

Aerosol light absorption and the role of extremely low volatility organic compounds

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Abstract

A month-long set of summertime measurements in a remote area in the Mediterranean is used to quantify aerosol absorption and the role of black and brown carbon. The suite of instruments included a high-resolution Aerosol Mass Spectrometer (HR-ToF-AMS), and a Scanning Mobility Particle Sizer (SMPS) both coupled to a thermodenuder and an aethalometer, a photoacoustic extinctionsmeter (PAX₄₀₅), a Multi-Angle Absorption Photometer (MAAP), and a Single Particle Soot Photometer (SP2).

The average refractory black carbon (rBC) concentration during the campaign was 0.14 $\mu\text{g m}^{-3}$, representing 3% of the fine aerosol mass. The measured light absorption was two or more times higher than that of fresh black carbon (BC). Mie theory indicated that the absorption enhancement due to the coating of BC cores by non-refractory material could explain only part of this absorption enhancement. The role of brown carbon (BrC) and other non-BC light-absorbing material was then investigated. A good correlation ($R^2=0.76$) between the unexplained absorption and the concentration of extremely low volatility organic compounds (ELVOCs) mass was found.

1. Introduction

Atmospheric aerosol may influence climate in two ways: directly through scattering and absorbing radiation, and indirectly through acting as cloud condensation nuclei (IPCC, 2011) Black carbon is the dominant light absorbing aerosol component. BC is a distinct type of carbonaceous material that is formed mainly during combustion processes. In addition, some

32 organic aerosol (OA) has the ability to absorb sunlight. The OA with strong light absorption is
33 called brown carbon (BrC) (Andreae et al., 2006).

34 The absorption of BC depends on its mixing state (Liu et al., 2015, 2017). Usually, BC is
35 coated with scattering material causing its light absorption to increase due to the lensing effect
36 (Fuller et al., 1999; Jacobson, 2001; Bond et al., 2006; Lack and Cappa, 2010). The absorption
37 enhancement (E_{abs}) is defined as the ratio of the aerosol absorption coefficient (b_{abs}) over the b_{abs}
38 of the pure BC particles. The E_{abs} accounts for the combined effects of lensing and the presence of
39 BrC. This enhancement can also be calculated by the ratio of the equivalent mass absorption cross-
40 sections (MAC). The MAC is defined as the ratio of b_{abs} over the BC mass. The MAC of pure,
41 uncoated BC at a specific wavelength depends both on the size distribution of the particles, but
42 also on their morphology (fresh aggregates versus collapsed more spherical structures). Forestieri
43 et al. (2018) measured the size dependent MAC for particles with a diameter lower than 160 nm.
44 Radney et al. (2014) and Dastanpour et al. (2017) showed that the MAC is proportional to particle
45 mass. Based on Bond et al. (2006) pure BC particles in the 10-350 nm diameter range have an
46 expected $MAC_{550}=3.9-6.8 \text{ m}^2 \text{ g}^{-1}$ and $MAC_{405}=5.3-9.2 \text{ m}^2 \text{ g}^{-1}$. The E_{abs} can be measured by using
47 a thermodenuder (TD) for the removal of the non-refractory coating material from the BC-
48 containing particles or it can be estimated using theoretical models (for example Mie or Rayleigh-
49 Debye-Gans theory) if the size of the primary spherules is known. Incomplete removal of the non-
50 refractory material in the TD can lead to underestimation of the E_{abs} (Healy et al., 2015;
51 McMeeking et al., 2014).

52 Previous studies have demonstrated that lensing has a wide range of effects on the light
53 absorption of BC. Liu et al. (2015) quantified the E_{abs} in a rural area near London during the winter.
54 Using the TD method, they found a campaign-average E_{abs} of 1.3 at 405 nm and 1.4 at 781 nm.
55 They also estimated the E_{ab} combining their measurements with a reference MAC from the
56 literature. At the large wavelength the two methods agreed, but this was not the case at the lower
57 wavelength. In addition, they showed that there was continuous change in E_{abs} with increasing R_{BC} .
58 The R_{BC} was defined as the ratio of the non-rBC mass to the rBC mass in BC-containing particles,
59 measured by the SP-AMS. Finally, this study suggested that the lower volatility BrC had stronger
60 absorption than the semi-volatile BrC. Knox et al. (2009) performed measurements in downtown
61 Toronto, during the wintertime using a TD at 340 °C. They reported an average E_{abs} of 1.43 for
62 fresh particles, based on thermal OC/EC and photoacoustic measurements. Liu et al. (2017)

63 combined laboratory experiments with diesel exhaust emissions and ambient measurements to
64 show that particles with a ratio of R_{BC} less than 1.5 (typical for traffic emissions) had a negligible
65 lensing effect. The R_{BC} was measured by combining an AMS and an SP2. When the R_{BC} was
66 above 3, lensing caused significant enhancement of the absorption of BC. They observed a
67 continuous change in E_{abs} with increasing R_{BC} . Zhang et al. (2018a) presented three years of
68 measurements in a suburban site outside Paris, France, influenced by both fresh and aged air
69 masses. On average they found an $E_{abs}=2.07$ at 370 nm, and an $E_{abs}=1.53$ at 880 nm. They
70 calculated the E_{abs} by measuring the absorption coefficient (b_{abs}) with an aethalometer, while the
71 elemental carbon (EC) was measured with thermal methods using daily filters. Zhang et al. (2018b)
72 presented measurements in Beijing, China during wintertime. Using aethalometer measurements,
73 refractory and non-refractory particle size distributions and Mie theory, they calculated that the
74 lensing effect led to an E_{abs} ranging from 1.5 to 2 on average at 880 nm. Zanatta et al. (2018) found
75 an average E_{abs} equal to 1.54 at 550 nm for measurements at the Zeppelin Arctic Station. Lack et
76 al. (2012) analyzed measurements of biomass burning plumes near Boulder, CO during
77 summertime. Using a TD at 200 °C, they found E_{abs} values as large as 2.5 at 404 nm and 1.7 at
78 532 nm. Using the absorption Angström exponent (AAE) and Mie theory calculations they showed
79 the presence of BrC.

80 Some studies have argued that Mie theory may overestimate the E_{abs} . Cappa et al. (2012)
81 suggested that the absorption enhancement of BC in California in the summertime was low with
82 average values equal to 1.13 at 405 nm and 1.06 at 532 nm for $R_{BC}>10$. For their measurements
83 they used a TD operated at 225-250 °C. Healy et al. (2015) also reported practically no
84 enhancement in the BC absorption at 781 nm and an average $E_{abs}=1.19$ at 405 nm in Toronto,
85 Canada during summertime. During a period associated with wildfires the same authors measured
86 $R_{BC}=6.9$ and an average $E_{abs}=1.39$ at 405 nm. They argued that there was little evidence of the
87 lensing effect, and that BrC was driving the E_{abs} . Cappa et al. (2019) performed measurements in
88 Fresno, CA during wintertime and Fontana, CA during summertime. They found that in Fresno
89 there was absorbing OA, BrC, which was related to biomass burning OA and nitrate-associated
90 OA. In Fresno, they reported average E_{abs} of 1.37, 1.22 and 1.1 at wavelengths equal to 405, 532,
91 781 nm, respectively, for R_{BC} ranging from 1 to 4. In Fontana the E_{abs} was lower with values of
92 1.1 at 405 nm and 1.07 at 532 nm. Laboratory measurements of McMeeking et al. (2014) for
93 biomass burning aerosol indicated higher absorption in lower wavelengths compared to higher

94 ones and thus the presence of BrC. The enhancement of the absorption was negligible at $R_{BC}<10$.
95 On average they found an E_{abs} equal to 1.25 at 781 nm with a maximum value of $E_{abs}=4$ for $R_{BC}>10$.

96 Recent studies have suggested that the absorption efficiency of OA could be related to its
97 volatility. Saleh et al. (2014) in their laboratory biomass burning experiments showed that almost
98 all absorbing OA was associated with extremely low volatility compounds (ELVOCs), with an
99 effective saturation concentration C^* of $10^{-4} \mu\text{g m}^{-3}$. In addition, Saleh et al. (2018) using controlled
100 combustion experiments showed that the absorption activity of BrC is proportional to its molecular
101 size.

