local component 674 that is characterized by high concentrations at low wind speeds (hourly average 1.4 m s⁻¹) when winds are originating from the 675 W-SW sector (Fig. S14a,c), pointing to the busy highway connecting Nicosia to the other major cities in the island while 676 integrating the highly populated residential areas of Strovolos and Lakatamia municipalities. (Fig. 1c). As discussed earlier, 677 the HOA-2 factor, apart from its local influence (also in the W-SW sector), exhibits significant concentrations related to higher 678 wind speeds from the NW and the E-NE sectors that could originate from power plants but also possibly from long-range 679 transport. Interestingly, a small local contribution from the city, still within the W-SW sector, can be also be observed for 680 both LO-OOA and MO-OOA, consistent with the peaks observed that could originate from local night-time chemistry. Still, 681 high-concentrations of MO-OOA (and, to a lesser extent LO-OOA) are observed with high wind speeds and Eastern directions 682 (Fig. S14e,d). Although the contribution of the power plant PP5 located in the East sector (Fig. S13c) cannot be excluded, 683 PSCF analysis points out that the hotspots of MO-OOA can be traced in neighbouring countries (eg. Syria, Lebanon and, South 684 Turkey) in the middle East (Fig 10a). These areas also represent hotspots of SO₄²⁻ according to PSCF analysis (Fig. S17a). 685 Warm period: Given the generally higher wind speeds recorded, in comparison to the cold season (average of 1.93 m s⁻¹ vs. 686 1.36 m s⁻¹ in the cold period), all OA factors show elevated concentrations coupled with higher wind speeds. The most strieking 687 result is the major influence of the E-SE sector for all OA sources, However, although this sector is upwind of Nicosia and, 688 therefore, poorly influenced by local city emissions. As noted previously, for the cold period, long-range transported OA from 689 the Middle East is expected to be the main driver here-to explain the influence of the E-SE sector, at least for LO-OOA and 690 MO-OOA (Fig. S16c,d). This is once again confirmed from by the PSCF results reported in Fig. 10b for the warm period. The 691 HOA-1 factor still shows maxima for low wind speeds (<5 km h⁻¹) characteristic of local emissions and the SW-S direction, 692 but also exhibits significant contribution related to the E-SE sector. Although the influence of the power plant PP5 on HOA-2 693 is expected, the contribution of this source can not be excluded for HOA-1 as well. On the other hand, quantification of the 694 Middle Eastern contribution to the HOA-2 factor remains to be assessed, since the current dataset cannot provide sufficient

information on separating the contribution of power plants on the island versus more regional Middle East emissions (Fig.
S165b). Although this hypothesis needs further investigation, the presence of HOA-2 in the Middle East would be consistent
with recent findings highlighting the importance of OC emissions from diesel generators used in Lebanon as a means of
complementary power generation (Fadel et al., 2022).

699



700

 Figure 10: PSCF plots for MO-OOA during the cold and warm periods. Colorscale <u>The color scale</u> represents the probability of air parcels arriving at the receptor site (white dot) for measured concentrations higher than the 75th percentile. PSCF values while the sampling site is denoted with a white dot.

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In conclusion, based on the relative contribution of OA factors (Fig. 9) and the NWR analysis (Fig. S14, S16), it can be reasonably assumed that <u>a</u> significant amount of measured OA in Nicosia <u>is originatingoriginates</u> from long-range transport with the Middle East being the major source region, during both cold and warm periods. This is the first time that such <u>a</u> high contribution of OA from the Middle East is highlighted over Cyprus. Assuming that biomass combustion and biogenic emissions of OA in the desert regions of the Middle East are relatively limited, these results suggest that most of <u>the</u> primary and secondary OA originating from the Middle East could be of fossil fuel origin, which is consistent with the previously reported <u>large-extensive</u> use of oil in this region.

