Vertical and horizontal variability of PM₁₀ source contributions in Barcelona during SAPUSS

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1 Abstract

During the SAPUSS campaign (Solving Aerosol Problems by Using Synergistic 2 Strategies) PM₁₀ samples at twelve hours resolution were simultaneously 3 collected at four monitoring sites located in the urban agglomerate of Barcelona 4 (Spain). A total of 221 samples were collected from 20 September to 20 5 October 2010. The Road Site (RS) site and the Urban Background (UB) site 6 were located at street level, whereas the Torre Mapfre (TM) and the Torre 7 Collserola (TC) sites were located at 150 m a.s.l. by the sea side within the 8 9 urban area and at 415 m a.s.l. 8 km inland, respectively. For the first time, we are able to report simultaneous PM₁₀ aerosol measurements allowing us to 10 study aerosol gradients at both horizontal and vertical levels. The complete 11 chemical composition of PM₁₀ was determined on the 221 samples, and factor 12 analysis (Positive Matrix Factorisation, PMF) was applied. This resulted in eight 13 14 factors which were attributed to eight main aerosol sources affecting PM₁₀ concentrations in the studied urban environment: (1) vehicle exhaust and wear 15 $(2-9 \ \mu g \ m^{-3}, 10-27\% \ of \ PM_{10} \ mass \ on \ average), \ (2) \ road \ dust \ (2-4 \ \mu g \ m^{-3}, \ 8-3)$ 16 12%), (3) mineral dust (5 µg m⁻³, 13-26%), (4) aged marine (3-5 µg m⁻³, 13-17 20%), (5) heavy oil (0.4-0.6 μ g m⁻³, 2%), (6) industrial (1 μ g m⁻³, 3-5%), (7) 18 sulphate (3-4 μ g m⁻³, 11-17%) and (8) nitrate (4-6 μ g m⁻³, 17-21%). Three 19 20 aerosol sources were found enhanced at the ground levels (confined within the urban ground levels of the city) relative to the upper levels : (1) vehicle exhaust 21 22 and wear (2.8 higher), (2) road dust (1.8 higher) and (3) local urban industries / crafts workshops (1.6 higher). Surprisingly, the other aerosol sources were 23 relatively homogeneous at both horizontal and vertical levels. However, air 24 25 mass origin and meteorological parameters also played a key role in influencing

the variability of the factors concentrations. The mineral dust and aged marine 1 2 factors were found to be a mixture of natural and anthropogenic components and were thus further investigated. Overall, three types of dust were identified to 3 affect the urban study area: road dust (35% of the mineral dust load, 2-4 μ gm⁻³ 4 on average), Saharan dust (28%, 2.1 µgm⁻³) and background mineral dust 5 (37%, 2.8 µgm⁻³). Our results evidence that although the city of Barcelona 6 broadly shows a homogeneous distribution of PM₁₀ pollution sources, non-7 exhaust traffic, exhaust traffic and local urban industrial activities are major 8 coarse PM₁₀ aerosol sources. 9

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11 **1 Introduction**

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The study of ambient particulate matter (PM) concentrations in urban 13 environments is of particular interest due to its adverse effects on human health 14 (Pope and Dockery, 2006; Pope et al., 2009; Lim et al., 2012). However, the 15 identification and quantification of PM sources - the basis for the design of 16 mitigation strategies - remains challenging due to their complex nature. Indeed, 17 aerosols have multiple sources, and emissions from the same source will 18 change with time and operating conditions. Source apportionment of airborne 19 particulate matter has assumed increasing importance in recent years (Viana et 20 al., 2008; Beddows et al., 2015). This is because current legislation has 21 highlighted the need for reliable quantitative knowledge of the source 22 apportionment of particulate matter in order to devise cost-effective abatement 23 24 strategies. To apportion PM sources, many tools have been used for their

identification and quantification (Viana et al., 2008). Chemical speciation of 1 2 ambient PM coupled to receptor modelling is currently considered one of the most powerful tools for this purpose (Srimuruganandam and Shiva Nagendra, 3 2012; Waked et al., 2014). Within factor analytical models, the positive matrix 4 factorization (PMF) one is highly recommended when sources are not formally 5 known (Paatero and Tapper, 1994). PMF has been used in many studies 6 7 dealing with air pollution in urban areas in America, Europe, and Asia. Many sources such as road transport, industrial emissions, sea salt, and crustal dust 8 were identified using this method (Viana et al., 2008). Apportioning correctly the 9 10 sources of airborne particulate matter is important because there are most probably differences in the toxicity of particles according to their chemical 11 composition. In other words, PM from different sources may have a very 12 13 different potency in affecting human health (Kelly and Fussell, 2012).

Urban PM₁₀ concentrations show significant variability across Europe. In 14 northern European countries, the road dust originated by pavement abrasion 15 due to the use of studded tires (Norman and Johansson, 2006), high sea salt 16 concentrations in the coastal areas (Yin et al., 2005) and the use of biomass 17 burning for heating purposes are significant sources of PM (Puxbaum et al., 18 2007). By contrast, in southern European countries - due to the drier climate 19 conditions - mineral dust, resuspension and Saharan dust outbreaks 20 substantially contribute to PM concentrations (Putaud et al., 2010; Kassomenos 21 22 et al., 2014; Amato et al., 2015). Broadly, the elevated concentrations usually observed in Southern Europe are attributed to the combination of diverse 23 emission sources including industry, traffic, resuspended dust, shipping 24 25 emissions and African dust outbreaks. Recently, a huge effort has been made

in assessing the PM trends in the Mediterranean basin, including the MED-1 2 PARTICLES project (Karanasiou et al., 2014) and the AIRUSE LIFE project (Amato et al., 2015). It is important to remember that most of the PM urban 3 studies are based on ambient measurements taken at a single sampling point in 4 a city, but the pollutants concentrations may vary across the city. On this regard 5 it is worth reporting the studies of the ESCAPE project (Eeftens et al., 2012) 6 7 which aimed to investigate long-term effects on human health of exposure to air pollution in Europe, and showed large spatial variability of trace elements and 8 sources for improved exposure assessment (Minguillon et al., 2014). 9

The work presented in this manuscript is part of the FP7-PEOPLE-2009-10 IEF SAPUSS project (Solving Aerosol Problem by Using Synergistic 11 Strategies), where for the first time in the Mediterranean basin both the spatial 12 vertical and horizontal distributions of air pollutants were investigated (Dall'Osto 13 et al., 2013a). SAPUSS allows to better understand the complex interactions 14 between these pollutants and different meteorological variables, as well as their 15 influence on air quality. As reported by Han et al. (2015), with the increase of 16 vertical height the influence of source emissions on local air quality is 17 weakening, but the characteristics of regional pollution gradually become 18 obvious. The novelty of SAPUSS relies in the fact that by simultaneously 19 20 measuring across the vertical and horizontal urban scale, local and regional PM sources can be better apportioned. Unfortunately, there are very few studies 21 22 conducted in European urban vertical columns specifically looking at chemically resolved aerosol sources (Ferrero et al., 2010; 2014; Harrison et al., 2012; 23 Curci et al., 2015). A larger number of aerosol vertical studies is found in Asia, 24 25 perhaps because a larger number of people live in high-rise buildings. For

example, an estimated 57.7% of the 6.5 million residents in the Taipei 1 2 metropolis lived on or above the third story (Wu and Lung, 2012). Most studies of vertical gradients of PM in urban environments consisted of a tower with 3 aerosol and meteorological instrumentation deployed at different levels, up to 4 320m (Chan et al., 2005; Tao et al., 2007; Zhang et al., 2011; Shi et al., 2012; 5 Xiao et al., 2012; Öztürk et al., 2013; Sun et al., 2013; Tian et al., 2013; 6 Moeinaddini et al., 2014; Wu et al., 2014). Higher concentrations of primary 7 emitted contaminants were found at the lower levels, reflecting the closeness to 8 the emission sources (Tian et al., 2013). However, at some sites high PM 9 10 concentrations were recorded at the top level due to long-range transport of secondary pollutants (Harrison et al., 2012; Shi et al., 2012; Xiao et al., 2012; 11 Sun et al., 2013; Moeinaddini et al., 2014). 12

Atmospheric aerosol characteristics in the city of Barcelona and its 13 surrounding area have been studied in great detail. The main sources of PM₁₀ 14 are mineral dust sources and road traffic emissions (Amato et al., 2009; Pérez 15 et al., 2010). Other minor sources comprise shipping, both due to the city 16 harbour and regional sources in the Mediterranean Basin (Querol et al., 2009; 17 Pey et al., 2010). Industry represents usually less than 10% of PM₁₀ mass, 18 being V and Ni common tracers both for shipping and industrial activities 19 (Querol et al., 2009; Viana et al., 2014). Biomass burning contribution to PM_{10} is 20 relatively low in the study area (Minguillón et al., 2011; Reche et al., 2012). 21 22 Secondary aerosol components affect PM₁₀ especially during regional recirculation episodes, were the stagnant conditions lead to the accumulation of 23 pollutants (Pandolfi et al., 2014). In addition, due to the coastal location of the 24 25 study area, sea breeze (during the day) and mountain breeze (during the night)

also influence pollution transport from/towards the urban area. However, the
urban vertical column of Barcelona - and other Mediterranean cities - are not
characterised at all. Within the scope introduced in the presenting overview
paper of this special issue (Dall'Osto et al., 2013a), the main objectives of this
work are:

To interpret the variability of aerosol levels and composition in the urban
 Mediterranean environment of Barcelona, the second largest city in Spain and a
 major metropolitan agglomerate in Europe.