102 Despite the significant progress in understanding the absorption of atmospheric fine aerosol
103 there are still remaining questions regarding both the absorption enhancement of black carbon and
104 the absorption of OA as the aerosol evolves in the atmosphere. In this study we try to address these
105 issues for aerosol that has been aged in the atmosphere for at least a few days before arriving at
106 the island of Crete in the Eastern Mediterranean.

107

108 **2. Experimental Methods**

109 A remote location in the Eastern Mediterranean was used for the study of the absorption
110 and volatility of aged carbonaceous aerosol. The area is characterized by intense photochemistry,
111 especially during the summer (Pikridas et al., 2010) and is affected by pollutants transferred from
112 continental Europe, Turkey, Greece and Africa (Mihalopoulos et al., 1997; Lelieveld et al., 2002;
113 Kalivitis et al., 2011; Bougiatioti et al., 2014). Previous measurements have shown that the OA
114 reaching the area is highly oxidized regardless of its origin (Hidlebrandt et al., 2010; 2011). Lee
115 et al. (2010) showed that these oxidized organic compounds have much lower volatility than fresh
116 SOA. Long-term measurements in the region have revealed relatively high light absorption and
117 scattering by aerosol during the summer (Kaliivitis et al., 2011; Vrekoussis et al., 2005).

118 The FAME-16 field campaign took place from May 9 to June 2, 2016. Measurements were
119 conducted at the Finokalia Station ($35^{\circ} 20' N$, $25^{\circ} 40' E$, 250 m asl), a remote site on the island of
120 Crete in Greece (Mihalopoulos et al., 1997). The nearest large city is Heraklion with 150,000
121 inhabitants located 50 km west of Finokalia (Kouvarakis et al., 2000). There are no local sources
122 near the station, allowing the investigation of aged OA from different source regions. During this
123 study, two Saharan dust events occurred from May 12 till May 15 and May 21 till May 22.

124 A Scanning Mobility Particle Sizer (SMPS, TSI classifier model 3080, CPC model 3775)
125 was used to measure the number and the size distribution of the particles. The aerosol flow was
126 set at 1 L min⁻¹ and the sheath flow at 5 L min⁻¹. The sampling time was 3 min.

127 The mass concentration and the chemical composition of the particles were monitored
128 using a High-Resolution Time-of-Flight Aerosol Mass Spectrometer (HR-ToF-AMS, Aerodyne
129 Research, Inc.). SQUIRREL 1.56D and PIKA v1.15D were used for the data analysis, while for
130 the elemental ratio calculations the improved ambient calculation approach of Canagaratna et al.
131 (2015) was used. The HR-ToF-AMS was operated in V-mode with a sample time of 3 min. The
132 collection efficiency of the HR-ToF-AMS was calculated using the algorithm of Kostenidou et al.
133 (2007). The average CE was 0.64±0.2. Positive matrix factorization (PMF) analysis (Lanz et al.,
134 2007; Paatero and Tapper, 1994; Ulbrich et al., 2009) was performed using as input the high
135 resolution OA mass spectra and the mass-to-charge ratios (m/z) from 12 to 200.

136 A Single Particle Soot Photometer (SP2, Droplet Measurement Technologies) was used to
137 measure the BC size distribution and concentration. The SP2 had 8-channels and included a 1064
138 nm Nd:YAG laser operating at 4 V and 3400 A. The instrument was calibrated using fullerene
139 soot (Alfa Aesar, stock 40971, lot L20W054, SSA=0.4) (Gysel et al., 2011). The calibration of the
140 SP2 was verified in separate experiments using a centrifugal particle mass analyzer (CPMA,
141 Cambustion). The SP2 mode mass and the CPMA mode mass were in good agreement with an
142 $R^2=0.99$ (Saliba et al., 2016). The data were analyzed using the Probe Analysis Package for Igor.
143 In our measurements, the number concentration of BC was low (<10,000 particles cm⁻³), and thus
144 it was assumed that there were no coincidence artifacts in our measurements. The scattering
145 measurement was calibrated using monodisperse polystyrene latex (PSL) spheres. The BC number
146 concentration distributions measured by the SP2 were fitted using a log-normal distribution to
147 account for particles smaller than the SP2 detection limit of approximately 50 nm (Ditas et al.,
148 2018). This extrapolation resulted in an increase of the BC mass concentration by 3-8% (Figure
149 S9), therefore the uncertainty introduced by BC outside the measurement window was minor.

150 A photoacoustic extinctionsmeter (PAX, Droplet Measurement Techniques) with a blue (405
151 nm) laser was used to measure the absorption (b_{abs}) and the scattering (b_{scat}) coefficients. Fullerene
152 soot and PSL spheres were used to calibrate the absorption and scattering signals, respectively. An
153 activated carbon denuder was placed in front of the PAX to remove NO₂. The b_{abs} measurement
154 by the PAX has an uncertainty of less than 10% (Nakayama et al., 2015). Furthermore, a seven

155 wavelength aethalometer (AE31, Magee Scientific) was used to measure the b_{abs} at 370, 470, 520,
156 590, 660, 880 and 950 nm and to calculate the absorption Angstrom exponent (AAE), which
157 describes the wavelength dependence of the b_{abs} . The aethalometer measurements were corrected
158 for scattering and multiple scattering artifacts following Saleh et al. (2014) and Tasoglou et al.
159 (2017) using the corrections suggested by Weingartner et al. (2003) and Kirchstetter and Novakov
160 (2007). High relative humidity (>70%) can introduce measurement artifacts in the measurement
161 absorption coefficient by filter- based techniques or photoacoustic methods (Arnott et al. 2003). A
162 diffusion drier was used upstream of the optical measurements. The campaign average temperature
163 and relative humidity were 22 ± 4 °C and 53 ± 19 %, respectively. The measurement station had a
164 temperature-control system maintaining the temperature at approximately 25 °C.

165 The thermodenuder (TD) used in this study, was placed upstream of the HR-ToF-AMS and
166 the SMPS. The TD design was similar to that developed by An et al. (2007) and is described by
167 Louvaris et al. (2017). The TD was operated at temperatures ranging from 25 °C to 400 °C using
168 several temperature steps from 25 °C to 200 °C over several hours and then rapidly (in 20 min)
169 increasing its temperature to the 375-400 °C to investigate the presence of ELVOCs. One complete
170 cycle from 25 to 400 °C and back to 25 °C lasted approximately 10 h. Sampling was alternated
171 between the ambient line and the TD line every 3 minutes using computer-controlled valves.
172 Changes in particle mass concentration, composition, and size due to evaporation in the TD were
173 measured by the HR-ToF-AMS and the SMPS resulting in thermograms of OA mass fraction
174 remaining (MFR) as a function of TD temperature. The OA MFR was calculated as the ratio of
175 organic mass concentration of a sample passing through the TD at time t_i over the average mass
176 concentration of the ambient samples that passed through the bypass line at times t_{i-1} and t_{i+1} . The
177 sample residence time in the centerline of the TD was 14 s at 25 °C, corresponding to an average
178 residence time in the TD of 28 s. The MFR values were corrected for particle losses in the TD due
179 to diffusion and thermophoresis. To account for these losses, sample flow rate as well as size- and
180 temperature-dependent loss corrections were applied following Louvaris et al. (2017)
181 corresponding to the operating conditions during the campaign. Less than 20% of the particulate
182 matter was lost in the TD at temperatures up to 100°C. The losses increased at higher temperatures.
183 and at 400 °C approximately 50% of particles larger than 50 nm was lost. The uncertainty
184 introduced by the loss correction was approximately 20% (Gkatzelis et al., 2016; Louvaris et al.,
185 2017). The final step of the data analysis was to average the corrected for CE and TD losses MFR

186 data based on temperature bins of 10°C. The MFR calculation assumes implicitly that the OA
187 concentration remains constant during the measurement period. To ensure that this condition is
188 satisfied, if two consecutive OA ambient mass concentrations differed by more than 25%, the
189 corresponding MFR was not included in the analysis. Also, in order to ensure that the temperature
190 was constant during the measurement, the absolute difference between the two samples had to be
191 less than 5°C. If this difference for a TD sample was higher, then the sample was not included in
192 our analysis. The same approach was used also for the factors resulting from the PMF analysis of
193 the AMS spectra. However, in this case a minimum concentration threshold of 0.1 $\mu\text{g m}^{-3}$ was used
194 for the ambient concentrations together with the criterion of the stability of the ambient
195 concentrations during the sampling period. MFR values corresponding to concentrations of the
196 PMF factors below this threshold were not included in the dataset. Approximately 75% of the OA
197 samples satisfied all these constraints and were used in the analysis. The corresponding
198 percentages were 65% and 70% for the two identified PMF factors. In the present work the
199 complete datasets will be analyzed together, averaging the corresponding measurements. More
200 details regarding the data analysis and the sensitivity tests of the TD measurements are provided
201 in the supplementary information.