711 3.7. Spatial and seasonal variability of BC sources

712 The above conclusion on the influence of primary and secondary OA sources from the Middle East region, and its strong fossil 713 fuel origin, motivates a careful examination of the geographic origin and sources of BC concentrations recorded in Nicosia. 714 Baseline (i.e., lowest) BCff concentrations are typically observed in the middle of the night and in the middle of the day, when 715 local emissions are at their minimum (See Fig. 4). As such, these background concentrations can be considered as a first 716 qualitative indicator of background BC_{ff} concentrations of regional origin. Interestingly, these baseline BC_{ff} concentrations 717 appear to be in phase with those of sulphate sulfate (Fig. 11), as well as the MO-OOA factor derived from the OA PMF 718 analysis. This observation points to the possible use of MO-OOA as a tracer for regional BCfr. Hence, it brings further evidence 719 on of the importance of regional emissions on carbonaceous aerosol concentrations in Nicosia.







The assumption that transported regional pollution can affect BC_{ff} concentrations in Nicosia can be further supported by investigating the BC_{ff} NWR polar plots for both the cold and warm seasons (Fig. S18a,b). Elevated concentrations related to 123 local emissions were observed for calm conditions with low wind speeds ($<5 \text{ km h}^{-1}$) in the SW sector, as previously observed 124 for HOA-1. Interestingly, BC_{ff} NWR plots show a distinct contribution at higher wind speeds ($\sim15 \text{ km h}^{-1}$) and the NE-SE 125 (Middle East) sector, during both the cold and warm periods, with estimated concentrations of roughly 1.5 µg m⁻³, further 126 support the major role of the Middle East in the observed BC concentration levels in Nicosia (Fig S18 a,b).

727BC source apportionment: In order to better assess the relative contributions of the multiple primary OA sources (HOA-1,728HOA-2) and to quantify the contribution of long-range transport from the Middle East to BCff, a multilinear regression (MLR)729model was tentatively performed using the principle of co-emission of BCff and organic species by the different sources730(Chirico et al., 2010; Laborde et al., 2013). This approach, used recently by Poulain et al. (2021), assumes that at any given731time (t), BCff mass concentration is the sum of BC from traffic (traced by HOA-1), from a mixed combustion source (traced732by HOA-2), and from long-range transport (traced by MO-OOA), as follows:

733

 $[BC]_{\rm ff} = [BC]_{\rm traffic} + [BC]_{\rm mix\ combustion} + [BC]_{\rm regional} (2)$

734 With:

735 736 $[BC]_{traffic} = a x [HOA-1] (3)$

 $[BC]_{regional} = c x [MO-OOA] (5)$

 $[BC]_{mix combustion} = b x [HOA-2] (4)$

737

738 Where a, b, and c are coefficients derived from the multi-linear regression model.

The above approach assumes that primary HOA-1 and HOA-2 can trace $BC_{traffic}$ and $BC_{mix \ combustion}$, respectively. This is 739 740 somewhat expected for traffic which with has a typical HOA-1/BCtraffic ratio with little variations. For HOA-2, this assumption 741 is valid for the fraction that is assumed to originate from power plant emissions, and for some of the cooking activities (e.g., 742 when using charcoal combustion) but not necessarily all. As such, the uncertainties of this approach is are expected to be 743 higher for HOA-27 compared to HOA-1. The use of MO-OOA to trace the regional source of BC would probably lead to even 744 higher uncertainties due to the fact that because MO-OOA is also sensitive to atmospheric photochemical processes and does 745 integrate multiple sources. Nevertheless, this latter assumption is believed to be acceptable given the good agreement reported 746 above between baseline concentrations of BC_{ff} and MO-OOA (Fig. S19), and the above conclusions that carbonaceous aerosols \underline{s} 747 originating from the Middle East are expected to be dominated by fossil fuel combustion. Note that MO-OOA was preferred 748 here to LO-OOA to trace regional emissions due to the latter's somewhat more local character.

Combing equations 2-5 provides the multilinear regression model with the free regression parameters a, b, c_{1} which are fitted to the time-resolved BC_{ff} mass concentration measured by the Aethalometer and PMF results for the ACSM data:

751

 $[BC]_{ff} = a x [HOA-1] + b x [HOA-2] + c x [MO-OOA] (6)$

Previous studies have shown that MLR models seem to have enhanced explanatory power when primary emissions are dominantdominate (Laborde et al., 2013). To reduce this potential bias, the MLR model was applied distinctly for the two seasons separately.