9 - To study aerosol particle mass in terms of the sources and the physico10 chemical transformations occurring simultaneously at the road, background,
11 tower and regional background sites.

To apply receptor modelling to PM₁₀ organic and inorganic species, in
 order to allow the identification of emission sources and their respective
 contributions to the PM mass in ambient air.

Our unique approach relies in the fact that - for the first time - both horizontal and vertical PM₁₀ measurements were taken simultaneously in four different sites across the urban agglomerate of Barcelona. Special emphasis is given in describing gradients ground/ tower and the variability of each factor concentrations under different air masses.

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1 2 Methodology

2 **2.1 Location**

Measurements were performed in the urban agglomerate of Barcelona, a 3 4 city located in the NE of Spain in the Western Mediterranean Basin (WMB). It is geographically constrained by the Llobregat and Besòs valleys (to the SW and 5 NE, respectively) and the coastal range of Collserola to the N. The city has a 6 population of 1.7 million, and the metropolitan area exceeds 4 million. These 7 conditions result in a highly populous urban area and one of the highest car 8 densities in Europe (6,100 cars km⁻², DGT 2015). More detailed information 9 about the SAPUSS study area can be found elsewhere (Dall'Osto et al., 2013a). 10 The structure of the planetary boundary layer (PBL) above Barcelona was 11 12 monitored by simultaneous measurements of ceilometers (Pandolfi et al., 2013). The field campaign took place in Barcelona from 20 September to 20 October 13 2010, including six sampling sites (Dall'Osto et al., 2013a). From those, the 14 following four monitoring sites are considered for the present study: 15

• Road Site (RS): It represents the average conditions of a trafficked road in the city centre (about 17,000 vehicles per day). The monitoring site was located in the Urgell Street, a street canyon with four vehicle lanes (one direction) and cycling lanes in both directions.

Urban Background (UB): it represents the urban background
 environment of Barcelona. It is located NW of the city centre in a small park 300
 m away from the busy Diagonal Avenue, a nine-lanes road used primarily by
 commuters (62, 000 vehicles per day).

Torre Mapfre (TM): It is a skyscraper located 300 m from the coast in the
Olympic Port of Barcelona, close to a recreational harbour and leisure area.
There is a tunnelled motorway ring road (four lanes) at 50 m distance from the
building and two three-lanes roads at ground level. The measurements were
taken at the rooftop terrace of the tower (150 m a.s.l.).

Torre Collserola (TC): This site was found at the Fabra Observatory (415 6 7 m a.s.l.), about 450 m from the Collserola tower (TC), where additional instrumentation was deployed (see Dall'Osto et al., 2013a). Due to logistical 8 9 reasons at the TC site (limited access and storage space), the Fabra Observatory (TC_g) was used as a monitoring site for the PM₁₀ aerosol chemical 10 measurements. In this study the TC_g is named TC for simplicity. Measurements 11 in this monitoring site were taken at ground level (10m above ground) but at an 12 overall height of 425 m a.s.l. 13

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It is important to note that whilst TM is well within the Barcelona urban city 15 centre, TC is located in the hills of the urban background of Barcelona. It is 16 worthwhile to stress that the two monitoring towers are the tallest buildings 17 within several kilometres of the sites, with good exposure to winds from all 18 directions (Dall'Osto et al., 2013a). The mean ceilometer surface mixed layer 19 (SML) and decoupled residual/convective layer (DRCL) heights over the whole 20 SAPUSS campaign were found to be 904±273 m a.g.l. and 1761±363 m a.g.l., 21 respectively (Pandolfi et al., 2013), all well above all the four measurements 22 sites described in this study. PBL maximum height (H) and daily variations (DV) 23 were strongly dependent on air mass types, ranging from the highest H -24 strongest DV during the Atlantic air masses to the lowest H - weakest DV 25

during the North African air masses, but overall always above both
 measurements towers and other monitoring sites.

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4 2.2 Measurements

High volume samplers DIGITEL DHA-80 and MCV CAV-A/mSb $(30m^3 h^{-1})$ equipped with PM₁₀ heads collected 12 h samples (from 11:00 to 23:00 and from 23:00 to 11:00, local time) on quartz fibre filters (Pallflex 2500QAT-UP) at the four monitoring sites. A total of 221 filter samples were collected, from which 93% were collected simultaneously at the four monitoring sites (54 concurrent samples per site).

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Meteorological parameters (temperature, relative humidity, wind speed and direction, solar radiation and pressure) were measured at the same sampling sites or at the nearest available meteorological station, as described elsewhere (Dall'Osto et al., 2013a).

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17 **2.2.1 PM concentration and chemical composition**

PM₁₀ mass concentrations were determined gravimetrically. The samples were analysed following the methodology described by Querol et al. (2001). Briefly, a quarter of the filter was acid digested (HNO₃:HF:HCIO₄), and the resulting solution was analyzed for Al, Ca, K, Mg, Fe, Ti, Mn, P, S, Na and 25 trace elements by inductively coupled plasma atomic emission and mass spectrometry (ICP-AES and ICP-MS), respectively; a quarter of the filter was water-extracted and the concentrations of SO₄²⁻, NO₃⁻ and Cl⁻, and NH₄⁺ were

determined by ion chromatography (IC) and selective electrode, respectively. A
section of 1.5 cm² of the filter was used to determine organic carbon (OC) and
elemental carbon (EC) by a thermal–optical transmission technique (Birch and
Cary, 1996) using a Sunset Laboratory OCEC Analyzer following the EUSAAR2
temperature protocol (Cavalli et al., 2010). Laboratory and field blanks were
analysed following the same procedure. Ambient concentrations were
calculated based on the samples and the blanks concentrations.

8 Crustal and sea salt aerosols concentration were also estimated. Moreno et al. (2006) reported the average composition of mineral dust originated in the 9 10 North African region that later reached the WMB. Based on the average Na/AI ratio of the North African dust, the mineral Na (or non-sea salt Na, nss Na) can 11 be calculated from the Al concentrations, hence the remaining Na is attributed 12 13 to sea salt (ss Na). The sea salt load for each sample was calculated based on the ss Na and the standard sea salt composition which includes Na, Cl⁻, ss Mg, 14 ss Ca and ss SO_4^{2-} (Mészáros, 1999). 15

Mineral matter was then calculated as the sum of SiO₂, CO₃²⁻, Al₂O₃, nss Ca, Fe₂O₃, K₂O, nss MgO and nss Na₂O. SiO₂ concentrations were estimated as $3*Al_2O_3$. CO₃²⁻ concentrations were estimated as 1.5*Ca, assuming that all Ca is present as CaCO₃ (Karanasiou et al., 2011). Organic matter (OM) was estimated as OC multiplied by a factor 1.4 at the RS, 1.6 at the UB and TM and 2.1 at the TC, according to Turpin and Lim (2001).

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1 **2.2.2 Source apportionment**

A constrained Positive Matrix Factorization (PMF, Paatero and Tapper, 1994) model was applied using the Multilinear Engine (ME-2) (Paatero, 1999) to assess the source apportionment.

