1	Long-term real-time chemical characterization of submicron aerosols
2	at Montsec (Southern Pyrenees, 1570 m a.s.l.)
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Abstract. Real-time measurements of inorganic (sulfate, nitrate, ammonium, chloride and black carbon (BC)) and organic submicron aerosols (aerosols with an aerodynamic diameter less than 1 µm) from a continental background site (Montsec, MSC, 1570 m a.s.l.) in the Western Mediterranean Basin (WMB) were conducted for 10 months (July 2011 - April 2012). An Aerosol Chemical Speciation Monitor (ACSM) was co-located with other on-line and off-line PM<sub>1</sub> measurements. Analyses of the hourly, diurnal, and seasonal variations are presented here, for the first time for this region.

25 Seasonal trends in PM<sub>1</sub> components are attributed to variations in: evolution of 26 the planetary boundary layer (PBL) height, air mass origin, and meteorological 27 conditions. In summer, the higher temperature and solar radiation increases 28 convection, enhancing the growth of the PBL and the transport of anthropogenic 29 pollutants towards high altitude sites. Furthermore, the regional recirculation of air 30 masses over the WMB creates a continuous increase in the background concentrations 31 of PM<sub>1</sub> components and causes the formation of reservoir layers at relatively high 32 altitudes. The combination of all these atmospheric processes results in a high 33 variability of  $PM_1$  components, with poorly defined daily patterns, except for the organic 34 aerosols (OA). OA was mostly composed (up to 90%) of oxygenated organic aerosol 35 (OOA), split in two types: semi-volatile (SV-OOA) and low-volatile (LV-OOA), the rest 36 being hydrocarbon-like OA (HOA). The marked diurnal cycles of OA components 37 regardless of the air mass origin indicates that they are not only associated with 38 anthropogenic and long-range-transported secondary OA (SOA), but also with recently-39 produced biogenic SOA.

40 Very different conditions drive the aerosol phenomenology in winter at MSC. 41 The thermal inversions and the lower vertical development of the PBL leave MSC in 42 the free troposphere most of the day, being affected by PBL air masses only after 43 midday, when the mountain breezes transport emissions from the adjacent valleys and 44 plains to the top of the mountain. This results in clear diurnal patterns of both organic and inorganic concentrations. OA was also mainly composed (71%) of OOA, with 45 contributions from HOA (5%) and biomass burning OA (BBOA; 24%). Moreover, in 46 47 winter sporadic long-range transport from mainland Europe is observed.

The results obtained in the present study highlight the importance of SOA formation processes at a remote site such as MSC, especially in summer. Additional research is needed to characterize the sources and processes of SOA formation at remote sites.

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Keywords: high altitude, mountain, remote, continental background, ACSM.

#### 55 **1** Introduction

56 Earth's climate system is modulated by atmospheric aerosols. Submicron 57 particles (< 1 µm in aerodynamic diameter) play a dominant role in both cloud 58 formation and scattering or absorbing solar radiation (IPCC, 2013). The complexity of 59 aerosol sources and processes results in an uncertainty in the radiative forcing of 60 climate. Aerosol optical properties are connected to direct and indirect climate forcing 61 effects, and they are dependent on particle composition. Moreover, aerosol 62 composition may provide valuable information on aerosol sources and processes. 63 Consequently, long-term measurements of PM<sub>1</sub> chemical composition are needed to 64 better understand aerosol sources, to quantify their lifetime in the atmosphere and to 65 constrain the uncertainties of their climatic influence.

66 Long-term PM<sub>1</sub> chemical composition measurements are relatively scarce both 67 off-line and on-line. In the last decade, on-line PM1 chemical composition 68 measurements have been performed using aerosol mass spectrometers (AMS) at a 69 number of locations. Measurements of on-line chemical composition are useful to study 70 hourly variations and daily patterns. Most of these studies, however, correspond to 71 short-term measurement campaigns (typically a month) (e.g. Crippa et al., 2014; 72 Jimenez et al., 2009; Lanz et al., 2010; Ng et al., 2010; Zhang et al., 2007) given the 73 intensive instrument maintenance required and the need of highly-qualified personnel 74 for a good quality dataset.

75 In contrast to the use of the AMS in relatively short campaigns, the more 76 recently developed Aerodyne Aerosol Chemical Speciation Monitor (ACSM) is 77 becoming a widely used on-line instrument for long-term measurements of PM1 78 chemical composition (Budisulistiorini et al., 2014; Canonaco et al., 2013; Petit et al., 79 2014; Tiitta et al., 2014). The ACSM is built upon the same technology as the AMS, in which an aerodynamic particle focusing lens is combined with high vacuum thermal 80 81 particle vaporization, electron impact ionization, and mass spectrometry. Modifications 82 in the ACSM design (e.g. lack of particle sizing chamber and components, use of 83 simple and compact RGA mass spectrometer detector), however, allow it to be smaller, 84 lower cost, and simpler to operate than the AMS (Ng et al., 2011c). The ACTRIS 85 (Aerosols, Clouds, and Trace gases Research InfraStructure) European network is 86 evaluating the use of the ACSM as a reliable instrument, which will provide the 87 opportunity to study long-term datasets of PM<sub>1</sub> chemical composition across the 88 continent.

Recent publications have investigated most of the existing worldwide AMS
databases (e.g. Crippa et al., 2014; Jimenez et al., 2009; Lanz et al., 2010; Ng et al.,
2010; Zhang et al., 2007) and reflected a prevalence of organic aerosols (20 to 90%) in

92 the submicron fraction, largely independent of the region and type of environment. However, our knowledge on organic aerosol (OA) formation, sources, and atmospheric 93 94 processing is still very incomplete, especially for secondary organic aerosols (SOA) 95 formed from chemical reactions of gas-phase species (e. g. Donahue et al., 2014; 96 Hallquist et al., 2009; Kroll and Seinfeld, 2008; Robinson et al., 2007; Volkamer et al., 97 2006). Recent progress has been made in identifying primary organic aerosols (POA) 98 sources (e.g. Elbert et al., 2007; Zhang et al., 2005), but significant gaps still remain in 99 our understanding on the atmospheric evolution of POA after emission (de Gouw and 100 Jimenez, 2009). For these reasons, OA measurements and analysis are required to 101 better understand its chemical evolution in the atmosphere.