755 During the cold period, a very good correlation between measured and modelled BC_{ff} was obtained ($r^2 = 0.70$; N = 2942), with the modelled BC_{ff} explaining 84 % of the measured one (Fig. S20a). The regression coefficients a (HOA-1), b (HOA-2) and 756 757 c (MO-OOA) were found to be 1.11 ± 0.03 , 0.15 ± 0.01 and 0.41 ± 0.01 , respectively. Regarding the warm period, it was not 758 possible to obtain a positive value for b (HOA-2). A correlation between long-range transported HOA-2 and MO-OOA is, 759 among other, a reason that can be proposed to explain why it has not been possible to extract a BC_{mix source} factor here. Therefore, 760 BC_{ff} was only apportioned using HOA-1 and MO-OOA. A good correlation between measured and modelled BC_{ff} was 761 obtained (r²=0.62; N=1251), with the modelled BC_{ff}, explaining 83% of observations (Fig S20b). The regression coefficients a (HOA-1) and c (MO-OOA) were found to be 3.05 ± 0.07 and 0.19 ± 0.01 , respectively. 762

The combination of the Aethalometer model (apportioning BC_{ff} and BC_{wb}) and the MLR model (apportioning BC_{traffic}, BC_{mix}
 source, and BC_{regional}) was performed to obtain an integrated picture of BC sources in Nicosia for both periods (see Fig. 12).



765

766 Figure 12: BC sources during the cold and the warm period in Nicosia

767 Spatial and seasonal variability of BC sources: During the cold period, BC was found to originate from four different sources 768 denoting the complexity of combustion sources of different origins in Nicosia. BCregional is the dominant source of BC (37%), while traffic, wood burning, and mix source are estimated to contribute to 26 %, 26% and 11% of BC, respectively. From the 769 770 perspective of BC_{ff} sources, long-range transport, traced by MO-OOA, remains the largest source of BC_{ff} during the cold period, contributing 63 %, while BCff from local emissions constrained with HOA-1 and HOA-2 represents 24% and 13%, 771 respectively (Fig S21). In other words, more than half of BC_{ff} in Nicosia was found to be regional and most probably 772 773 originatingprobably originated from the Middle East during the cold period. This high contribution of regional BCff is quite 774 unexpected for a medium-sized European city like Nicosia, where local traffic is likely to be the main contributor to $BC_{\rm fr}$. 775 Nevertheless, extra caution should be taken here, and tThe obtained contribution of 63% for BCff regional should be seen as 776 an upper limit since a fraction of MO-OOA was shown to be of local origin during the cold period. During the warm period, 777 the picture remains similar, with traffic and wood burning- representing two_thirds of BC (56 % & 18 %). Here, BC regional 778 contributed 26 % to total BC. From the perspective of BC_{ff} sources during the warm period, the long-range transport, 779 contributed 41 %, while BC_{ff} from local emissions constrained with HOA-1 represents 59 % (Fig S21). Although the two models (Aethalometer and MLR) are associated with non-negligeable uncertainties, the BC source apportionment obtained, 780

shows that local emissions cannot be considered only for BC, with demonstrated significant contribution of Middle East fossilfuel emissions.

783 4. Conclusions

784 Near-real-time chemical composition of submicron aerosols and source apportionment of carbonaceous aerosols was 785 performed for the first time in Nicosia, a medium-sized European capital city (circa 250,000 inhabitants) in Cyprus located in 786 the Eastern Mediterranean and surrounded by Middle East countries with fast-growing population and increasing emissions 787 of air pollutants. Continuous observations were performed at an urban background site; for approximately 6 months (between 788 7 December 2018 and 31 May 2019), in order to obtain a large and representative dataset capturing specific features _ related 789 to both the cold and warm periods -, such as domestic heating and regional transport. Measurements of the major fractions of 790 PM1 were carried out with a Q-ACSM and an Aethalometer complemented by a comprehensive suite of collocated instruments 791 (e.g., filter sampling, SMPS) to further assess the quality of the acquired data further.

Unlike many European cities, no clear PM₁ pollution episodes of several consecutive days could be observed over Nicosia. However, very intense peaks (above 40 μ g m³_{a, t} 1h averages) were recorded systematically every evening during the cold period. Carbonaceous aerosols (BC and OA) were identified as the main components of these peaks and were mostly attributed to local emissions from heating with only-little contribution from local meteorology (PBL height did not show significant diurnal variability during the cold period). Furthermore, a significant portion of PM₁ was found to be related to long range transported aerosol, while the influence of shipping emissions was estimated to be rather low (less than 8%).