5 PMF is a widely used receptor model based on the mass conservation 6 principle:

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$$x_{ij} = \sum_{k=1}^{p} g_{ik} f_{jk} + e_{ij} \quad i=1,2,...,m \quad j=1,2,...,n \quad (1)$$

8 Where x_{ij} is the i^{th} concentration of the species j, g_{ik} is the i^{th} contribution of the 9 source k and f_{jk} is the concentration of the species j in source k, and e_{ij} are the 10 residuals. Equation (1) can be also expressed in matrix form as $X=GF^{T}+E$. 11 PMF solves equation (1) minimizing the object function Q:

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$$Q = \sum_{i=1}^{n} \sum_{j=1}^{m} (e_{ij} / s_{ij})^2$$
 (2)

13 Where s_{ii} are the individual data uncertainties. The uncertainty estimates were based on the approach by Escrig Vidal et al. (2009) and Amato et al. (2009) and 14 provided a criterion to separate the species which retain a significant signal 15 from the ones dominated by noise. This criterion is based on the signal-to-noise 16 S/N ratio defined by Paatero and Hopke (2003). Species with S/N < 2 are 17 generally defined as weak variables and downweighted by a factor of 3. 18 Nevertheless, since S/N is very sensitive to sporadic values much higher than 19 20 the level of noise, the percentage of data above detection limit was used as complementary criterion. All the samples collected at the four sites were 21 gathered in one data matrix. This data assembling allows exploring a larger 22

area of the N-dimensional source contributions space. The data matrix was 1 2 uncensored, i.e. negative, zero and below detection limit values were included as such in the analyses to avoid a bias in the results (Paatero, 2007). A total of 3 221 samples containing 32 different species were included in the PMF which 4 was run by means of the Multilinear Engine-2 program allowing to handle a 5 priori information as shown in the Results section. A bootstrap technique was 6 used to estimate the uncertainties of factor profiles, based on the EPA PMF 7 v3.0 script. It consisted on three different steps: re-sampling, reweighting and 8 random rotational pulling (Tukey, 1958; Efron and Tibshirani, 1986). A seed 9 10 value of 7 was used with 20 runs. Rotational ambiguity was reduced by means of the implementations of the constraints and evaluated through bootstrapping. 11 The final number of factors was chosen based on several criteria: Q values, 12 13 factors profiles and contributions, distribution of scaled residuals and G space plots. 14

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16 3 Results

3.1 PM₁₀ concentration and chemical composition in a three dimension
 (3D) scenario

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As presented in the introduction overview paper of this special issue (Dall'Osto et al., 2013a), the average PM_{10} concentrations at the four monitoring sites during SAPUSS followed a decreasing trend from the site closest to traffic sources to the one located in the suburban background at 415 m a.s.l. (RS 30.7, UB 25.9, TM 24.8 and 21.8 µgm⁻³ at TC; Figures 1 and 2).

This suggests that the city surface is enhanced in coarse particles. It is 1 important to note that the high PM₁ /PM₁₀ ratios at the towers (TM, TC) are not 2 only due to the low PM₁₀ concentrations, as the absolute values of PM₁ 3 detected at the tower sites were also higher than the urban background ground 4 levels. Deng et al. (2015) recently also reported that - overall - the vertical 5 distributions of PM₁₀, PM_{2.5}, and PM₁ decreased with height. The lapse rates 6 showed the following sequence: $PM_{10} > PM_{2.5} > PM_1$, which indicates that the 7 vertical distribution of fine particles is more uniform than that of coarse particles; 8 and the vertical distribution in summer is more uniform than in other seasons. 9

Whilst the chemical analysis of the PM mass was only briefly described in an earlier study (Dall'Osto et al., 2013a), in this section the chemical components building up the PM₁₀ mass concentrations are described in more details. The concentrations of each analysed species is shown in Table S1. Fig. 2 and Fig. 3 show the absolute and relative contributions to PM₁₀ at each site. Briefly:

Crustal elements accounted for 15-18% of the mass, suggesting an unexpectedly homogeneous relative contribution to the total PM₁₀ across the Barcelona urban area (Fig. 2). However, absolute mass concentrations of crustal matter (Fig. 3a) decreased from the city centre sites (5.5 µgm⁻³ at RS) to those located further inland (3.2 µgm⁻³ at TC), suggesting higher contribution of dust in the lower level monitoring sites.

Sea salt aerosols appeared aged due to the robust lower Cl⁻/Na and
 higher SO₄²⁻/Na ratio than sea water composition (Henderson and
 Henderson, 2009) and the relative contribution was homogenous at all

sites ranging from 7-11% (1.5-2.7 µgm⁻³, Fig. 2). Fig. 3b shows that the
highest concentrations were recorded near the sea side monitoring sites
(TM, RS), with a decreasing gradient recorded towards inland (UB, TC).
Despite TM being located a few meters from the sea side, it shows
similar concentrations to RS, 4 km distant. This might be due to the
elevation of the TM sampling site (150 m a.s.l.) causing dilution of
surface sea spray.

Secondary inorganic aerosols (SIA: sulphate SO42-, nitrate NO3 and 8 ammonium NH₄⁺) display altogether similar concentrations at the city 9 sites (6.3 μ gm⁻³ on average at RS, 5.7 μ gm⁻³ at UB and 5.8 μ gm⁻³ at TM) 10 and decrease around 25% at TC (4.4 µgm⁻³). Overall, SIA accounted for 11 20-23% of the mass (4.4-6.3 µgm⁻³, see Fig. 2). The contribution of 12 ammonium was similar at all sites, around 2-3% (0.5-0.9 µgm⁻³) of PM₁₀ 13 mass. Sulphate and nitrate contributed in similar proportions at all sites, 14 ranging from 7-11% each (1.6-2.6 μ gm⁻³ for nitrate and 2.3-2.8 μ gm⁻³ for 15 sulphate), although in absolute concentrations a decrease of 15-35% at 16 the TC was detected for SIA (see Fig. 3c). 17

Elemental carbon (EC) concentration was clearly influenced by traffic sources, as its contribution decreased with the distance to traffic hot spots (RS: 1.4 µgm⁻³, 5%; UB: 0.9 µgm⁻³, 3%; TM: 0.7 µgm⁻³, 3%; TC: 0.5 µgm⁻³, 2% - see Fig. 2 and 3d).

Organic matter (OM) accounted for 16-21% of the mass and was found
 in the highest relative proportion at the TC site due to its location on a
 nearby hill in a suburban environment. The lowest concentrations were

observed at UB and TM (4.2 μ gm⁻³), followed by TC (4.6 μ gm⁻³), while the highest ones were recorded at the RS (5.2 μ gm⁻³ - see Fig. 2 and 3 3d).

The unaccounted mass of PM (31-35%) is that resulting from the 4 difference of the gravimetric measurements of the filters and the sum of 5 all the components determined by chemical analysis. This unaccounted 6 mass is usually attributed to water molecules contained in potential 7 remaining moisture, and crystallisation and formation water (water 8 molecules in the structure of specific chemical species), as well as 9 heteroatoms contained in the organic species and not analysed (Querol 10 et el., 2001; Amato et al., 2015). 11

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Despite the low contribution of trace elements to the bulk PM_{10} (<1%), 13 their concentrations provide valuable information by tracing specific pollution 14 sources. Figure 3e-j show the concentration of a selection of trace elements 15 associated with four known sources affecting the metropolitan area of 16 Barcelona: crustal, road traffic, industry and shipping. The most abundant 17 crustal tracer is Ti followed by far by Rb, Y, Ce, La, Li, Nd, and Ga. Such 18 elements were found in higher concentrations at the ground sites (RS and UB) 19 relative to the tower sites (TM and TC). This might be due to the dust 20 resuspension by road traffic or other anthropogenic sources such as 21 construction works or parks, thus producing a vertical decreasing gradient. The 22 spatial variability of vehicle wear tracers such as (Cu, Sn and Ba; Schauer et 23 al., 2006) is driven by the proximity to traffic sources, being the RS the most 24

polluted site. The industrial emissions tracers for the city of Barcelona (Zn, Mn, 1 2 Pb, As and Ni; Amato et al., 2009) showed higher concentrations at the UB followed by the RS, TM and TC. This is likely to be due to the pollution plumes 3 originating from the industrial area along the Llobregat Basin (south-western 4 side of Barcelona metropolitan area) that first impacts the UB site (Dall'Osto et 5 al., 2013b). On the other hand, the remaining industrial tracers Cu and Sb 6 (showing a decreasing gradient with the distance to traffic sources) might 7 account for both industrial and traffic-related emissions. Both traffic and 8 industrial tracers showed significant higher values at the ground sites respect 9 10 the tower sites. On this regards, it is worth to remember that V and Ni (Fig. 3i-j) are usually associated with shipping emissions (Kim et al., 2008; Agrawal et al., 11 2008). Indeed, higher V levels were found at the sites closer to the coast (TM 12 13 and RS), in agreement with findings from Minguillón et al. (2014). However, higher Ni concentrations were recorded at the ground sites (UB and RS) 14 respect to the tower sites (TM and TC), probably due to the contribution of other 15 Ni sources reaching the ground sites (e.g. smelters). Viana et al. (2014) 16 reported a V/Ni ratio associated to shipping emissions of 2.3-4.5. In our study 17 the only site within this range was TM (2.4), located closest to the sea side. RS 18 and UB showed lower ratios (1.3 and 1.1, respectively) probably due to the 19 higher impact of an industrial plume containing Ni. TC ratio was 2.0, which 20 might indicate an additional V loading of crustal origin, due to its location on a 21 nearby hill. 22