102 The lack of long-term on-line PM<sub>1</sub> chemical composition measurements is 103 especially evident in the Western Mediterranean Basin (WMB), which is characterized 104 by particular atmospheric dynamics strongly influenced by its topography (Jorba et al., 105 2013; Millan et al., 1997). Over this region, arrival of natural and anthropogenic 106 aerosols as a result of long-range transport from Africa and Europe is frequent (e.g. 107 Pey et al., 2013; Querol et al., 2009; Ripoll et al., 2014; Rodríguez et al., 2001) and 108 accumulation and recirculation processes are frequently observed (Rodriguez et al., 109 2002). The sources and meteorological controls of PM in the regional background of 110 the WMB have been recently investigated during the DAURE study (Pandolfi et al., 2014a) using an AMS and <sup>14</sup>C analyses (Crippa et al., 2014; Minguillón et al., 2011). 111 112 Furthermore, Ripoll et al. (2015) studied the  $PM_1$  and  $PM_{10}$  chemical composition with 113 daily time resolution in the continental and regional background environments in the 114 WMB. In that study, a higher mineral contribution was identified in the continental 115 background due to the preferential transport of African dust at high altitude layers and 116 to the increased regional dust resuspension enhanced by the drier surface and higher 117 convection. Nevertheless, aerosol chemical characterization with higher time resolution 118 is needed to study the origin of specific PM components and the local and/or regional 119 processes, in particular to exploit the information contained in diurnal cycles that is 120 typically not accessible with off-line measurements.

121 In this study we deployed an ACSM at a high altitude site (Montsec, 1570 m a.s.l.) in the NE of the Iberian Peninsula (42° 03' N, 0° 44' E), representative of the 122 123 continental background conditions of the Western Mediterranean Basin (WMB) (Ripoll 124 et al., 2014). This environment is under free tropospheric (FT) influence most of the 125 time, although it is exposed to regional pollutants during the summer time and/or under 126 the influence of mountain breezes, and it is affected by trans-boundary incursions of 127 natural and anthropogenic aerosols from Europe and North Africa (Ripoll et al., 2014, 128 2015). Co-located on-line and off-line PM<sub>1</sub> measurements were also carried out to

129 complement the ACSM dataset. Hence, the work presented here interprets the real-130 time variation of inorganic and organic submicron components during 10 months (July 131 2011 - April 2012), and the types of OA are also studied. Special emphasis is placed 132 on the analysis of diurnal pattern and seasonal variations of chemical components and 133 the main factors influencing these variations.

## 134 2 Methodology

## 135 2.1 Sampling site

Montsec\_site (MSC) is located on the highest part of the Montsec d'Ares
mountain, at an altitude of 1570 m a.s.l., in a plain near to the edge of a 1000 m cliff to
the south, with no wind obstructions present around. It is located in the NE of the
lberian Peninsula (42°03'N, 0°43'E), 50 km S of the Pyrenees and 140 km NW of
Barcelona (Fig.S1). A detailed description of this site can be found in Ripoll et al.
(2014).

142 The daily classification of atmospheric episodes affecting MSC was made 143 following the procedure described by Ripoll et al. (2014) using HYSPLIT model from 144 the NOAA Air Resources Laboratory (ARL). Air masses reaching MSC are mainly from 145 the Atlantic (62% of the days) all along the year. From March to October, North African 146 (NAF) episodes are more frequent (17% of the days) and very often are alternated with 147 the summer regional (SREG) scenarios (12% of the days). The winter regional 148 (WREG) scenarios are detected from October to March (11% of the days), as well as 149 the European (EU) episodes (11% of the days). Conversely, the Mediterranean (MED) 150 episodes are detected sporadically (4% of the days).

151 The boundary layer height was calculated using the <u>Global Data Assimilation</u> 152 <u>System (GDAS)</u> model from the NOAA Air Resources Laboratory 153 (<u>http://www.ready.noaa.gov/READYamet.php</u>) (Fig.S2).

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## 2.2 ACSM sampling and data analysis

155 The aerosol chemical speciation monitor (ACSM) (Aerodyne Research Inc.) was measuring continuously from July 2011 to April 2012. The ACSM provides real-156 157 time mass concentration of submicron particulate organics, nitrate, sulfate, ammonium 158 and chloride via thermal vaporization and electron impact ionization, with detection by a 159 quadrupole mass spectrometer (Ng et al., 2011c). The mass concentration of a given 160 species is determined from the sum of the ion signals at each of its mass spectral 161 fragments and its Ionization Efficiency (IE) (Canagaratna et al., 2007). Since calibration of IEs for all species is not feasible, the Relative Ionization Efficiency (RIE) (compared 162 163 to that of nitrate) is used (Jimenez et al., 2003). The ammonium nitrate calibration 164 described by Ng et al. (2011b) was performed using an atomizer (TSI, Constant Output 165 Atomizer Model 3076) for primary aerosol generation, followed by a silica gel diffusion 166 dryer, a differential mobility analyzer (DMA) model TSI 3936, and a condensation 167 particle counter (CPC, TSI 3772). Monodisperse 300 nm ammonium nitrate aerosol 168 particles were used, covering a range of nitrate concentrations from 2 to 15  $\mu$ g m<sup>-3</sup>. 169 Several calibrations were conducted throughout the sampling period, and average 170 values of 2.2 x 10<sup>-11</sup> for nitrate IE and 5.4 for RIE for ammonium were used for the 171 whole dataset. The RIE values used in this study for the rest of the species were those 172 usually applied in AMS ambient concentrations: 1.4 for OA and 1.1, 1.2, and 1.3 for 173 nitrate, sulfate, and chloride, respectively (Canagaratna et al., 2007). RIE for sulfate 174 was experimentally determined one year later and was found to be 1.26, although the 175 default value was used for the current dataset. A time resolution of 62 minutes was 176 used as a result of 12 scans (1 open and 1 filtered) per data point with a scan speed of 177  $1 \, \text{s amu}^{-1}$ .