798 Source apportionment of OA has been used to derive a local biomass burning OA (BBOA_{cv}) mass spectrum_t in order to 799 properly apportion the contribution of domestic wood burning properly. A total of five OA sources were identified during the cold period, among which four are typically reported within urban environments (HOA-1, BBOA, LO-OOA, MO-OOA). An 800 801 additional one (HOA-2) was assigned as a mixture of several combustion sources, such as cooking as well as a significant 802 contribution from power plants located in the Northern part of the island. These power plants in addition, represent major 803 island-based hotspots of NOx, as evidenced from by satellite observations. Interestingly, a similar HOA-2 source was identified 804 at our regional background site (40 km distance from Nicosia; Chen et al., 2022), pointing to a possible influence from these 805 power plants to an extended part of the island. The impact of this specific source brings the OA contribution of primary sources 806 up to 40 % over Nicosia during the cold period. Few additional features were noticed for the other OA sources with 1) a typical 807 traffic-related (HOA-1) source observed during both seasons, 2) a biomass burning source (BBOA) related to domestic heating 808 enhanced at night during the cold season and accounting for 12 % of the total OA, 3) a less oxidized secondary (LO-OOA) 809 source of a semi-volatile character, and influenced by local night-time chemistry, that shows wasto be more oxidized (andi.e., 810 of a more regional originless local character) during the warm period, and 4) a secondary (MO-OOA) source mostly of regional 811 origin but also influenced by night-time chemistry during the cold period.

The geographic origin of each OA source was assessed for both seasons. With the exception of Except for MO-OOA, which shows-systematically shows a strong regional component, HOA-1, HOA-2, and LO-OOA exhibit a clear local origin during both seasons, and a more pronounced influence from the Eastern wind sector during the warm period. The prevalence of this sector is systematically observed for MO-OOA highlighting the major role of Middle East emissions in contributing to almosten. half of OA concentrations in Nicosia during both cold and warm seasons.

817 To further elucidate the influence of this complex mixture of OA sources on BC levels, a source apportionment of BC was 818 performed by combining i) the aethalometer model to separate BC into its fossil fuel (BCff) and wood burning components 819 (BC_{wb}) , and ii) a multi-linear regression model to apportion the contribution to BC_{ff} from traffic (constrained by HOA-1), mix 820 combustion sources from cooking and power plants (constrained by HOA-2), and long-range transport from the Middle East 821 (constrained by MO-OOA). Although several assumptions and uncertainties are associated with this approach, it has shown to 822 provide an interesting tool to reconstruction reconstructing the BC concentrations derived experimentally. Such BC 823 apportionment performed for both cold and warm seasons solidified the conclusions reached through the OA source 824 apportionment, with almost half of BCff being of regional origin, with the Middle East playing an important role. This result 825 is quite unexpected given that local traffic emissions are usually considered the dominant contributor to BC_{ff} in urban 826 background environments. These conclusions have numerous implications related to PM regulation and the efficiency of local 827 abatement strategies (in particular regarding traffic emissisions), health (combustion aerosols being considered as particularly 828 adverse for human health), and climate (major influence of light_-absorbing aerosols from the Middle East fossil fuel 829 emissions).

More accurate OA and BC source apportionment i) with more co-located high_resolution measurements of specific trace metal and organic tracers, ii) better resolved OA mass spectra (e.g., from HR-ToF-AMS), iii) the use of various source_specific mass spectra fingerprints (e.g., from cooking or power plants), and iv) multi-site measurements (incl. both urban and regional background) will enable a more accurate estimation of local vs_ regional fossil fuel emissions in Cyprus while better constraining the current regional efforts on air quality modelling and forecasting.

836	Data availability: All data used in this study can be accessed here:
837	https://doi.org/10.5281/zenodo.7802065https://doi.org/10.5281/zenodo.7186341. More details on the analyses are available
838	upon request to the contact author Aliki Christodoulou (a.christodoulou@cyi.ac.cy).
839	
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