202 The concentrations of gas-phase pollutants were measured using a Proton-Transfer
203 Reaction Mass Spectrometer (PTR-QMS 500, Ionicon Analytik) and gas monitors. The PTR-MS
204 was calibrated with a standard gas mixture of VOCs. The concentration of O₃ was measured using
205 a continuous O₃ analyzer (Thermo Scientific, 49i) and the concentrations of nitrogen oxides were
206 measured using a NO/NO₂/NO_x analyzer (Thermo Scientific, 42i-TL).

207

208 **3. Theoretical Analysis Methods**

209 The dynamic TD evaporation model of Riipinen et al. (2010) together with the uncertainty
210 estimation algorithm of Karnezi et al. (2014) were used for the determination of the OA volatility
211 distribution. Inputs for the model included the ambient OA concentration, the OA density
212 calculated by the algorithm proposed by Kostenidou et al. (2007), the initial average particle size,
213 TD temperature, the MFR values, and TD residence time. In the volatility basis set framework of
214 Donahue et al. (2006), the volatility distribution is represented with a range of logarithmically
215 spaced C* bins along a volatility axis. In this study 6 bins with variable mass fractions were chosen.
216 For this 6-bin solution, the best 2% of the mass fraction combinations with the lowest error were

217 used to estimate the average mass fraction along with their corresponding standard deviation for
218 the uncertainty of each bin. Additionally the parameters that affect indirectly the calculated
219 volatility such as the effective vaporization enthalpy (ΔH_{vap}), and the effective accommodation
220 coefficient (a_m) were estimated following Karnezi et al. (2014). Additional information regarding
221 the data analysis of the TD measurements are in the supplemental information (sections S1 and
222 S2).

223 A Mie theory model based on the work of Bohren and Huffman (1983) was used to
224 calculate the theoretical MAC and E_{abs} at 405 nm assuming a spherical core-shell morphology
225 (China et al., 2015). The assumption that all BC particles were coated and had obtained a relatively
226 spherical shape can be justified by the lack of any local sources and the fact that all particles
227 reaching the site had been heavily processed in the photochemically active summertime
228 atmosphere of the Eastern Mediterranean. PMF analysis of the OA AMS spectra did not show any
229 fresh emissions like HOA or BBOA and no acetonitrile was detected. FLEXPART analysis
230 confirmed that the air masses measured were transferred from long distance areas (30% continental
231 Greece and the Balkans, 13% Aegean, 24% Africa, 33% Italy/Sicily). The measurements
232 suggested a positive correlation ($R=0.31$) between the MAC_{405} and the ratio of non-refractory PM_{10}
233 to rBC with an intercept of $12.1 \text{ m}^2 \text{ g}^{-1}$ (Figure S10).

234 For the BC core we assumed a refractive index of the core $n_{\text{rBC}}=1.85+0.71i$ (Bond et al.,
235 2006) and a density of 1.8 g cm^{-3} (Mullins and Williams, 1987; Park et al., 2004; Wu et al., 1997).
236 A non-absorbing coating of the BC core was assumed, with a refractive index of $n_{\text{OA}} = 1.55$ (Bond
237 and Bergstrom, 2006). The total aerosol effective density used was calculated based on the SMPS
238 and HR-ToF-AMS distributions. The average effective density for the campaign was $1.66 \pm 0.11 \text{ g}$
239 cm^{-3} . The size distribution of the BC cores was provided by the SP2 and the corresponding
240 measured rBC size distributions including the extrapolation to smaller sizes outside the SP2
241 measurement window.

242 The coating thickness in the base case calculation was estimated based on the assumption
243 that the BC material is internally mixed with the non-refractory aerosol species (Saliba et al.,
244 2016). The SMPS size distributions were used to estimate the ratio of the total aerosol mass over
245 the BC mass as measured. 1-hour averaged data were used as inputs in the model. In order to
246 provide a better constraint for the analysis, the average coating thickness for each period of the
247 corresponding mean BC cores ($\pm 5 \text{ nm}$) was estimated using the leading-edge-only (LEO) fit

248 method (Gao et al. 2007) and the SP2 data. The thicknesses as expected were lower than those
249 resulting from the internal mixing assumption, but were still significant (Figure S11). Mie theory
250 calculations using the coating thickness based on the LEO fit method were also performed.

251

252 **4. Results and discussion**

253 The average rBC concentration of the campaign was $0.14 \mu\text{g m}^{-3}$ and the average OA
254 concentration was $1.5 \mu\text{g m}^{-3}$ (Figure 1). The two major Saharan dust events affected, as expected,
255 the aerosol optical properties. In the present study we focus only on the non-dust periods. The
256 dominant PM_{10} components were sulfate and OA, accounting for 46% and 34% of the PM_{10} ,
257 respectively. The O:C ranged from 0.65 to 1, with an average value of 0.83, revealing the absence
258 of fresh OA. These values are typical in Finokalia during the spring and summer periods
259 (Hidlebrandt et al., 2010).

260

261 **4.1 OA Volatility**

262 The estimated volatility distribution for the total OA in Finokalia during FAME-16 is
263 depicted in Figure 2. Use of OA with $C^* = 10^{-8} \mu\text{g m}^{-3}$ was needed to capture the behavior of the
264 OA at 400°C . Almost 40% of the OA consisted of semi-volatile organic compounds (SVOCs),
265 35% of low volatility organic compounds (LVOCs), and the rest was extremely low volatility
266 organic compounds (ELVOCs).

267 The estimated value of the effective vaporization enthalpy was $80 \pm 20 \text{ kJ mol}^{-1}$. This value
268 was in agreement with the reported value by Lee et al. (2010) of 80 kJ mol^{-1} for the FAME-08
269 campaign. The estimated accommodation coefficient was 0.27, ranging from 0.1 to 0.8. This value
270 was a little higher than the 0.05 value reported in the earlier study. However, both suggest only
271 moderate resistances to mass transfer during the evaporation in the TD.

272 The corresponding measured and predicted thermograms are depicted in Figure 3. Almost
273 30% of the OA had not evaporated even after heating at 400°C . The temperature at which half of
274 the OA evaporated was $T_{50}=120^\circ\text{C}$, a value similar to that observed by Lee et al. (2010). The
275 composition of the OA leaving the TD changed significantly as temperature increased according
276 to the model. At 125°C the LVOCs and ELVOCs contributed equally to the remaining OA mass.
277 For further temperature increases the LVOC fraction was reduced until 375°C , at which point only
278 the ELVOCs remained.

279 PMF analysis resulted in a two-factor solution (Florou et al., in prep.). Factor 1
280 corresponded to more oxidized oxygenated OA (MO-OOA) and Factor 2 to a less oxidized
281 component (LO-OOA). The average contribution of the two factors was 47% for the MO-OOA
282 and 53% for the LO-OOA. The O:C for the MO-OOA was 0.95 ($OS_C = 0.59$) and for the LO-
283 OOA it was 0.56 ($OS_C = -0.27$). There was a weak positive correlation between LO-OOA, MO-
284 OOA and rBC. The R^2 between the hourly concentrations of MO-OOA and rBC was 0.12 (Figure
285 S12), and for LO-OOA and rBC 0.17 (Figure S13). This is not unexpected given that Finokalia is
286 far away from the corresponding sources of both BC and organic compounds and significant
287 physical and chemical processing has taken place during the transport of the aerosol from the
288 sources to the receptor.

289 The OA in the Eastern Mediterranean during the summer has significant contributions from
290 both anthropogenic and biogenic sources but their contributions remain uncertain. Based on our
291 recent work (Drosatou et al., 2019) the LO-OOA and MO-OOA do not reflect different sources,
292 but rather different degrees of chemical aging. The Finokalia area is characterized by the absence
293 of local sources and as result the absence of fresh OA. The area is characterized by dry land and
294 dry vegetation (dry bushes), thus no high biogenic emissions are expected. The OA during FAME-
295 08, was also found to consist entirely of OOA with no primary OA (POA) present (Hildebrandt et
296 al., 2010). POA evaporates and gets oxidized rapidly in the photochemically active environment
297 of the Eastern Mediterranean during its transport from its sources to this remote site. FLEXPART
298 simulations during the measurement campaign revealed that the air masses arriving at the site
299 during the study were originating from continental Greece and the Balkans (30% of the time),
300 Aegean (13% of the time), Africa (24%), and Italy/Sicily (33%).