178 The ACSM data were analyzed with the standard ACSM data analysis software 179 version 1.5.3.2 (Aerodyne Research Inc.) written in Igor Pro 6 (WaveMetrics, Inc., Lake 180 Oswego, OR, USA). As the ACSM was measuring continuously for a long time, the 181 standard correction for instrument sensitivity drifts was applied to the dataset based on 182 the inlet pressure and N<sub>2</sub> signal. Finally, mass concentrations were corrected using a 183 Collection Efficiency (CE) to account for the particle bounce of aerosols on the 184 vaporizer. The composition-dependent CE was calculated as described by Middlebrook 185 et al. (2012) and was close to 0.45 for most of the time. Since for most ambient studies 186 a 0.5 CE value is found to be representative with data uncertainties generally within 187 ±20% (Canagaratna et al., 2007), and since our ACSM concentrations using CE=0.5 188 were in good agreement with concentrations from other co-located instruments, a CE 189 of 0.5 was used.

190 The organic components were further investigated by applying Multilinear 191 Engine (ME-2) (Paatero, 1999) to the organic mass spectra. With the ME-2, the user 192 can introduce a priori information about sources with the so-called a-value approach. 193 Hence, the user inputs one or more factor profiles and a constraint defined by the a-194 value, which determines the extent to which the output profile can differ from the profile 195 fed to the model. The source apportionment of OA was performed applying ME-2 using the custom software tool of Source Finder (SoFi) version 4.8 developed by Canonaco 196 197 et al. (2013). The ME-2 analysis was carried out separately for the summer period (14 Jul 11 – 24 Sep 11) and the winter period (10 Jan 12 – 7 Mar 12). Only  $m/z \le 100$  were 198 199 used for source apportionment of OA because: a) the signals of m/z>100 account for a 200 minor fraction of the total organic mass (on average, 2 %), b) the m/z>100 have larger

uncertainties, and c) the large interference of naphthalene signals (used for m/zcalibration of the ACSM) at these m/z (e.g., m/z 127, 128, and 129) (Sun et al., 2012).

203 2.3

### 2.3 Co-located measurements used in this study

204 MSC has been permanently equipped with aerosol monitoring instrumentation 205 since January 2010 and some of these measurements were used in this study. 24-h 206 PM<sub>1</sub> samples were collected every 4 days on 150 mm quartz micro-fiber filters (Pallflex QAT) using high volume samplers (30 m<sup>3</sup> h<sup>-1</sup>, MCV CAV-A/MSb) equipped with MCV 207 PM1 cut-off inlets. Daily PM1 mass concentrations were determined by off-line 208 209 gravimetric procedures according to the EN 12341 standard (CEN, 1999), i.e. at 20°C 210 temperature and 50% relative humidity. Furthermore, PM<sub>1</sub> chemical composition was 211 obtained as described by Ripoll et al. (2015) using inductively coupled plasma atomic 212 emission spectroscopy (ICP-AES) and mass spectrometry (ICP-MS) for major and 213 trace elements, respectively, ion high performance liquid chromatography (HPLC) and 214 selective electrode for ions concentrations, and thermal-optical method (using the 215 EUSSAR 2 protocol) for elemental carbon (EC) and organic carbon (OC) 216 concentrations. Real-time PM<sub>1</sub> mass concentrations were continuously measured by an 217 optical particle counter (OPC, Model GRIMM 1.107 calibrated with different latex PSL 218 refraction index 1.59). PM<sub>1</sub> 30-minute data were daily averaged and subsequently 219 corrected by comparison with 24-h standard gravimetric mass measurements. The 220 absorption coefficient was measured continuously at 637 nm using a Multi Angle 221 Absorption Photometer (MAAP, model 5012, Thermo). Equivalent black carbon (BC) 222 mass concentrations (Petzold et al., 2013) were calculated by the MAAP instrument 223 software by dividing the measured absorption coefficient  $\sigma_{ap}$  ( $\lambda$ ) by 6.6 m<sup>2</sup> g<sup>-1</sup>, which is 224 the instrument default mass absorption cross section (MAC) at 637 nm (Müller et al., 225 2011; Petzold and Schönlinner, 2004). Particle scattering ( $\sigma_{sp}$ ; 0°-360°) and 226 hemispheric backscattering ( $\sigma_{bsp}$ ; 90°-270°) coefficients at three wavelengths (450nm, 227 525nm, 635nm) were measured with a LED-based integrating nephelometer (model 228 Aurora 3000, ECOTECH Pty, Ltd, Knoxfield, Australia).

Finally, all meteorological data were measured by the Catalonian Meteorological Service from the Montsec d'Ares station. Gaseous pollutants (O<sub>3</sub>, NO, NO<sub>2</sub>, CO and SO<sub>2</sub>) were measured using real-time monitors belonging to the Department of Environment of the Autonomous Government of Catalonia. <u>NO and NO<sub>2</sub></u> <u>concentrations were measured using a Thermo Scientific instrument, Model 42i-TL;</u> <u>CO using a Teledyne 300 EU Gas filter correlation analyzer; O<sub>3</sub> using a MCV 48AV UV photometry analyzer; and SO<sub>2</sub> using a Teledyne 100 EU UV fluorescence analyzer.</u> 236 In addition to these routine measurements, 2 intensive campaigns were 237 performed in July - August 2011 and January - February 2012. During these intensive 238 campaigns PM<sub>1</sub> filters were collected daily and a scanning mobility particle sizer 239 (SMPS) was installed to measure particle number size distribution of mobility diameters 240 11-350 nm in the summer campaign, and 8-450 nm in the winter campaign. The SMPS 241 system comprises a classifier unit (Model TSI 3080) and a differential mobility analyzer 242 (DMA, Model TSI 3081) connected to a condensation particle counter (CPC, Model TSI 243 3772). The SMPS data for the winter campaign were also used to estimate the mass 244 concentration to compare with the ACSM data. To this end, the volume size 245 distributions were calculated from the measured particle number distributions assuming 246 sphericity. The total volume concentrations were computed by integrating over the 247 measured particle range and converted to mass concentration using the estimated 248 composition-dependent density, calculated using the chemical composition given by 249 the ACSM and the equation of Salcedo et al. (2006). Average concentrations shown in 250 the whole paper are arithmetical averages unless otherwise specified.