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3.2 Source apportionment and temporal variability in a 3D scenario

2 The PMF analysis was applied to the PM₁₀ data matrix, which contained 221 samples (56 at RS, 54 at UB, 55 at TM and 56 at TC). The method herein 3 used is the same of the one recently reported by Amato et al. (2015), aiming at 4 characterising similarities and heterogeneities in PM sources and contributions 5 in urban areas from Southern Europe. This method is not new, as it was already 6 proven to increase considerably the statistical significance of the analysis 7 8 (Amato et al., 2009). Different constraints were added into the PMF model (Paatero and Hopke, 2008), in order to reduce rotational ambiguity. The PMF 9 solution chosen was an eight factor solution, with a Q (constrained) value of 10 6,627, being the Q/Qexp ratio 1.17, and with an increase dQ of 7.5% with 11 respect to the base run, due to the implementation of auxiliary equations. 12

One factor (road dust) was pulled towards the composition of local road dust (average of city centre samples, as reported by Amato et al., 2009). The road dust emission profile was introduced by means of auxiliary equation (pulling equations, Paatero and Hopke, 2008), consisting in pulling a f_{jk} toward the specific target value *a*:

$$Q_{aux} = \frac{(f_{jk} - a)^2}{\sigma_{aux}^2}$$

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where σ_{aux} is the uncertainty connected to the pulling equation, which expresses the confidence of the user on this equation. In the present study, a pulling equation was used for each specie in order to pull concentrations of a factor toward the target concentration in the road dust emission profile as obtained by Amato et al, (2009). An average profile of 4 road dust samples

collected at 4 different points of the Diagonal Avenue was selected as a
representative emission profile. The Na/AI in the mineral factor ratio was
pulled towards the value reported for earth's crust composition by Mason (1966)
in form of auxiliary equation (Paatero and Hopke, 2009).

Overall, a total of eight factors were identified by the application of PMF: 5 (1) "vehicle exhaust and wear", (2) "road dust", (3) "mineral", (4) "aged marine", 6 (5) "heavy oil", (6) "industrial", (7) "sulphate" and (8) "nitrate". The PMF result 7 has proven to be very robust (the sum of the contributions of the chemical 8 9 species for each source are close to one, see Table S2) and the PM₁₀ mass has been well reconstructed by the PMF model (see Figure S1). The average 10 concentrations of each factor registered at each site are shown in Figure 5 and 11 12 Table 1, while Figure S2 displays the temporal variation of each factor for all sites. To further complete the analysis and interpretation of the results, Polar 13 Plots were obtained using the OPENAIR software package of R (Carslaw and 14 15 Ropkins, 2012; R Development Core Team, 2012). These plots display the different factors concentrations depending on the blowing wind direction and 16 speed, thus allowing deducing the main pollution sources origin (Fig. S3). The 17 eight identified aerosol PMF factors characteristics can be seen in Fig. 4 and 18 are summarised as: 19

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The vehicle exhaust and wear factor profile (Fig. 4a) was dominated by
 EC and OC originating from vehicle exhaust emissions. Other chemical
 elements include Cu, Sb, Cr, Fe and Sn (67%, 53%, 46%, 41% and 39%
 of the variation, respectively) which are usually present in the brake and

tyre wear (Sternbeck et al., 2002; Ntziachristos et al., 2007; Amato et al., 1 2009). Due to its direct traffic origin, this factor accounts for the highest 2 mass contribution at the RS (27%, 8.7 μ g m⁻³), followed by UB (18%, 5.0 3 ug m⁻³). TM (11%, 2.9 µg m⁻³) and TC (10%, 1.9 µg m⁻³) (see Table 1 4 and Fig. 4a). A clear horizontal and vertical gradients can be seen for the 5 PM₁₀ contributions of this source. It originated at the traffic hot spots near 6 RS, UB and TM and was later transported upslope towards TC by the 7 sea breeze, where the maximum concentrations were recorded under SE 8 winds from the city (Fig. S3). 9

The road dust PMF factor profile (Fig. 4b) was constrained using the 10 emission profile reported for the city of Barcelona by Amato et al. (2009) 11 by means of a pulling equation. It contained high concentrations of AI_2O_3 , 12 Ca, Fe, Li, Ti but also explained around 20% of the variation of Cu and 13 Sb (Table 1 and Fig. 4b). As expected, this factor concentration followed 14 also a decreasing trend from RS (12%, 3.8 μ g m⁻³) and UB (12%, 3.3 μ g 15 m⁻³) to TM (9%, 2.3 μ g m⁻³) and TC (8%, 1.6 μ g m⁻³). The road dust was 16 transported from the nearby busy streets towards the sites, as seen in 17 Fig. S3. 18

The mineral PMF factor profile (Fig. 4c) was mainly traced by of Al₂O₃,
 Ti, Rb, La, Li and Se (42%, 45%, 36%, 27%, 26%, 22% of the variation,
 respectively). Unexpectedly, average absolute concentrations were very
 homogeneous across the city (4.6 to 5.1 µg m⁻³, 13-26%) pointing to a
 source affecting the whole urban area but also due to the important and
 robust enrichment in S and C with respect to the average crust
 composition (Taylor,1964 and Mason, 1966) indicating the mixing of

mineral dust with regional/local plumes and the neutralization of
 sulphuric acid by mineral cations through heterogeneous reactions.

The aged marine PMF factor profile (Fig. 4d) contribution is 3 characterised by Na and CI (75% and 52% of the variation, respectively) 4 and also a proportion of Mg and SO_4^{2-} (48% and 22% of the variation, 5 respectively). The ratio SO_4^2 /Na exceeded the calculated sea salt ratio, 6 indicating its aged nature. As expected, the highest concentrations were 7 reached at the sites located closer to the sea (TM, 20%, 5.2 μ g m⁻³; RS, 8 14%, 4.6 µg m⁻³; UB, 13%, 3.6 µg m⁻³; TC, 13%, 2.6 µg m⁻³). The highest 9 concentrations at all sites were reached under E-SE blowing winds 10 (caused by sea breeze, Fig. S3). 11

The heavy oil PMF factor profile (Fig. 4e) was characterized by V and Ni 12 (71% and 45% of the variation, respectively) and showed a relevant 13 concentration of SO_4^{2-} and EC. It was attributed to fuel oil combustion 14 from shipping emissions since power generation around Barcelona is 15 only allowed by using natural gas since 2008. Furthermore, 98% of 16 domestic heating systems use natural gas, and the spatial distribution of 17 18 V concentrations evidenced higher levels as we approach the coast (Table S1). Average concentrations varied between 0.4-0.6 μ g m⁻³, 19 20 representing on average about 2% of the load at each site (Table 1).

The industrial PMF factor profile (Fig. 4f) was defined by Pb, Zn, Mn and
Cd (50%, 44%, 31% and 19% of the variation, respectively). It was
related to the smelters and cement kilns located along the nearby
Llobregat valley, NW of the city (Amato et al., 2009; Moreno et al., 2011,
Dall'Osto et al., 2013b). The emission plume was transported towards

the city by the night land breeze (Fig. S3), reaching the ground sites UB
and RS in a greater measure due to proximity of the sources (RS, 4%,
1.2 μg m⁻³; UB, 5%, 1.4 μg m⁻³; TM, 3%, 0.7 μg m⁻³; TC, 5%, 0.9 μg m⁻³).
The highest peaks (6-8 μgm⁻³) were recorded for the UB site the 5
October and the 10 October (Fig. S2), showing the highest
concentrations at all sites under NW winds (Fig. S3).

The sulphate PMF factor profile (Fig. 4g) was defined by SO₄²⁻ and NH₄⁺
(39% and 73% of the variation, respectively). OC is also present in
significant concentration (around 10%), suggesting the contribution of
secondary organic aerosols to this factor. As a consequence of its
regional origin and secondary nature, it shows homogeneous
concentration values at the four sites (3.3-4.2 µgm⁻³, 11-17%; see Table
1).

The nitrate PMF factor profile (Fig. 4h) was mainly traced by NO₃⁻ (97% of the variation), but NH₄⁺, OC and Cl⁻ also contributed in a minor proportion (10%, 11% and 6%, respectively). It also shows homogeneous concentration values at the three city sites (around 5.4 µgm⁻³), whereas at TC concentrations were 33% lower (3.6 µg m⁻³, see Table 1). This revealed a dominant local urban origin of this factor, as nitrate was diluted while being transported to the suburban area.