301 The volatility distributions of the two PMF factors were estimated following the same
302 approach as that for the total OA. The measured thermograms for the two PMF factors are shown
303 in Figure 4. Almost 50% of both the MO-OOA and the LO-OOA evaporated at 150 °C. Almost
304 30% of the MO-OOA mass, and about 20% of the LO-OOA did not evaporate even at temperatures
305 as high as 400°C. The model reproduced the observed MFR values for both factors. Both factors
306 contained components with a wide volatility range. The MO-OOA exhibited a bimodal volatility
307 distribution with peaks at effective saturation concentrations of 10^{-8} and $10 \mu\text{g m}^{-3}$. Its effective
308 enthalpy of vaporization was approximately $90 \pm 35 \text{ kJ mol}^{-1}$ and its accommodation coefficient
309 was 0.27. The estimated volatility distribution of the LO-OOA was a little more uniform peaking

310 at an effective saturation concentration of $1 \mu\text{g m}^{-3}$. The average calculated saturation
311 concentration of LO-OOA at 298 K was $0.016 \mu\text{g m}^{-3}$, an order of magnitude higher than that of
312 the MO-OOA. The LO-OOA enthalpy of vaporization was $70 \pm 20 \text{ kJ mol}^{-1}$, 20 kJ mol^{-1} lower
313 than that of the MO-OOA. Its accommodation coefficient was approximately 0.1 indicating small
314 mass transfer resistances. MO-OOA consisted of approximately 40% SVOCs, 30% LVOCs, and
315 30% ELVOCs. On the contrary, LO-OOA consisted of almost 45% SVOCs, 40% LVOCs and only
316 15% ELVOCs. ELVOCs can be produced both by primary sources (combustion of fossil fuels but
317 also biomass burning) and secondary processes. Given the intense chemical processing of the
318 organic compounds from all sources in their way to Finokalia, we cannot assign based on our
319 measurements to a specific source or process.

320 The fitting of the individual factor thermograms implicitly assumes that each factor had the
321 same size distribution as the total OA and also that the two factors were externally mixed. The
322 uncertainty introduced by these two assumptions was implicitly evaluated comparing the estimated
323 total OA of volatility distribution with the composition-weighted average of the volatility
324 distributions of the two OA factors. The two distributions agreed within a few percent for $10^{-3} < C^* <$
325 10^0 and within 10% for the lowest and highest volatility bins.

326

327 4.2 Aerosol optical properties

328 The PAX was used to measure the b_{abs} and b_{scat} at 405 nm. The measured $b_{scat,405}$ ranged
329 from 11.5 to 47 Mm^{-1} with an average campaign value of 26.5 Mm^{-1} (Figure 5). The corresponding
330 $b_{abs,405}$ ranged from 0.64 to 5.6 Mm^{-1} with an average value of 2.1 Mm^{-1} .

331 The absorption coefficients measured by the aethalometer at $\lambda=370$ nm and at $\lambda=470$ nm,
332 were compared with the absorption coefficient measured by the PAX at $\lambda=405$ nm. The
333 measurements of the two instruments were highly correlated with $R^2=0.9-0.91$. The $b_{abs,370}$
334 measured by the aethalometer was higher than the $b_{abs,405}$ of the PAX. Their relationship is
335 described by the equation $y=3.32x - 3.24$. Similarly, the $b_{abs,450}$ measured by the aethalometer was
336 higher than that of the PAX at 405 nm and their relationship was described by $y=1.85x - 0.93$
337 (Figure S14). Part of these differences are due to the artifacts associated with filter-based
338 absorption measurements (Cappa et al., 2008; Lack et al., 2008).

339 Acetonitrile is a known biomass burning marker and can help identify the potential
340 influence of the site by biomass burning events or wildfires during the campaign. The acetonitrile

341 concentration measured by the PTR-MS remained close to 0.4 ppb during the campaign, which is
342 the local background level. This together with the low BC levels indicate that the site was not
343 impacted by nearby biomass burning during the study.

344 The $b_{abs,405}$ variation followed that of the rBC ($R^2=0.74$ for the hourly averages). The R^2 at
345 $\lambda=370$ nm was lower at 0.67 than that at 405 nm. This is consistent with the presence of some BrC.
346 However, the $b_{abs,370}$ is measured by the aethalometer and the $b_{abs,405}$ by the PAX. For the b_{abs} at
347 higher wavelengths measured by the aethalometer the R^2 at $\lambda=450$ nm was 0.68, at $\lambda=520$ nm 0.67,
348 at $\lambda=590$ nm 0.65, at $\lambda=660$ nm 0.65, and at $\lambda=950$ nm 0.51 (Figure S15). These values are
349 inconclusive regarding the existence or absence of BrC. Part of the explanation for this behavior
350 could be that a fraction of the BrC is associated with rBC either from the emissions or from the
351 associated gas-phase pollutants that react in their way to the site.

352 The ratio of the $b_{abs,405}$, measured by the PAX, over the rBC mass, was equal to 16 ± 2.2 m²
353 g⁻¹. In this study the average rBC size distribution had a number mode diameter of 67 nm and a
354 mass mode diameter equal to 185 nm. The measured MAC₄₀₅ is clearly higher than the expected
355 MAC₄₀₅ of uncoated BC particles (Bond et al., 2006). The difference between the measured and
356 the reference value of MAC₄₀₅ can be due to the coating of BC by other PM components (lensing
357 effect) and/or the existence of other absorbing material. These two potential explanations will be
358 explored in the following paragraphs.

359 The AAE of the aerosol was calculated using a power-law fitting of the b_{abs} measured by
360 the aethalometer in all seven wavelengths (370, 470, 520, 590, 660, 880, 950 nm). The campaign
361 average AAE was 0.97 ± 0.22 . Lack and Cappa (2010) suggested that an AAE>1.6 should confirm
362 the presence of non-BC absorbing material, however an AAE<1.6 does not exclude its presence.
363 The AAE of coated BC cores can deviate from the typical AAE=1 with values greater or lower
364 than unity. Gyawali et al. (2009) have shown using a core-shell model and assuming spherical
365 particles that for relatively small cores, the coating thickness increase leads to the increase of the
366 AAE while for large cores, AAE is not affected by the changes of the coating thickness. Liu et al.
367 (2018) showed that the AAE of coated BC is highly sensitive to the particle size distribution and
368 demonstrated that AAE decreases as particle size increases. They showed that an AAE as low 0.8
369 is possible and demonstrated the importance of various parameters on the BC AAE and the
370 potential problems introduced by assuming BC AAE as being equal to 1.0. Based on the relatively
371 large aged particles present in Finokalia relatively low values of AAE should be expected.

372 According to simulation presented by Gyawali et al. (2009), the average AAE for this study can
373 range from 1 to 1.25, based on the core diameter measured and the estimated coating thickness.
374 Therefore, the relatively low average AAE does not preclude the presence of some absorbing
375 organic aerosol.

376 Mie theory calculations were performed in order to estimate the MAC_{405} and the E_{abs} due
377 to the lensing effect of the shell covering the BC core. In our base case the coating thickness was
378 estimated based on the ratio of the total aerosol mass over the BC mass (Saliba et al., 2016).
379 Initially, a non-absorbing shell was assumed. The predicted MAC_{405} had an average value of 15.8
380 $m^2 g^{-1}$. The average E_{abs} due to the lensing effect was 2.13. The predicted average MAC_{405} using
381 the measured BC size distribution and assuming pure uncoated spherical particles was $7.4 m^2 g^{-1}$
382 varying from 6.1 to $7.8 m^2 g^{-1}$ during the study (hourly averages). This is consistent with the
383 expected 5.3-9.2 $m^2 g^{-1}$ for BC core diameters in the 10-350 nm size range (Bond et al. 2006). The
384 average predicted MAC_{405} was lower than the average measured $MAC_{405} = 16 m^2 g^{-1}$ but the p-
385 value was equal to 0.079. Taking into account the measurement uncertainty, the measured average
386 MAC_{405} could range from 14.4 to $17.6 m^2 g^{-1}$. However, here were several periods during which
387 the measured MAC was much higher than the predicted values with the differences as high as 8.5
388 $m^2 g^{-1}$ (Figure 6). These differences suggest the average contribution of BrC to absorption during
389 the measurement period was relatively small, but there were periods with high levels of absorbing
390 OA.