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3

#### **Results and discussion**

#### 252 3.1 Submicron aerosol mass concentrations

253 In order to establish the consistency of the different measurements during this 254 study, the sum of the ACSM species (= sulfate + nitrate + ammonium + OA + chloride) 255 and the BC mass concentrations was compared with the co-located PM1 and light 256 scattering measurements (Fig. 1). The scatter plots of ACSM plus BC concentrations 257 versus PM<sub>1</sub> concentrations from the OPC and SMPS showed strong correlations 258 (R<sup>2</sup>=0.72 and R<sup>2</sup>=0.87, respectively) and slopes close to unity (slope=0.94 and 259 intercept=0.09 for the ACSM+BC vs. OPC, and slope=1.21 and intercept=0.86 for the 260 ACSM+BC vs. SMPS) (Fig.S3). The differences in the particle size range measured by the different instruments needs to be considered when assessing these comparisons. 261 262 Moreover, ACSM plus BC concentrations were also highly correlated with light scattering at 525 nm determined by the nephelometer (R<sup>2</sup>=0.85, Fig.S3). The high 263 264 degree of agreement is also apparent in the time series plots of  $PM_1$  shown in Fig. 1.

The average concentration ( $25^{th}$ ,  $50^{th}$ ,  $75^{th}$  percentiles) of the ACSM + BC mass during this study (July 2011 - April 2012) was 4.9 µg m<sup>-3</sup> (0.9, 2.8, 7.9 µg m<sup>-3</sup>) (Table S1), which is similar to the 2010-2012 average reported by Ripoll et al. (2014) from OPC measurements ( $5.0 \mu g m^{-3}$ ). For the sake of brevity only summer (<u>14 Jul 11 – 24</u> Sep 11) and winter (<u>10 Jan 12 – 7 Mar 12</u>) hourly variation will be discussed in the following sections, given that hourly variation in spring was similar to summer and that 271 in fall was similar to winter. The seasonal average PM<sub>1</sub> concentrations were higher in summer (7.5  $\mu$ g m<sup>-3</sup> (3.4, 7.1, 10.5  $\mu$ g m<sup>-3</sup>), Table S1) than in winter (4.1  $\mu$ g m<sup>-3</sup> (0.8, 272 1.7, 5.6 µg m<sup>-3</sup>), Table S1). A similar seasonal pattern has been described at other high 273 274 altitude sites in Europe (e.g. Carbone et al., 2014; Cozic et al., 2008; Freney et al., 275 2011; Tositti et al., 2013), being associated with differences in the air mass origin from 276 summer to winter, and also to variations in the planetary boundary layer (PBL) height. 277 The seasonal variation at MSC has been described in detail in recent works (Pandolfi 278 et al., 2014b; Ripoll et al., 2014, 2015), and it has been principally attributed to the 279 seasonal variation of the PBL (Fig.S2). In summer, the stable anticyclonic conditions 280 over this continental area enhance convection increasing the development of the PBL 281 and favoring the transport of anthropogenic pollutants towards high altitude sites such 282 as MSC. The situation in winter is notably different, as the lower vertical development 283 of the PBL over this area leaves high altitude sites in the FT, isolating MSC from 284 polluted air masses. The seasonal variation of PM concentrations at MSC has been 285 also connected to mesoscale and synoptic processes. At MSC, southern flows and 286 regional recirculation episodes are more frequent in summer, whereas clean Atlantic 287 advections and northeastern winds from mainland Europe are more common in winter 288 (Fig.S4 and Table S2) (Pandolfi et al., 2014b; Ripoll et al., 2014, 2015). Moreover, the 289 summer maximum has been ascribed to the more vigorous photochemistry in the 290 atmosphere that enhances the formation of secondary inorganic and organic aerosols 291 (Querol et al., 1999).

#### 292 3.2 Submicron aerosol chemical composition

293 Concentrations of ACSM species were daily averaged and compared with off-294 line measurements from 24-h PM<sub>1</sub> samples, and all species showed strong correlations 295 (R<sup>2</sup> between 0.77 and 0.96, Fig.S5). Different slopes (ACSM vs off-line measurements) 296 were found for each of the species: 1.12 for sulfate, 1.31 for ammonium and 1.35 for 297 nitrate. The relatively higher slope for nitrate, with respect to sulfate, could be attributed 298 to a sampling negative artifact due to the volatilization of nitrate on the off-line samples 299 (Schaap et al., 2004). Ammonium is present as a counterion for sulfate and nitrate, and 300 thus its slope is in between those of the two species.

For the OA, the slope could be interpreted as the OM-to-OC ratio, since the offline measurements determined organic carbon. This slope was found to be 3.39, however values of 2.2 are more common for aged aerosol (e.g. Aiken et al., 2008; Minguillón et al., 2011; Takahama et al., 2011). This disagreement of a factor of 1.54 may be attributed to different reasons. A negative volatilization artifact may occur in the filters, hence resulting in an underestimation of OC. Alternatively it is possible that the 307 RIE for OA in the ACSM is larger than the value of 1.4 determined for the AMS, a topic 308 currently being investigated by the ACSM manufacturer, which would result in an 309 overestimation of OA. Similar series of intercomparisons with a similar discrepancy for 310 OA has been found for a one-year dataset (June 2012 – July 2013) with the same 311 instrument at Montseny site (Minguillón et al., 2015) and for a recently reported study in 312 Atlanta, US by Budisulistiorini et al. (2014). Assuming that the disagreement was due 313 to the overestimation by the ACSM, the OA concentrations were corrected dividing by 314 the disagreement factor (1.54) to compare the results with co-located measurements 315 (Fig.S3). The resulting slopes were very similar and hence OA concentrations reported 316 in the present paper were not corrected since further research is needed to better 317 estimate the RIE for OA in the ACSM.