21

A nine factor solution was attempted and showed the same factors shown with the eight solution, with an additional nine one called "Se-SUL", composed mainly of Selenium and Sulphate and lacking of any clear temporal trends (see Supporting Information).

1

2 4. Discussion

3

4 4.1 Variability of aerosol sources across sites and ground-tower ratio

The distance to the emission source strongly influences the concentration 5 levels of certain PMF factors detected at the sampling sites. This is the case of 6 the exhaust and wear and road dust factors, presenting a decreasing 7 8 concentration gradient with the distance to the traffic hot spots, as the highest concentrations were found at the RS, followed by UB, TM and TC. Regarding 9 the marine factor, the distance to the sea influenced the average 10 11 concentrations, being highest at TM, followed by RS, UB and TC. In the case of 12 the industrial factor, whose main source is located SW of the city, the UB was the first site impacted by this plume as it showed the highest concentrations, 13 followed by RS, TC and TM. On the other hand, fairly homogeneous 14 concentrations were recorded at the sites for mineral and heavy oil factors, 15 evidencing that its sources affect the whole urban area. Regarding sulphate, 16 similar concentrations were recorded at the sites, although the higher 17 concentrations displayed at the UB site remain unexplained at this stage. 18 19 Concerning the nitrate factor, concentrations were found to decrease with the distance to traffic sources from RS to TC. However, at the TM site higher 20 concentrations than at the UB site were recorded due to the elevation of the 21 22 urban tower site above the ground favouring the formation of particulate nitrate due to colder temperature. Curci et al. (2015) also showed that an important 23 player in determining the upper planetary boundary layer (PBL) aerosol is 24

particulate nitrate, which may reach higher values in the upper PBL (up to 30%)
than in the lower PBL. Overall, the trends are in line with a recent study in a
Chinese Tower (Han et al., 2015), suggesting that the impact of primary
sources from the ground decreased with the increase of height, while the impact
of secondary sources mainly influenced by regional sources becomes more
prominent.

7 To further study the vertical variability in the factors concentrations, the ratios between the average concentrations at the "ground" sites (RS, UB) and 8 9 "tower" sites (TM, TC) were calculated for each factor, and the result are presented in Fig. 5. Three sources were found to be different between ground 10 and tower levels: exhaust and wear, road dust and industry. The highest 11 differences were found for the exhaust and wear and road dust factors, where 12 the concentrations at the ground sites were 2.8 and 1.8 times those at the 13 towers, respectively. The industrial factor concentration at the ground sites were 14 on average 1.6 times higher than at the towers, pointing towards the greater 15 impact of the SW industrial plume on ground levels; although a contribution of 16 the small industrial facilities spread within the city should not be discarded. On 17 the other hand, the remaining factors (mineral factor, aged marine and heavy 18 oil) showed similar contributions at ground and tower sites. A vertical 19 20 distribution for various chemical species was also previously reported in a number of other studies. For example, Han et al. (2015) recently showed similar 21 22 percentage levels at the four different heights for AI and Si. However, for the Ca and EC fractions, higher values were observed at lower sampling sites. The 23 percentages of nitrate, sulphate and OC, and the OC/EC ratios were higher at 24 25 the higher sites. Source apportionment for ambient PM₁₀ showed that the

percentage contributions of secondary sources obviously increased with height, 1 2 while the contribution of cement dust decreased with height. Ho et al., (2015) also reported that vertical variations were observed for mineral and road dust 3 (Si, Ti and Fe) in the PM_{2.5} region. Similarly, Wu et al. (2014) reported traffic 4 related aerosol (in particular resuspended road dust - traced with Si) and 5 industrial ground activities vertically stratified. In summary - consistent with this 6 SAPUSS study - exhaust traffic, non-exhaust traffic and industrial aerosol 7 sources were the ones mostly affecting the aerosol vertical gradients. 8

9

10 **4.2** Aerosol sources variability relative to air mass category

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12 The variability in PM₁₀ concentrations and air mass scenario during SAPUSS (according to the classification presented by Dall'Osto et al., 2013a) 13 shown in Fig. 1 was already briefly discussed in section 3.1. Fig. 6 shows the 14 average concentrations at each site under the five air mass scenarios identified 15 during SAPUSS: Atlantic (ATL), Regional (REG), North African West (NAF W), 16 North African East (NAF_E) and European (EUR). REG scenarios were related 17 to the recirculation of air masses over the study area, thus favouring the 18 19 accumulation of both primary (vehicle exhaust and wear, industrial) and secondary pollutants (sulphate, nitrate). Overall, concentrations were 32% 20 higher under these REG air masses due to low pollutants dispersion (Fig. 6). Air 21 22 masses with an African origin (NAF W and NAF E) were influenced mainly by the mineral factor. The NAF_E air mass also crossed the WMB, blowing 23 easterly winds inland and also causing an increase in aged marine aerosols 24

concentration. Under North African air masses (NAF_W and NAF_E), average 1 concentration levels nearly doubled with respect to the average levels for non-2 African days (9.2 µg m⁻³ vs 4.9 µg m⁻³, Fig. 5 and Fig. 6e-f). The cruise and 3 commercial port of the city is located south of the city (Fig. S3), and thus under 4 NAF_W air masses and S-SW winds the highest heavy oil concentrations were 5 recorded at all sites, pointing towards direct port emissions as the main 6 contributor to this factor. Road dust also showed the highest concentrations 7 during NAF E scenarios due to the increase of its loading and subsequent 8 resuspension. During the study period EUR air masses were related to a rainfall 9 10 event, thus wet deposition caused a radical decrease in road dust concentrations under this scenario. A decrease in concentrations of 90% on 11 average, due to wet deposition and less resuspension, was recorded under 12 13 EUR air masses, as they were associated with rain during the study period (0.3) μ g m⁻³ vs 2.7 μ g m⁻³, see Fig. 6c-d). On the other hand the aged marine factor 14 concentrations were found to increase, due to the trajectory followed by this air 15 mass over the Mediterranean Sea. ATL air masses were generally related to 16 low concentrations for the different factors (due to pollutants dilution) except for 17 18 the industrial factor, which might be explained similarly by the accompanying westerly winds under this scenario. 19

As discussed in Dall'Osto et al. (2013a), a number of possible REG stagnant different scenarios were classified during SAPUSS. As a case study, we consider two different ones: REG_1 (4 days between 29 September and 2 October) and REG_2 (4 days between 14-17 October). Figure S4 shows the meteorological diurnal profiles of the two scenarios. Whilst REG_1 shows warmer temperatures and a daily sea breeze circulation, REG_2 is

characterised by colder temperatures, more stagnant air and an absence of a 1 2 sea breeze circulation. Overall, high PM₁₀ concentrations were recorded under REG air mass due to the accumulation of pollutants. REG_1 presented 19-31 3 $\mu g m^{-3} PM_{10}$ average mass concentrations across the four sites, whereas 4 REG 2 showed even higher loadings (30-41 µg m⁻³). The REG_1 episode 5 allowed the transport of heavy oil towards the city with the development of the 6 sea breeze, whereas in the REG_2 episode the poor development of the sea 7 8 breeze minimized the transport of the shipping emissions towards the city (Fig. S4). It is interesting to note that during the REG_2 recirculation episode (14-17) 9 October), the nitrate PMF factor concentrations were doubled (10.7 vs 4.9 µgm⁻ 10 ³ overall SAPUSS average) at the four monitoring sites, reaching occasionally 11 higher levels at the tower sites (TM, TC) than at ground levels (RS, UB). By 12 13 contrast, the sulphate PMF factor did not show a larger variation among different REG scenarios. The PMF nitrate/ PMF sulphate ratio for was found to 14 be 1.2 and 2.3 for REG_1 and REG_2, respectively. As previously observed in 15 a vertical aerosol study in London (Harrison et al., 2012) the cooler 16 temperatures and higher relative humidity on the tower level during the REG_2 17 18 scenario can shift the gas/aerosol nitrate equilibrium towards the aerosol phase. In other words, during SAPUSS some aspects of nitrate behaviour were broadly 19 similar to those of sulphate, but other aspects proved very different. During 20 SAPUSS, Aerosol Time-Of-Flight Mass Spectrometer studies (Dall'Osto et al., 21 2013a) reported two types of nitrate aerosols. Briefly, the first appeared to be 22 associated with local formation processes and occurred at times outside of the 23 long-range transport episode. The second type of nitrate was regionally 24 transported and internally mixed with sulphate, ammonium and both elemental 25

and organic carbon (Dall'Osto et al., 2009). On this regards, it is worth to 1 2 remember that the nitrate radical (NO_3) is amongst the most important oxidants in the nocturnal boundary layer (NBL) (Benton et al., 2010). Little is known 3 about products between the formation of NO₃, its reactions with volatile organic 4 compounds (VOCs) and the formation of organic nitrate (Wayne et al., 1991; 5 Brown et al., 2009). The PMF method applied in this aerosol filter based PM₁₀ 6 concentrations shows OC being an important component (11%) for the PMF 7 nitrate factor, although 19% of the OC component was not described by the 8 PMF and found in the PMF residuals. It is likely that the high concentrations of 9 10 nitrate found in regional air masses during SAPUSS are a complex mixture of different types of aerosol nitrate, not been distinguished during this PMF 11 analysis and likely due to the poor time resolution (12 hours) of the off-line 12 aerosol filter techniques (Dall'Osto et al., 2013a). 13