391 In the next step, the Mie theory calculations were repeated assuming an absorbing shell
392 with a refractive index of $n_{OA} = 1.55 + ki$, where the imaginary part, k , was allowed to vary from
393 0 to 0.4. This range of k values was selected based on previous literature (Kirchstetter et al., 2004;
394 Alexander et al., 2008; Chakrabarty et al., 2010; Chen and Bond, 2010; Saleh et al., 2014;
395 Chakrabarty et al., 2016, Li et al., 2016; Saleh et al., 2018). Approximately half of the resulting k
396 values during the campaign were zero, suggesting a non-absorbing shell, while the other half were
397 were positive. More specifically, 24% of the estimated k values ranged from 0.01 to 0.1, 8% from
398 0.11 to 0.2, 3% from 0.21 to 0.3, and 10% of the k values ranged from 0.31 to 0.4. During these
399 periods, the campaign average $b_{abs,405}$ was equal to $2.4 Mm^{-1}$. BrC was estimated to lead to a 15%
400 increase of the campaign average $b_{abs,405}$.

401 The Mie calculations were repeated using a coating thickness estimated by the LEO fit
402 method and the SP2 data. The average coating thickness calculated by this approach was

403 approximately half of that calculated using the internal mixture assumption (Figure S11). This
404 resulted in lower predicted absorption and therefore a larger gap between measurements and
405 predictions assuming that the organic aerosol was not absorbing. The predicted MAC_{405} had an
406 average value of $14.4 \text{ m}^2 \text{ g}^{-1}$. The predicted average MAC_{405} assuming pure uncoated spherical
407 particles was $7.4 \text{ m}^2 \text{ g}^{-1}$. The E_{abs} was equal to 1.94. Assuming an absorbing shell, we found that
408 the imaginary part of the refractive index of the shell was 20% of the times in the range of 0.01 to
409 0.1, 16% between 0.11 to 0.2, 16% between 0.21 to 0.3, and 20% between 0.31 to 0.4. Only 28%
410 of the times the k was equal to 0 indicating no absorbing shell.

411 For both methods used to estimate the MAC_{405} it is noted that there was presence of
412 absorbing material in 45% up to 72% of the measurements.

413

414 **4.3 The role of ELVOCs**

415 ELVOCs can exhibit substantial larger light absorption than LVOCS or SVOCs. The
416 association of photochemically aged BrC with material of lower volatility has been reported in a
417 number of studies focusing on biomass burning (Saleh et al., 2014; Wong et al., 2019). The
418 hypothesis that the presence of ELVOCs could explain the higher aerosol light absorption was
419 tested. The unexplained MAC (ΔMAC) difference of measured and predicted values was
420 compared with the total ELVOC mass concentration. The ELVOC concentration was estimated
421 based on the results on the volatility analysis of the two PMF factors:

$$422 \quad [ELVOC] = 0.15 [LO-OOA] + 0.3 [MO-OOA]$$

423 The predicted MAC_{405} used was from the base case Mie theory calculations in which the coating
424 thickness was estimated from BC mass fraction. The unexplained b_{abs} at 405 nm (3 h average
425 values) was well correlated with the estimated ELVOC concentrations with an $R^2=0.76$ (Figure
426 7). Even after excluding the period with the highest unexplained absorption, the relationship is still
427 relatively strong with $R^2=0.41$. The corresponding R^2 between the 3-h average ΔMAC and the
428 ELVOCs was 0.66 (Figure S16). These results suggest that the ELVOCs were probably
429 contributing to the total absorption and could explain the difference in the MAC.

430 Correlation of the ΔMAC with the rBC, and with other parameters were lower than those
431 for the ELVOCs. For example, the R^2 with the rBC was 0.38, with the LO-OOA 0.44 and with the
432 MO-OOA 0.29. There was a relatively high correlation with the sulfate levels ($R^2=0.59$) that could
433 be interesting as high sulfate levels in this area correspond to high aerosol acidity which has been

434 shown to promote formation of oligomers in secondary organic aerosol. A comparison of the
435 difference of measured and the predicted $b_{abs,405}$ and the ELVOC was also conducted for all the
436 periods including those in which no BrC was present ($k=0$). The correlation was $R^2 = 0.52$. During
437 the $k=0$ periods the predicted $b_{abs,405}$ was equal or higher than the measured $b_{abs,405}$. This difference
438 was due to overestimation of the absorption values by the Mie theory or due to the uncertainties in
439 the measurements of the $b_{abs,405}$.

440 The calculations were repeated using the predicted MAC based on the Mie theory assuming
441 a coating thickness according to the LEO fit. The correlation between the unexplained absorption
442 and the ELVOC concentration remained high with $R^2=0.69$ (Figure S17), suggesting that despite
443 the uncertainty in the coating thickness our results are quite robust.

444

445 5. Sensitivity analysis

446 The analysis presented in the previous sections has been based on a series of assumptions
447 and, as expected is affected, by measurement uncertainties. We have tested the robustness of our
448 conclusion about the link between the unexplained absorption and ELVOC levels by repeating the
449 analysis for several cases. We focused on the estimation of the coating thickness of the BC
450 particles, the refractive index of black carbon, the uncertainty of the measurements by the SP2 and
451 PAX₄₀₅, the uncertainty of the results from the PMF analysis and finally the uncertainty of the
452 thermodenuder model. The results of the corresponding tests are summarized in the following
453 paragraphs.

454 The effect of the assumed BC refractive index on our results was tested by repeating the
455 Mie theory calculations for two additional values: a relatively high value of $1.95+0.79i$ and a
456 relatively low value $1.5 + 0.5i$ (Bond and Bergstrom, 2006). For the high refractive index, the
457 predicted average MAC_{405} using the measured BC size distribution and assuming pure uncoated
458 spherical particles was $7.7 \text{ m}^2 \text{ g}^{-1}$ varying from 6.3 to $8.2 \text{ m}^2 \text{ g}^{-1}$ during the study (hourly averages).
459 Even if the unexplained absorption was reduced its R^2 with the ELVOCs remained high and equal
460 to 0.72 (Fig. S18). For the low refractive index, the predicted average MAC_{405} of the uncoated BC
461 particles was $6.2 \text{ m}^2 \text{ g}^{-1}$ (range of 5.2 - $6.5 \text{ m}^2 \text{ g}^{-1}$ during the study). The R^2 between the unexplained
462 absorption, $\Delta b_{abs,405}$ and the ELVOC concentration was $R^2=0.55$ (Fig. S19). The link between the
463 unexplained absorption and the abundance of ELVOCs is quite robust with respect to the assumed
464 value of the BC refractive index.

465 We estimated that 92-97% of the BC mass concentration was inside the SP2 measurement
466 window therefore given that we also corrected for it fitting the measured size distribution, the
467 uncertainty introduced by this limitation of the SP2 was minor. Nakayama et al. (2015) reported
468 an uncertainty of the b_{abs} measurements by the PAX of less than 10%. Both of these uncertainties
469 did not have an important effect on the link between the unexplained absorption and the ELVOC
470 concentrations.

471 An assessment of the uncertainty of the concentrations of the two OA factors determined
472 by the PMF analysis was performed by bootstrapping 10 simulations (Ulbrich et al., 2009). The
473 estimated uncertainty for the LO-OOA concentrations was 2% and for the MO-OOA
474 concentrations was 3% (Fig. S20). These relatively small uncertainties suggest that the PMF
475 uncertainty regarding the determination of these factors does not affect significantly the
476 conclusions of this study.

477 The uncertainty related to the volatility distributions determined by the thermodenuder
478 results was assessed by estimating low and high limits of the ELVOC concentrations: $ELVOC_{low}$
479 and $ELVOC_{high}$. These were estimated based on the extreme mass fractions that were calculated
480 during the sensitivity analysis of the TD model (Supplemental information Section S2). The LO-
481 OOA ELVOC mass fraction ranged from 0.09 to 0.25 while that of the MO-OOA from 0.13 to
482 0.47. For the low ELVOC case we thus assumed that:

$$483 \quad [ELVOC_{low}] = 0.09 [LO-OOA] + 0.13 [MO-OOA]$$

484 The R^2 between the unexplained $b_{abs,405}$ and $ELVOC_{low}$ was $R^2=0.79$ (Fig. S21). For the high
485 ELVOC case:

$$486 \quad [ELVOC_{high}] = 0.25 [LO-OOA] + 0.47 [MO-OOA]$$

487 Once more the correlation between the unexplained absorption at 405 nm and the ELVOCs was
488 quite high with $R^2=0.78$ (Fig. S22).