318 The average of PM<sub>1</sub> chemical composition at MSC during this study (July 2011 -319 April 2012) is given in Fig. 2. On average, OA was the largest PM<sub>1</sub> constituent (50%), 320 followed by sulfate (20%), nitrate (14%), ammonium (12%), BC (4%) and chloride (1%). 321 As was the case of PM<sub>1</sub> concentrations, all chemical components increased in summer 322 and decreased in winter, with the exception of nitrate (Fig. 2 and Table S1). The higher 323 nitrate concentrations in winter than in summer were also observed in other studies in 324 the Mediterranean region (e.g. Pey et al., 2009; Querol et al., 2009; Ripoll et al., 2015) 325 and this variation was attributed to the high volatility of ammonium nitrate at low 326 humidity and high temperature (Zhuang et al., 1999). At MSC, the summer maximum of 327 the rest of PM components has been mainly ascribed to the higher temperature and 328 solar radiation in summer (Table S2), which enhances atmospheric photochemistry, 329 promoting the formation of secondary inorganic and organic aerosols. All these 330 seasonal characteristics are described in detail in Ripoll et al. (2015).

### 331 3.2.1 Summer trends

332 Time series of  $PM_1$  components during summer time (14 Jul – 24 Sep 2011) are 333 shown in Fig. 3. Wind direction and wind speed, temperature, precipitation and 334 concentrations of nitrogen oxides, sulfur dioxide, and ozone are also depicted. The 335 daily classification of atmospheric episodes affecting MSC is also illustrated in different 336 background colors. On average, during summer the lowest concentrations of all  $PM_1$ 337 components and gases were recorded under the Atlantic advection conditions since 338 these air masses are associated with precipitation, decreased temperature and solar 339 radiation, and strong winds, leading to cleaner atmospheric conditions. Conversely, 340 summer regional episodes lasted for 6 to 11 consecutive days and led to sustained 341 increases of the background concentrations of sulfate, OA and BC at MSC. Despite the 342 limited ACSM data availability during North African episodes, relatively high 343 concentrations of PM<sub>1</sub> components were observed under this type of episodes,
344 especially BC (Fig. 3).

345 The diurnal cycles of PM<sub>1</sub> components, gaseous pollutants and meteorological 346 variables are shown in Fig. 4. The summer concentrations of PM<sub>1</sub> components and 347 gases showed no clear diurnal patterns, except for ozone and OA. This lack of defined daily patterns is similar to the findings obtained at the high altitude Puy-de-Dôme 348 349 station in central France (Freney et al., 2011, the only similar study found at a remote 350 site). For a high altitude site as MSC, the lack of diurnal cycles can be explained by a 351 combination of factors. In summer, the recirculation of air masses over the WMB 352 induced by an abrupt orography (Fig.S1) causes the formation of reservoir layers at 353 any time at relatively high altitudes (Millan et al., 1997; Rodríguez et al., 2003). 354 Moreover, long-range transport from North Africa, which can be more intense at high 355 altitude layers (Ripoll et al., 2015), could also blur the daily patterns since the 356 occurrence of this transport does not depend on the time of the day. These factors 357 result in a lack of well-defined daily patterns but in a high variability of diurnal cycles 358 even within the same type of episode, which is reflected in the similar average daily 359 evolutions and the high standard deviations calculated for the average daily patterns 360 when separated by air mass origin (Fig.S6).

361 The ozone and OA concentrations had a marked diurnal cycle regardless of the 362 air mass origin (Fig. 4 and Fig.S6) in summer. These different daily patterns with 363 respect to the rest of the gases and chemical components points to the fact that ozone 364 and OA variations are strongly influenced by local/regional processes and not just 365 dominated by long-range transport. Minimum ozone concentrations were recorded 366 between 8:00 and 9:00 UTC (Coordinated Universal Time, which is local time - 1:00h 367 and local summer time - 2:00h), whereas maximum concentrations were measured 368 between 16:00 and 17:00 UTC. In contrast, the highest OA concentrations were 369 observed around 12:00 UTC, and the lowest during the night and in the early morning. 370 The ozone variations may influence those of OA, although the complete understanding 371 of the ozone diurnal evolution is outside the scope of this study. The average increase 372 in OA during the day is likely due to the photooxidation of volatile organic compounds 373 (VOCs). Given that MSC is a remote site, in summer VOCs are most likely dominated 374 by local biogenic emissions (BVOCs), as it was found in the Mediterranean forested 375 area of Montseny (Seco et al., 2011). Hence, the midday increase is likely due to 376 recently-produced biogenic SOA, and to a lesser degree, photooxidation of 377 anthropogenic VOCs.

378 Despite the marked diurnal cycle of OA regardless of air mass origin, the 379 average increase during the day with respect to average concentrations during the

night was higher under summer regional (2.6  $\mu$ g m<sup>-3</sup>) and North African (3.0  $\mu$ g m<sup>-3</sup>) 380 episodes than during Atlantic advections (1.3 µg m<sup>-3</sup>) (Fig.S6). This difference could be 381 382 caused by the higher SOA formation. This is due to the increase in BVOCs emissions 383 and atmospheric photooxidation caused by the higher temperature and solar radiation 384 (Paasonen et al., 2013; Seco et al., 2011) under summer regional and North African 385 episodes. Furthermore, under these episodes higher concentrations of ozone were 386 measured, which also favors the formation of SOA (via direct oxidation and also by 387 leading to higher OH concentrations). The SOA formation registered at MSC is 388 relatively high when compared to other high altitude sites such as Puy-de-Dôme 389 (Freney et al., 2011). This is in agreement with the modeled SOA emissions over 390 Europe, which identified higher SOA concentrations in Mediterranean environments 391 (Bessagnet et al., 2008). This higher SOA formation is probably due to the higher 392 emissions of BVOCs in the Mediterranean forested areas (up to 3 times higher than 393 Boreal forested areas) (Bessagnet et al., 2008; Lang-Yona et al., 2010; Steinbrecher et 394 al., 2009) and the comparable concentrations of tropospheric ozone with other high 395 altitude European sites (Chevalier et al., 2007). On the other hand, the extra formation 396 of SOA under summer regional and North African episodes might also have a 397 contribution from the photooxidation of anthropogenic VOCs, since Atlantic advections 398 are associated with cleaner atmospheric conditions.