14

15 **4.3 Additional aerosol source estimations**

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The PMF model was applied during this SAPUSS study and the results 17 were presented in section 3.2. However, PMF may not be able to separate 18 19 similar sources and, due to chemical reactions, apparently "natural" PMF factors like mineral and sea salt may also include anthropogenic contributions. In order 20 to elucidate the contributions to ambient PM₁₀ concentrations, a combination of 21 22 additional aerosol source estimation techniques were applied to further elucidate two main natural sources (mineral dust and sea salt sources) 23 contributing to the PM₁₀ mass during SAPUSS in Barcelona. 24

1

2 4.3.1 Mineral dust sources

3

Mineral sources in a Mediterranean urban environment are diverse. 4 5 Broadly, three main components of mineral dust have been reported in the literature: (1) urban-regional background dust, (2) local road dust and (3) 6 Saharan dust. Querol et al. (2001) reported a urban/regional background 7 mineral dust factor enriched in AI and Ca, which presented higher 8 concentrations in summer than in winter. A background source rich in Ca, Si, Al 9 10 and Ti was also attributed to regional anthropogenic and natural resuspension such as urban dust from construction/demolition works, unpaved areas and 11 parks, among other sources (Amato et al., 2009). Road dust is associated with 12 resuspended road dust by passing vehicles and wind, and is traced by Fe, Ca, 13 Al, Si, Ti, Cu, Sb, Sn, Ba, Zn, OC and EC (Schauer et al., 2006). The use of 14 15 constraints for the source apportionment PMF model by using pulling equations enabled to quantify the road dust fraction of the mineral dust. This PMF factor 16 was characterised by a clear decreasing concentrations gradient with the 17 distance to traffic sources and it contributed with 1.6 to 3.8 µgm⁻³. 18

Saharan dust outbreaks transporting dust (made of quartz, clays, calcium
carbonate and iron oxide and traced by Al, Si, Ti among others) regularly impact
the study area (Querol et al., 2001). Efforts have focused on quantifying this
contribution to the average mineral loading, both for air quality purposes (Querol
et al., 2009; Pey et al., 2013) and its impact on population's health (Pérez et al.,
2008). However, the PMF factor analysis could not efficiently separate Saharan

dust, background mineral and road dust. Hence, the methodology proposed by 1 2 Escudero et al. (2007) for estimating the Saharan dust daily contribution for different mass fractions was applied. Briefly, it consists in subtracting from the 3 average concentrations registered at the city (Barcelona) those ones 4 simultaneously measured at the nearest regional background site (Montseny, 5 720 m.a.s.l., 50 km NE of Barcelona). However, during SAPUSS the estimated 6 7 Saharan dust loadings calculated with this method often exceeded the real PM₁₀ concentrations registered at the SAPUSS sites. This is likely due to the 8 fact that Saharan dust outbreaks are different at the sea level Barcelona city 9 10 and its higher altitude regional background surrounding area (Escudero et al, 2007). Therefore, a different methodology was applied for subtracting the 11 Saharan dust load from the mineral factor. We calculated the in situ baseline of 12 13 mineral dust levels at each site during the Saharan outbreaks, taking into account the concentrations registered before and after the Saharan dust 14 episodes, and extracted these from the mineral dust load for each sample at 15 each site. The resulting concentration excedence of mineral dust was 16 interpreted as the Saharan dust contribution. Overall, it was found that the 17 18 average contribution of Saharan dust for the whole study period at the four sites was 2.1 μ gm⁻³ (28% of the PM₁₀ mineral load). Upon subtraction of the 19 estimated Saharan dust contribution at each site, the remaining mineral loading 20 corresponds to background mineral dust of urban or regional origin (2.7 to 2.9 21 μgm^{-3}). This narrow concentration range at the four sites (Fig. 7) -22 independently of the height and urban location - points towards a regional origin 23 of this background mineral matter. 24

In summary, during SAPUSS the three mineral dust sources (Fig. 7-9)
 could be summarised as follow:

1) Background dust: it presented a homogeneous distribution among the 3 sampling sites and was thus attributed to background mineral dust with a 4 possible urban or regional origin. A regional origin is thought to be more 5 6 probable due to the uniform distribution of this dust type at both horizontal and vertical levels for the whole study area. Average concentrations during 7 the SAPUSS study ranged from 2.7 to 2.9 µgm⁻³, resulting in the mineral 8 source with the highest contribution (37% of the mineral dust in PM₁₀ in the 9 study period). 10

2) Road dust: the concentrations decreased from RS to TC, contributing
3.8-1.6 µgm⁻³ on average (35% of the mineral dust in PM₁₀ during the study
period).

3) Saharan dust: African air mass incursions occurred on 20% of the days
during the study period. Under this scenario, the excess dust from the PMF
mineral factor was extracted and attributed to Saharan dust, thus obtaining
an average Saharan dust contribution of 2.1 µgm⁻³ (28% of the mineral dust
in PM₁₀ in the study period).

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- 20
- 21 4.3.2 Sea salt aerosols

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23 Sea spray aerosol is an important component of the aerosol population in 24 the marine environment, and given that 70% of the Earth's surface is covered

by oceans, it contributes significantly to the global aerosol budget (Vignati et al., 1 2 2010). Due to the high impact of anthropogenic activities on the WMB and the frequent recirculation of regional polluted air masses on the region, an 3 interaction between natural and anthropogenic sources is expected. Indeed, 4 22% of the variability of $SO_4^{2^2}$ was attributed to the aged marine PMF factor 5 (Fig. 4, Table S3), suggesting that this factor is internally mixed with 6 anthropogenic pollutants. The fresh sea salt is calculated as the sum of ssNa + 7 Cl + ssMg + ssCa+ ssSulphate. ssNa is calculated as the measured Na - nss 8 Na – Na from Na₂SO₄ in the PMF aged marine factor. The anthropogenic 9 marine sulphate is calculated as the difference between the PMF aged marine 10 and the fresh sea salt. The PMF aged marine factor (2.6-5.2 µgm⁻³) could be 11 broken down into calculated fresh sea salt (1.2-2.1 µgm⁻³, 40-47%) and 12 anthropogenic marine sulphate of regional origin (1.4-3.1 µgm⁻³, 53-60%). 13 These results evidence that both the calculated sea salt and the anthropogenic 14 marine sulphate aerosols contributed in a similar proportion to the aged marine 15 factor (Fig. 8, 9). The marine sulphate of anthropogenic origin derived from the 16 aged marine factor shows a different origin to the PMF sulphate factor. As can 17 18 be seen in Fig. S3 the highest concentrations of anthropogenic marine sulphate were recorded under eastern winds at all sites, whereas for the PMF sulphate 19 factor no dominant wind direction was found. Namely, the highest sulphate 20 factor concentrations were recorded under REG air masses while the 21 anthropogenic marine sulphate shows relatively concentrations. 22 low Conversely, under NAF E air masses the marine sulphate of regional origin 23 shows the highest concentrations, contrarily to the secondary sulphate factor 24 (Fig. S2). 25