489

490 6. Conclusions

491 A month-long campaign was conducted at a remote site, in Finokalia, Crete during May of
492 2016. The dominant PM_1 components were sulfate and aged organics with $O/C=0.81$. The average
493 ambient OA concentration was $1.5 \mu g m^{-3}$ and the rBC was $0.14 \mu g m^{-3}$. Continuous monitoring of
494 biomass burning markers revealed that there were no periods of enhanced biomass burning
495 influence on the site during the campaign. PMF analysis resulted in two secondary OA factors:

496 one more oxidized (MO-OOA) and one less oxidized (LO-OOA). Total OA consisted on average
497 of 40% SVOCs, 35% LVOCs and 25% ELVOCs. Both OA components with a wide range of
498 volatilities. Approximately 30% of the MO-OOA was ELVOCs, 30% LVOCs and 40% semi-
499 volatile material. The LO-OOA was more volatile on average with 40% consisting of LVOCs and
500 45% of SVOCs and 15% of ELVOCs.

501 Aerosol optical properties were measured. The average b_{scat} at 405 nm was 26.5 Mm^{-1} and
502 the average b_{abs} at 405 nm was 2.1 Mm^{-1} . Furthermore, the average AAE was 0.97 and the MAC_{405}
503 was 15.7. Mie theory calculations were able to reproduce less than half of the measured MAC_{405}
504 values assuming core-shell morphology and a non-absorbing shell ($k=0$). We estimated that the
505 non-absorbing shell was causing an enhancement of the absorption by a factor of 2.1. For the other
506 half of the measurements the presence of an absorbing shell with an average k of 0.18 was needed
507 to explain the measurements.

508 The ELVOCs mass concentration was estimated using the volatility distributions of the
509 two factors. The ELVOC concentration was highly correlated with the unexplained MAC_{405}
510 ($R^2=0.66$) and the unexplained $b_{abs,405}$ ($R^2=0.76$), defined as the differences of the parameters
511 measured and the ones predicted by Mie theory for $k=0$. These results suggest that the unexplained
512 absorption in this remote site could be due to a large extent to the extremely low volatility
513 components of the organic aerosol.

514
515 *Data availability.* The data in the study are available from the authors upon request
516 (spyros@chemeng.upatras.gr).

517
518 *Author contributions.* AT conducted the absorption measurements, analysed the results and wrote
519 the paper. EL performed the thermodenuder measurements and analysed the results. KF performed
520 the AMS measurements and analysed the results. AL performed the PTR-MS measurements and
521 analysis. EK was responsible for the OA volatility analysis. CK coordinated the field campaign
522 and assisted with all measurements. NW assisted with all measurements SNP was responsible for
523 the design and coordination of the study and the synthesis of the results. All co-authors contributed
524 to the writing of the manuscript.

525
526 *Competing interests.* The authors declare that they have no conflict of interest.

527

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538

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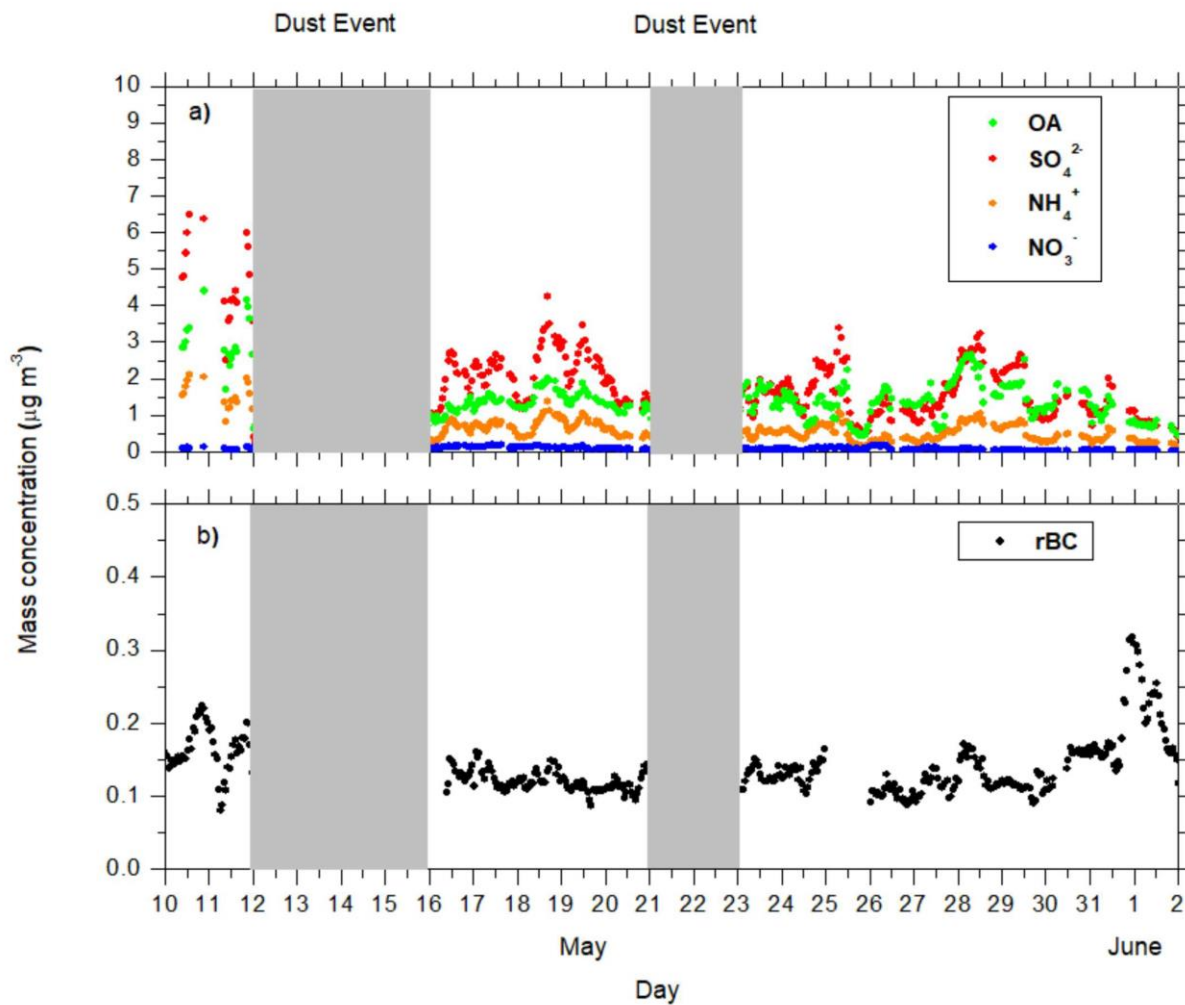
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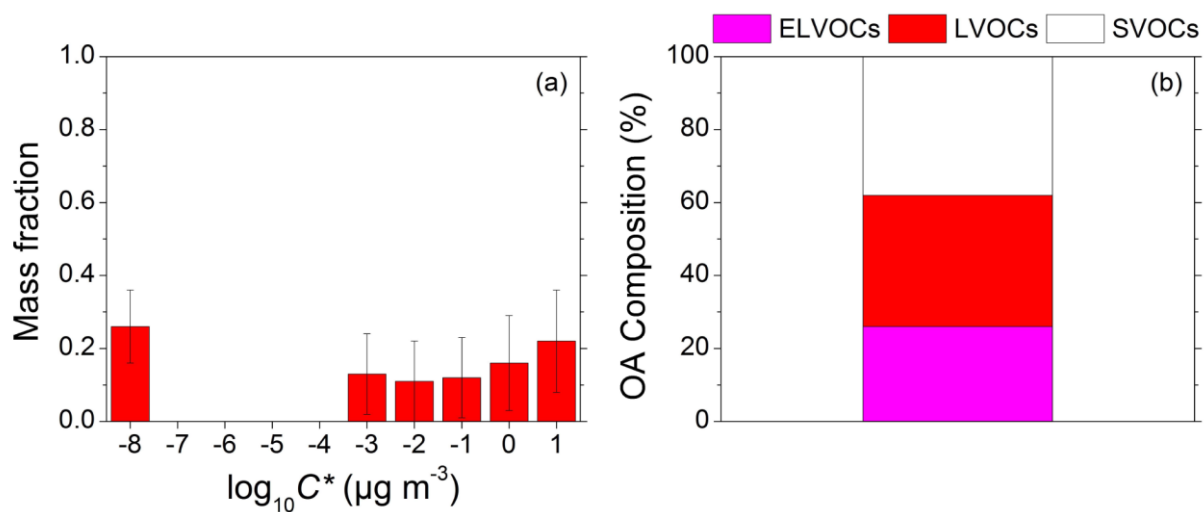


773

774 **Figure 1:** Evolution of the aerosol chemical composition based on the HR-TOF-AMS and SP2
 775 measurements during FAME-16: a) the non-refractory aerosol components; b) rBC concentration.