#### 399 3.2.2 Winter trends

Similar to summer, the lowest concentrations of all PM<sub>1</sub> components and gases in winter (10 Jan 12 – 7 Mar 12) were recorded under the Atlantic advections, whereas the highest were measured when MSC was affected by air masses from mainland Europe and sporadically under regional conditions (Fig. 5). Mediterranean air masses were detected very infrequently and therefore conclusions on their characteristics will not be drawn in the present paper. The relative contribution of different components was similar, with OA representing a somewhat smaller fraction than in summer.

407 In contrast to what was found in summer, in winter concentrations of most PM<sub>1</sub> 408 components and gaseous pollutants showed much clearer diurnal patterns, with a 409 minimum around 7:00 UTC and a maximum between 14:00 and 15:00 UTC (Fig. 4). 410 Similar patterns have been observed at the Puy-de-Dôme station during a winter 411 campaign (Freney et al., 2011). These daily cycles are probably caused by the fact that 412 MSC is located most of the day within the FT in winter, whereas PBL air masses are only injected upwards after midday (Fig.S2). Moreover, thermal inversions are very 413 414 frequent from 20:00 to 07:00 UTC. These situations prevent the transport of pollutants 415 from the lower populated areas towards higher altitudes, especially at night. During the 416 morning, the thermal inversions dissipate due to the radiative warming of the ground 417 and mountain upslope winds develop (e.g. Henne et al., 2004). These mountain winds 418 transport anthropogenic emissions from the adjacent valleys and plains to the top of 419 the mountain, with a maximum upslope transport in the afternoon. Moreover, biogenic 420 emissions influence cannot be ruled out as average winter temperatures are high 421 enough for them to occur (Seco et al., 2011; Steinbrecher et al., 2009). Thus, mountain 422 breezes play an important role in determining the diurnal variation of PM<sub>1</sub> components 423 in winter (Fig. 4), especially under regional conditions. A clear example of the  $PM_1$ 424 components diurnal pattern under winter regional episodes was observed from 22 to 25 425 February 2012 with  $PM_1$  concentrations (and  $NO_x$ ) increasing several fold during the afternoon (Fig. 5). 426

427 The study of the daily cycles as a function of air mass origin (Fig.S7) showed 428 clear diurnal patterns under winter regional episodes, as mentioned above, and less 429 marked daily patterns when MSC is affected by Atlantic advections and long-range 430 transport from mainland Europe. Under Atlantic episodes the concentrations of PM<sub>1</sub> 431 components were very low and the standard deviations with respect to the average 432 pattern were guite high, resulting in unclear diurnal patterns compared to those under 433 winter regional conditions. During European episodes, which can be more intense at 434 high altitude layers (Ripoll et al., 2015; Sicard et al., 2011), background concentrations 435 of PM<sub>1</sub> components were higher and the midday increment was lower compared to 436 those under winter regional conditions, resulting in less marked daily patterns. These 437 less-marked diurnal cycles are probably due to the fact that the increase of PM1 438 components occurs during these episodes regardless of the time of the day. A good 439 example of this less-marked diurnal variation during European episodes was observed 440 from 17 to 19 February 2012 (Fig. 5).

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# 3.3 Characterization of OA components

442 In order to better characterize the profiles of OA components, the ME-2 analysis 443 was performed separately for the summer period (14 Jul - 24 Sep 2011) and the winter 444 period (10 Jan 12 – 7 Mar 12), since OA components are expected to vary throughout 445 the year. The solution of ME-2 analysis selected for each season was based on several 446 tests, with different number of factors and different a-values, taking into account the 447 correlations with external tracers (including nitrate, sulfate, BC and ozone), the daily 448 patterns of each factor, and the residuals. As a result, a solution of 3 factors was 449 selected for each season. In summer, a hydrocarbon-like OA (HOA), a semi-volatile 450 oxygenated OA (SV-OOA) and a low-volatile oxygenated OA (LV-OOA) (Fig.S8) were 451 resolved. The HOA factor was constrained using an average HOA factor from different

452 datasets (Ng et al., 2011b), with an a-value of 0.1. The SV-OOA was characterized by 453 a high 43-to-44 ratio, and the LV-OOA was defined by having a dominant peak at m/z454 44. In winter the 3 factors identified were: hydrocarbon-like OA (HOA), biomass burning 455 OA (BBOA) and oxygenated OA (OOA) (Fig.S8). The HOA and the BBOA factors were 456 constrained based on the profiles from different datasets (Ng et al., 2011b), with an a-457 value of 0.1 in both factors, and including BC as an additional variable to the ME-2 458 analysis. The OOA was characterized by a 43-to-44 ratio between those found for the 459 LV-OOA and the SV-OOA in summer, and by having a dominant peak at m/z 44. This 460 high signal of m/z 44 in the winter OOA indicates a high degree of oxidation, and 461 therefore a dominant aged character during winter. A solution with 4 factors in winter 462 was investigated in order to split the OOA into SV-OOA and LV-OOA, but the resulting profiles did not represent two different OOA types, and the time series showed that one 463 464 of the factors was mainly representing noise. The HOA profiles obtained for the 465 summer and winter periods were similar and showed similar deviations from the Ng et 466 al. (2011b) spectrum (Fig. S9), since they were constrained with the same anchor HOA 467 profile.

On average, LV-OOA dominated the OA fraction in summer contributing 64%, 468 469 followed by SV-OOA (26%) and HOA (10%) (Fig. 6), whereas in winter OOA accounted 470 for 71%, BBOA contributed 24%, and HOA contribution decreased to 5% (Fig. 7). The 471 high contribution of OOA components confirms the initial hypothesis of MSC organic 472 components being mostly secondary in their origin, and it is in agreement with what 473 was found in other remote sites (Freney et al., 2011; Raatikainen et al., 2010). 474 Furthermore, the origin of OA has been recently investigated in the Mediterranean 475 forested area of Montseny, and it has been found that SOA accounted for 91% 476 (Minguillón et al., 2011). The low contribution of primary organic components at MSC is 477 in agreement with the location, since the primary organic emissions are mixed and 478 oxidized during their transport from industrial and urban areas to the remote site of 479 MSC.

480 The diurnal cycles of OA components were studied as a function of air mass 481 origin (Fig. 6 and Fig. 7). A clear daily pattern of OA components was found regardless 482 of the air mass origin, except for Atlantic advections in winter. In summer, the 483 maximum concentrations of LV-OOA and HOA were measured between 12:00 and 484 13:00 UTC, whereas those of SV-OOA were observed between 11:00 and 12:00 UTC. 485 The LV-OOA has been generally associated with highly oxidized, aged, and long-486 range-transported aerosol particles (Lanz et al., 2010). Conversely, the SV-OOA has 487 been described as the less oxygenated and semi-volatile fraction of OOA (Ng et al., 488 2011a) and therefore it has been mostly attributed to SOA formation from more local emissions (Jimenez et al., 2009). For this reason, the LV-OOA and HOA hourly
variations are more influenced by long-range transport than those of SV-OOA, which
are strongly influenced by local/regional processes. In winter, the maximum
concentrations of the OA components were observed simultaneously around 14:00
UTC. The different daily patterns between seasons can be attributed to the higher
production of SOA in summer as opposed to winter, when the maximum daily
concentrations are reached later driven by the mountain breezes.

#### 496 **4 Conclusions**

497 This work interprets the real-time variation of inorganic and organic submicron 498 components during 10 months (July 2011 - April 2012) at a high altitude site in 499 southern Europe (Montsec, 1570 m a.s.l.). The aerosol chemical composition was 500 obtained with an ACSM, and co-located on-line and off-line PM1 measurements were 501 also carried out. The average concentration of the ACSM plus BC mass during this 502 study was 4.9  $\mu$ g m<sup>-3</sup>, and on average OA was the foremost PM<sub>1</sub> constituent (50%), 503 followed by sulfate (20%), nitrate (14%), ammonium (12%), BC (4%) and chloride (1%). 504 Discrepancies of OA determined by ACSM with co-located measurements pointed to 505 an overestimation by the ACSM probably caused by the use of the default RIE for OA, 506 which could be lower than the actual one. Further research is needed to better address 507 this issue.

508 The seasonal variation of PM<sub>1</sub> mass and chemical components concentrations 509 showed similar patterns, with an increase in summer and a decrease in winter, except 510 for nitrate which has high volatility in summer. The seasonal variation was attributed to 511 the evolution of the PBL height throughout the year and to synoptic circulation and 512 meteorological factors. At MSC the higher temperature and solar radiation in summer 513 enhances the convection processes, incrementing the development of the PBL, and 514 augments atmospheric photochemistry, promoting the formation of secondary inorganic 515 and organic aerosols.

516 The diurnal variation of PM<sub>1</sub> components had no clear diurnal patterns in 517 summer, except for organics. This lack of defined daily patterns was ascribed to the 518 recirculation of air masses that causes the formation of reservoir layers at relatively 519 high altitudes, and to the long-range transport from North Africa. These factors result in 520 a high variability of diurnal cycles even within the same type of episode. Nevertheless, 521 organic concentrations had a marked diurnal cycle regardless of the air mass origin, with maximum concentrations around 12:00 UTC. The OA was dominated by LV-OOA 522 523 (64%), followed by SV-OOA (26%), and HOA (10%). Hence, the midday increase with 524 respect to average concentrations during the night was attributed to the formation of525 SOA.

In winter under regional conditions, concentrations of all PM<sub>1</sub> components showed much clearer diurnal patterns than in summer, with a maximum between 14:00 and 15:00 UTC. These daily cycles were connected to the fact that MSC is located most of the day within the FT, whereas PBL air masses are only injected upwards after midday. However, when MSC was affected by long-range transport from mainland Europe, less marked daily patterns of PM<sub>1</sub> components were observed.

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The OA in winter was also mainly secondary (71%), with contributions from BBOA (24%), and HOA (5%). The hourly variation of these factors showed a clear diurnal pattern regardless of the air mass origin, except for Atlantic advections.

535 To the authors' knowledge, this is one of the first times when real-time 536 submicron aerosol chemical composition is characterized and its variation is interpreted 537 during almost a year in a continental background environment. The results obtained in 538 the present study highlight the importance of the SOA formation processes at such 539 remote site as MSC, which could be the objective of further investigations.

540

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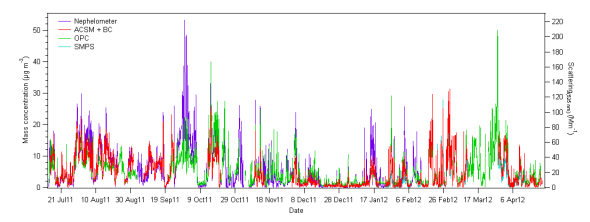
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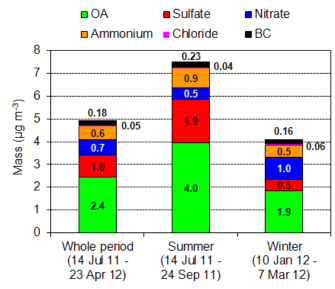
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Fig. 1 Time series of PM<sub>1</sub> total mass from co-located measurements and light scattering
at 525 nm.



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895 Fig. 2 Average concentrations of PM<sub>1</sub> chemical species measured at Montsec during the

896 whole study, in summer and in winter.