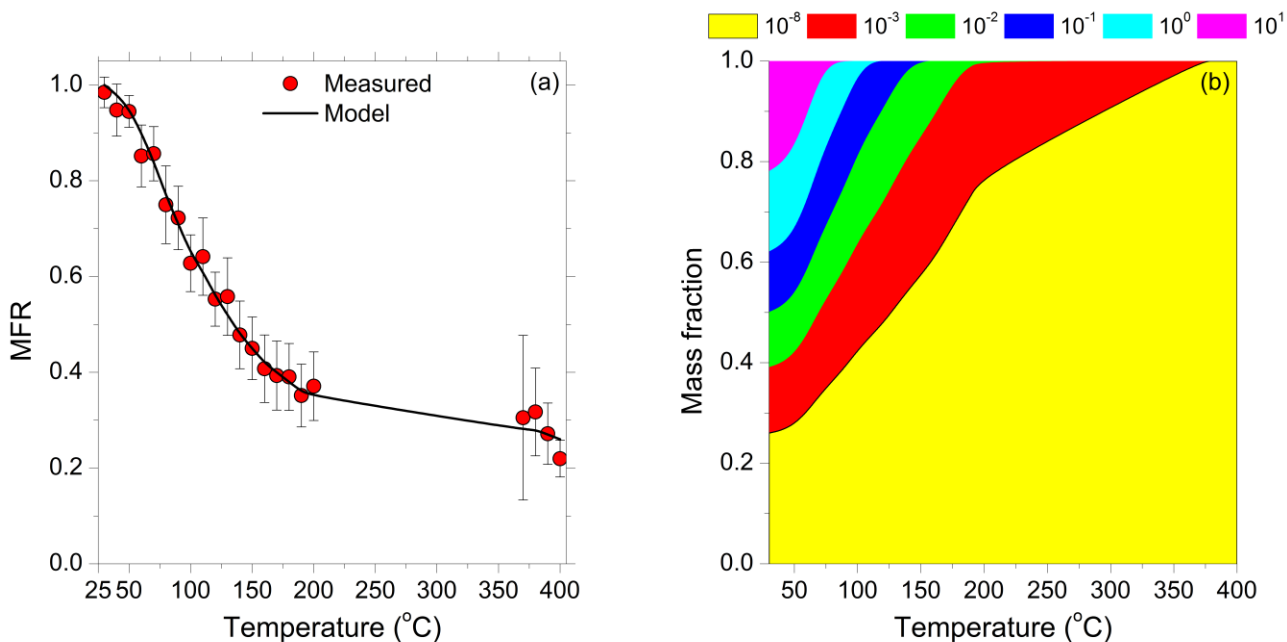
776 The shaded areas represent the dust events periods.

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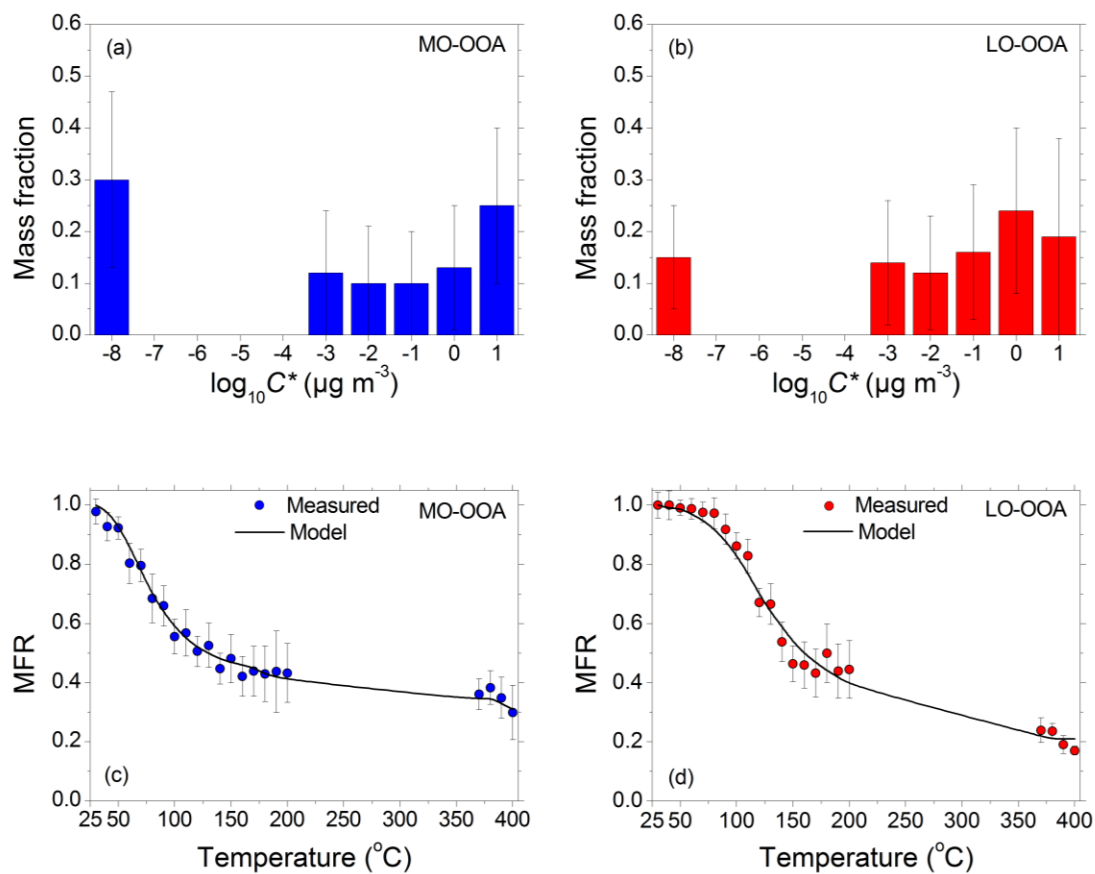
778

779 **Figure 2:** (a) Total OA volatility distribution along with its uncertainty estimated by the Karnezi
 780 et al. (2014) approach. The error bars represent the corresponding variability (± 1 standard
 781 deviation). (b) OA composition. Magenta color represents the ELVOCs, red the LVOCs, and white
 782 the SVOCs.



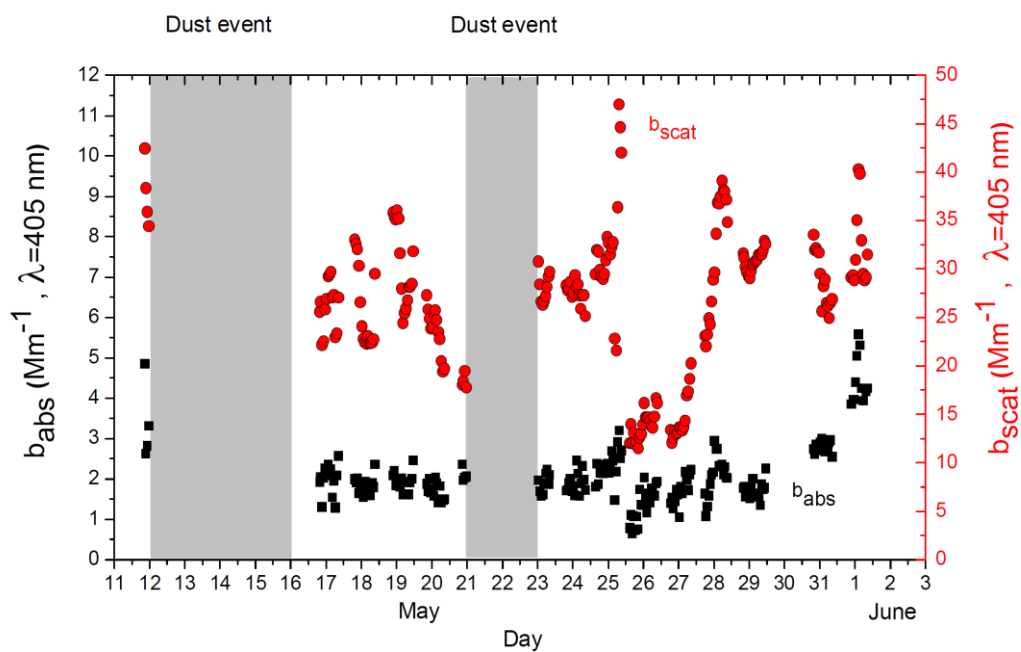
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784 **Figure 3:** (a) Average loss-corrected total OA thermograms. Red circles represent the measured
 785 total OA MFR and the error bars the corresponding variability (± 2 standard deviations of the
 786 mean). The solid lines are the model predictions (b) Mass fraction of the total OA for different
 787 effective saturation surrogate species with concentrations as a function of TD temperature. Yellow
 788 color represents the contribution of the effective saturation concentration $C^* = 10^{-8} \mu\text{g m}^{-3}$, red the
 789 contribution of the $C^* = 10^{-3} \mu\text{g m}^{-3}$, green the $C^* = 10^{-2} \mu\text{g m}^{-3}$, blue the $C^* = 10^{-1} \mu\text{g m}^{-3}$, cyan
 790 the $C^* = 10^0 \mu\text{g m}^{-3}$, and magenta the $C^* = 10 \mu\text{g m}^{-3}$.



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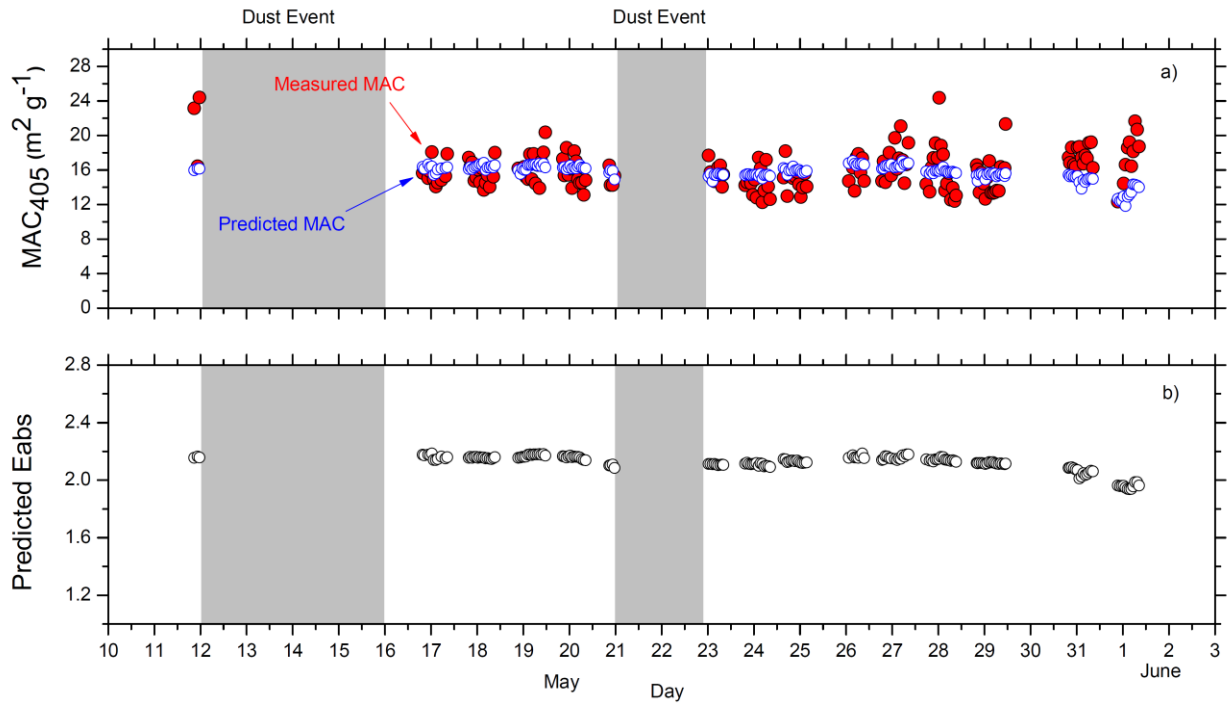
792 **Figure 4:** (a) Estimated volatility distribution of the MO-OOA factor along with its corresponding
 793 uncertainties by using the approach of Karnezi et al. (2014). (b) Estimated volatility distribution
 794 of the LO-OOA factor along with its corresponding uncertainties. (c) Measured (in circles) and
 795 predicted thermograms for the LO-OOA factor. The error bars represent ± 2 standard deviations
 796 of the mean. (d) Measured (in circles) and predicted thermograms for the LO-OOA factor. The
 797 error bars represent ± 2 standard deviations of the mean.



798

799 **Figure 5:** The timeseries of the aerosol optical properties at $\lambda=405$ nm. The black squares represent
 800 the absorption coefficient, b_{abs} , while the red circles represent the scattering coefficient, b_{scat} . The
 801 shaded areas represent the dust events periods.

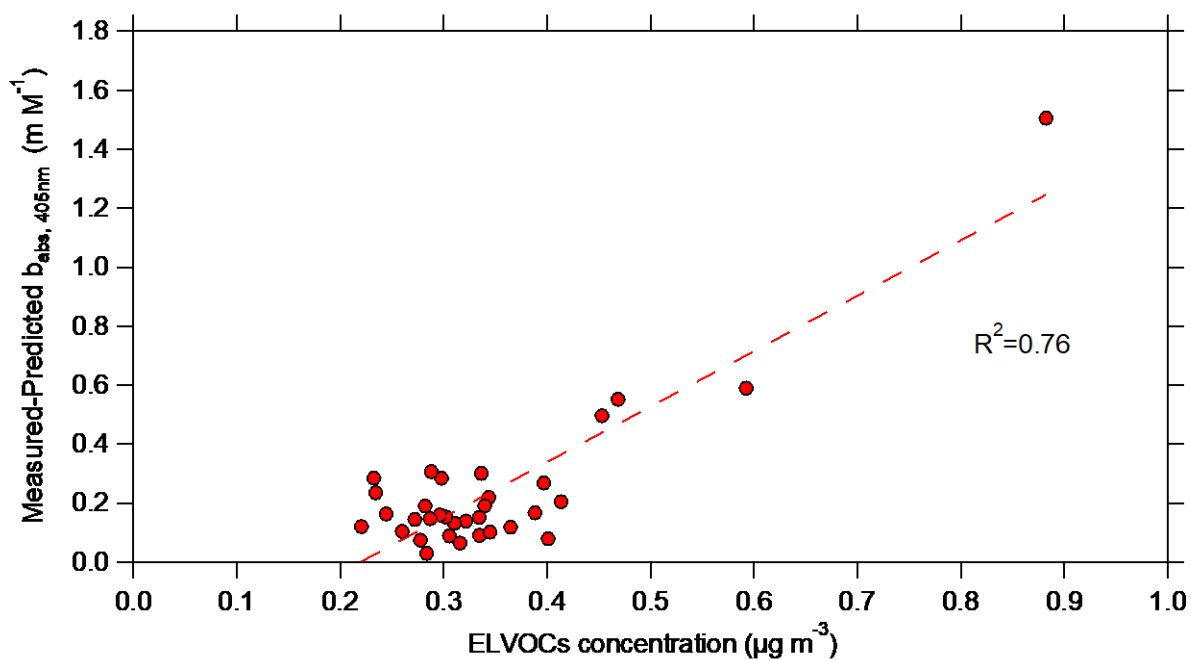
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804 **Figure 6:** Hourly averaged results from the Mie theory calculations, assuming a non-absorbing
 805 shell ($k=0$): a) predicted (blue circles) and measured MAC₄₀₅ (red circles). b) Hourly averaged
 806 Predicted E_{abs} values. The shaded areas represent the dust events periods.

807



808

809 **Figure 7:** The difference between the measured and the predicted unexplained $b_{\text{abs},405}$ as a function of the
810 estimated concentration of the ELVOCs. The data shown represent 3-hours averaged values.