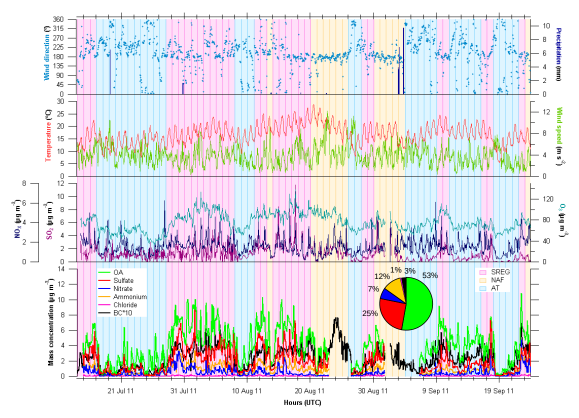
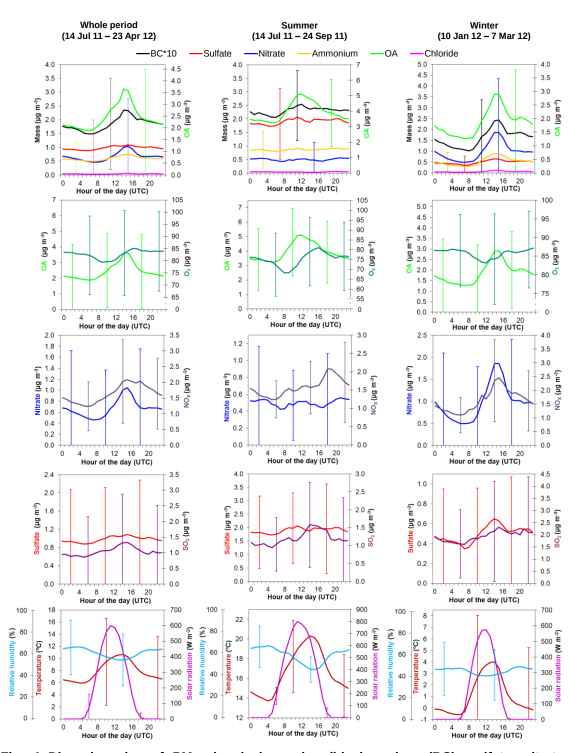
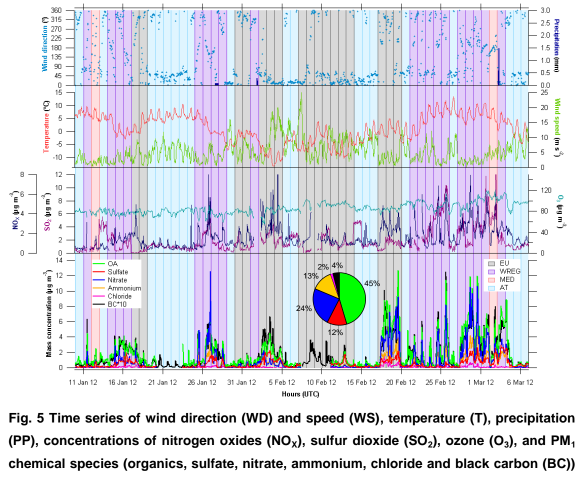


Fig. 3 Time series of wind direction, precipitation, temperature, wind speed, concentrations of nitrogen oxides ( $NO_x$ ), sulfur dioxide ( $SO_2$ ), ozone ( $O_3$ ), and  $PM_1$ chemical species (OA, sulfate, nitrate, ammonium, chloride and black carbon (BC)) in summer (14 Jul 11 – 24 Sep 11). Background colors correspond to daily classification of atmospheric episodes (summer regional (SREG), North African (NAF), and Atlantic (AT)) and the pie chart correspond to the average chemical composition for the summer period.



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Fig. 4 Diurnal cycles of  $PM_1$  chemical species (black carbon (BC), sulfate, nitrate, ammonium, chloride and OA), gaseous pollutants (ozone (O<sub>3</sub>), nitrogen oxides (NO<sub>x</sub>), and sulfur dioxide (SO<sub>2</sub>)), and meteorological parameters (relative humidity, temperature and solar radiation) averaged for the whole period, summer and winter. Variation bars indicate ± standard deviation.



917 in winter (10 Jan 12 – 7 Mar 12). Background colors correspond to daily classification of

- 918 atmospheric episodes (European (EU), winter regional (WREG), Mediterranean (MED) and
- 919 <u>Atlantic (AT)</u> and the pie chart correspond to the average chemical composition for the
   920 winter period.

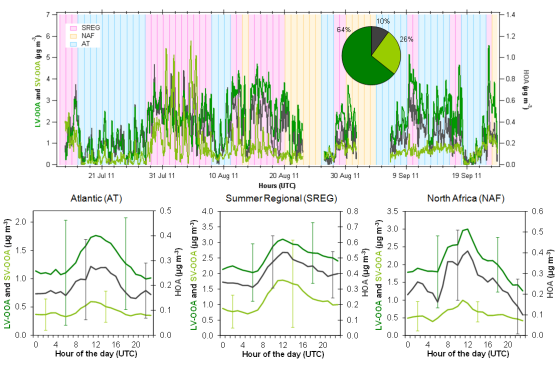


Fig. 6 Top: Time series of organic species (hydrocarbon-like organic aerosol (HOA),
semi-volatile oxygenated organic aerosol (SV-OOA) and low-volatility oxygenated
organic aerosol (LV-OOA)) concentrations in summer (14 Jul 11 – 24 Sep 11).
Background colors correspond to daily classification of atmospheric episodes and the
pie chart correspond to the average organic species composition for the summer period.
Bottom: Diurnal cycles of organic species concentrations averaged as a function of
atmospheric episode for the summer period. Variation bars indicate ± standard deviation.

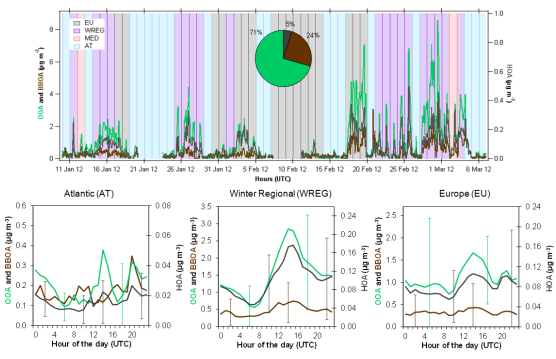


Fig. 7 Top: Time series of organic species (hydrocarbon-like organic aerosol (HOA), biomass burning organic aerosol (BBOA) and oxygenated organic aerosol (OOA)) concentrations in winter (10 Jan 12 – 7 Mar 12). Background colors correspond to daily classification of atmospheric episodes and the pie chart correspond to the average organic species composition for the winter period. Bottom: Diurnal cycles of organic species concentrations averaged as a function of atmospheric episode for the winter period. Variation bars indicate ± standard deviation.