- 2 5 Conclusions
- 3

1

With the aim of assessing and evaluating the vertical and horizontal spatial 4 5 variability of PM₁₀ concentrations in a Southern European urban environment, 221 PM₁₀ samples (12 hours resolution) were simultaneously collected at four 6 7 monitoring sites strategically located within the city of Barcelona during one month (SAPUSS campaign, 20 September to 20 October 2010). A decreasing 8 PM₁₀ concentration gradient from road traffic hot spots to the background areas 9 10 was recorded. Overall, both the proximity to traffic sources and the different types of air mass scenarios lead to a wide variability in concentrations and 11 chemical composition of PM₁₀ across the vertical and horizontal scale in 12 Barcelona during SAPUSS. When a PMF factor analysis was run on the 221 13 filters sampled collected, the optimal chosen solution contained eight factors: 14 15 (1) vehicle exhaust and wear, (2) road dust, (3) mineral, (4) aged marine, (5) heavy oil, (6) industrial, (7) sulphate and (8) nitrate. Overall, primary traffic 16 emissions (exhaust and wear and road dust) accounted for 18-39% of PM₁₀ 17 mass, primary inorganic aerosols (mineral dust and aged marine) 27-39%, 18 industry (heavy oil and industrial) 5-7% and secondary aerosols (sulphate and 19 nitrate) 28-36%. The main factors influencing the different sources 20 concentration at each site were: air mass origin, proximity to the emission 21 source and meteorological parameters, such as wind speed and direction 22 23 (influencing the sea breeze development for both dispersion and transport of specific pollutants) and temperature (causing the volatilization of nitrate under 24

high temperatures). Special emphasis was put in trying to further apportion the 1 dust aerosol sources. Overall, three sources of dust were identified in the urban 2 area of Barcelona: road dust (3.8-1.6 µg m⁻³, average 35%), Saharan dust (2.1 3 μ g m-3, average 28%) and mineral dust of regional origin (2.7-2.9 μ g m⁻³, 4 average 38%). Regarding the aged marine aerosol factor, it was found to be 5 internally mixed with sulphate of regional origin, as the calculated fresh sea salt 6 (1.8 µgm⁻³, 45% of the aerosol marine load) was aged by the mixing with 7 anthropogenic marine sulphate of regional origin (2.2 µgm⁻³, 55% of the aerosol 8 marine load). As expected, it was found that non vehicle exhausts, vehicle 9 10 exhausts, and local industries located in the city centre were contributing to the PM₁₀ ground concentrations levels. However, surprisingly the 11 PM_{10} concentrations of secondary aerosols were found more homogeneous than 12 13 expected. On the whole, our results show that although a higher homogeneity than expected was found in the horizontal and vertical variability of pollution 14 levels in the Barcelona urban atmosphere, primary emission factors related to 15 vehicle exhaust emissions and road dust resuspension decrease with the 16 distance to traffic hot spots. Road traffic emissions comprise not only tailpipe 17 18 exhaust emissions but also non-exhaust emissions derived from the vehicleinduced resuspension of dust deposited on the road, and from the direct 19 emissions from vehicle wear (brakes, tyres, discs etc.). This study confirms that 20 - for the coarse PM₁₀ fraction - road traffic is still a major source of ground level 21 PM₁₀ aerosol mass. Furthermore, this study shows that local industries-small 22 workplaces are also a source of PM₁₀ aerosol mass within urban ground levels. 23

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10 LIST OF TABLES AND FIGURES

Table 1: Mean concentrations (µgm⁻³) of PMF factors at the Road Site (RS),

- 13 Urban Background (UB), Torre Mapfre (TM) and Torre Collserola (TC).

PMF Factors (µgm ⁻³)	RS	UB	ТМ	тс
Exhaust & wear	8.7 (27%)	5.0 (18%)	2.9 (11%)	1.9 (10%)
Road dust	3.8 (12%)	3.3 (12%)	2.3 (9%)	1.6 (8%)
Mineral	4.6 (13%)	5.1 (18%)	4.8 (19%)	4.9 (26%)
Aged marine	4.6 (14%)	3.6 (13%)	5.2 (20%)	2.6 (13%)
Heavy oil	0.5 (2%)	0.6 (2%)	0.6 (2%)	0.4 (2%)
Industrial	1.2 (4%)	1.4 (5%)	0.7 (3%)	0.9 (5%)
Sulphate	3.5 (11%)	4.2 (15%)	3.8 (15%)	3.3 (17%)
Nitrate	5.7 (17%)	4.9 (17%)	5.5 (21%)	3.6 (19%)
	32.6 (100%)	28.1 (100%)	25.8 (100%)	19.2 (100%)

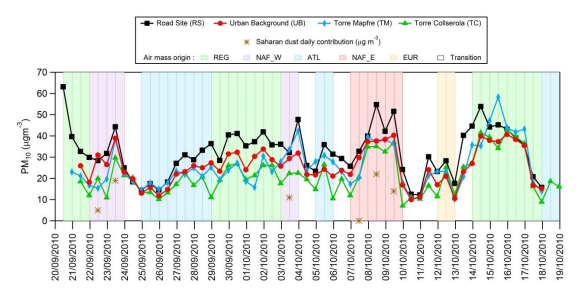
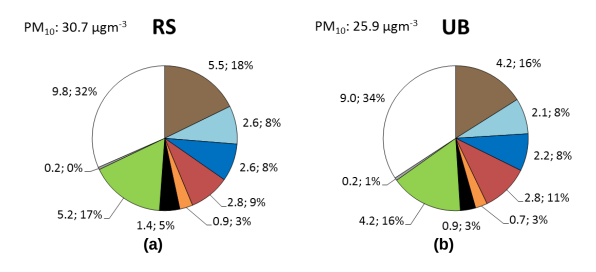


Figure 1: PM₁₀ variation at the 4 sites (RS, UB, TM, TC) during the SAPUSS
campaign under different air mass origin (Regional (REG), North African West
(NAF_W), Atlantic (ATL), North African East (NAF_E), European (EUR)).
Saharan dust daily contribution to PM₁₀ is indicated.



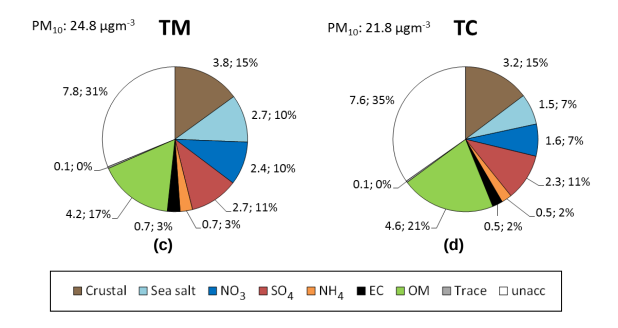
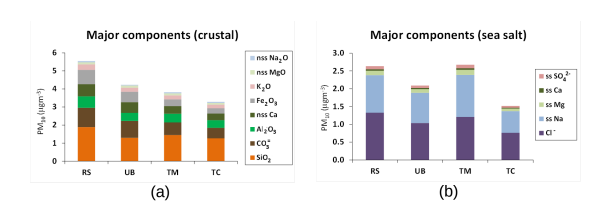
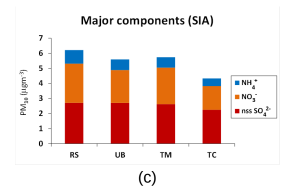
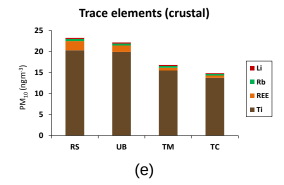


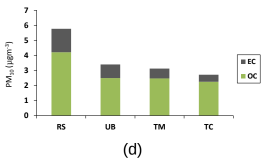
Figure 2: Mean composition of PM₁₀ concentration in µg m⁻³ measured during
the SAPUSS campaign at: a) Road Site (RS), b) Urban Background (UB), c)
Torre Mapfre (TM) and d) Torre Collserola (TC). Data are given in µgm⁻³ and %.
On the top right of each graph average gravimetric PM₁₀ concentration are
represented.



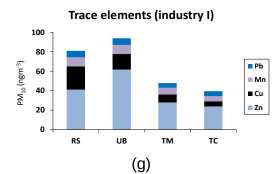




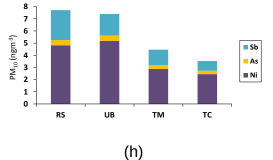


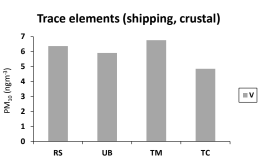


Trace elements (road traffic) 50 40 PM₁₀ (ngm⁻³) Ba 30 Sb 20 🛛 Sn 🔳 Cu 10 0 RS UB ТΜ тс (f)



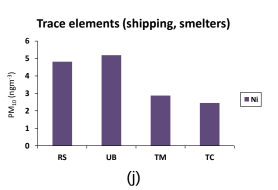
Trace elements (industry II)





(i)

RS

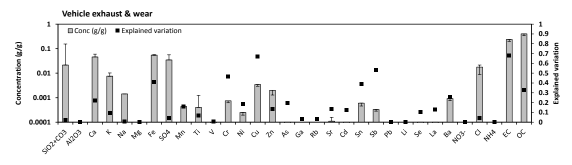


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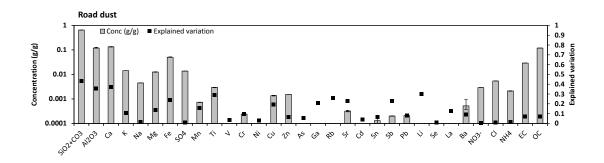
Figure 3: Average PM₁₀ concentration of main and trace elements for different 2 3 emission sources at each site (RS: Road Site, UB: Urban Background, TM:

4 Torre Mapfre, TC: Torre Collserola). REE denote Rare Earths elements.

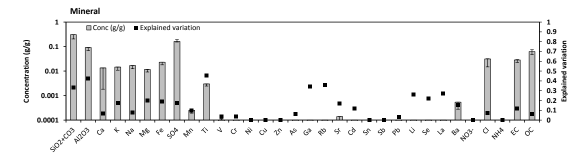
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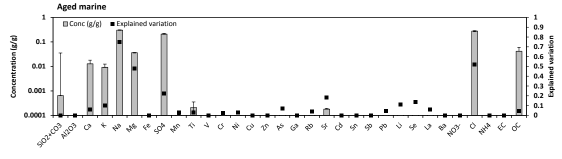
(a)



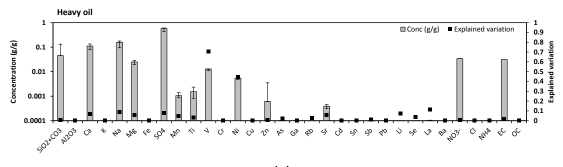
(b)



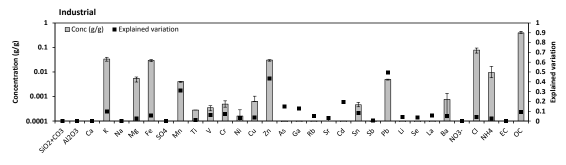
(c)



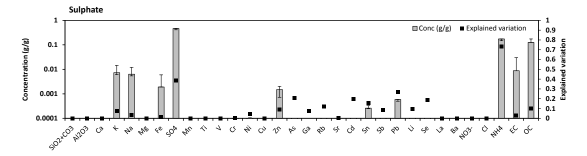
(d)











(g)

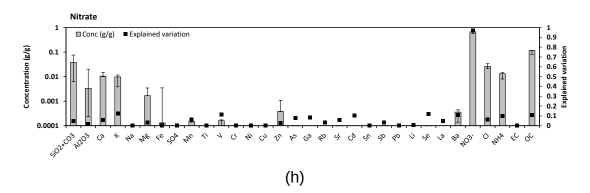


Figure 4: PMF sources profiles for PM₁₀ during the SAPUSS campaign: a)
vehicle exhaust and wear, b) road dust, c) mineral, d) aged marine, e) heavy oil,
f) industrial, g) sulphate, h) nitrate. Uncertainties were obtained by
bootstrapping.

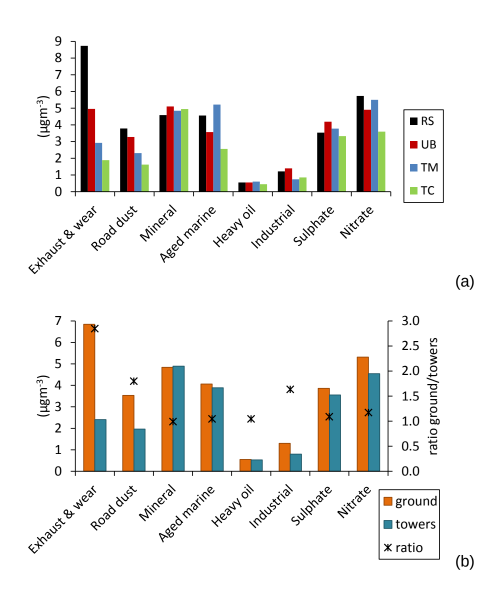
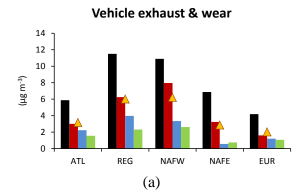
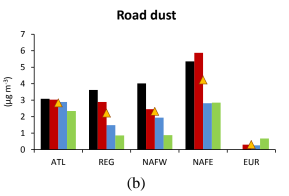
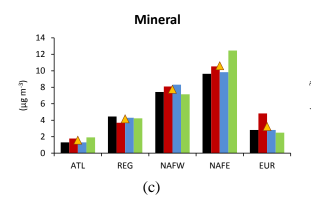


Figure 5: Contribution to PM₁₀ concentration levels of each of the eight factors:
a) at each of the 4 sites (RS, UB, TM, TC) and b) at ground (RS and UB) and
tower levels (TM and TC) and the concentration ratio between ground and
tower sites during the SAPUSS campaign.

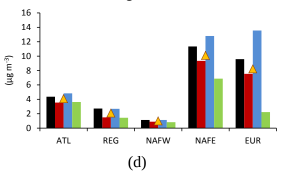
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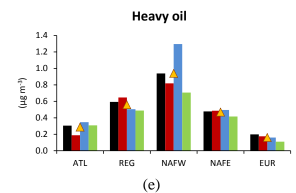




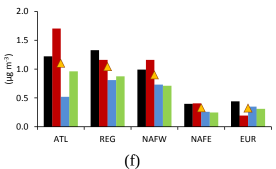


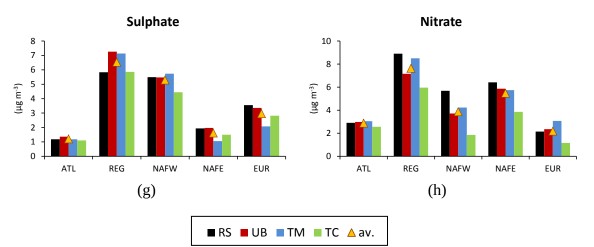
Aged marine





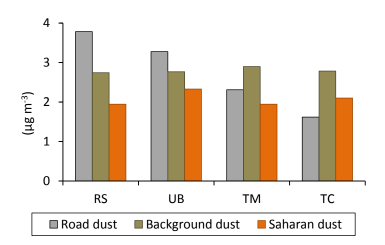
Industrial





- 1 Figure 6: Average PM₁₀ contributions from the eight PMF factors at each of the
- 2 sites (RS, UB, TM and TC) under different atmospheric scenarios (Atlantic,

- 1 ATL; Regional, REG; North African West, NAFW; North African East, NAFE and
- 2 European, EUR) during the SAPUSS campaign.
- 3



- 4 Figure 7: Average sources contributing to the mineral dust load during
- 5 SAPUSS at the four monitoring sites RS, UB, TM and TC. Mineral background
- 6 dust contributes on average 2.8 μ g m⁻³, road dust 2.7 μ g m⁻³ and Saharan dust

7 2.1 µg m⁻³.

8

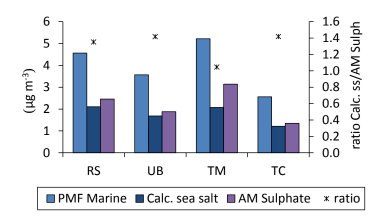
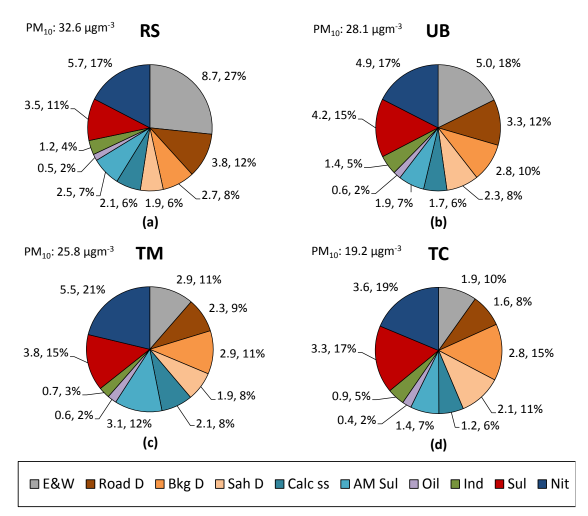


Figure 8: Average sources contributing to the sea salt factor during SAPUSS at
the four monitoring sites RS, UB, TM and TC. Calculated sea salt contributes on
average 1.8 μg m⁻³ and marine sulphate from regional pollution 2.2 μg m⁻³. (AM:
anthropogenic marine).

13

14



1 **Figure 9:** Sources contributing to the PM₁₀ load extracted with the PMF tool and subcomponents at each monitoring: a) RS, b) UB, c) TM and d) TC. 2 Exhaust&Wear (E&W), Road dust (Road D), Heavy oil (Oil), Industrial (Ind), 3 Sulphate (Sul) and Nitrate (Nit) are direct PMF factors. The mineral factor was 4 broken into Background dust (Bkg D) and Saharan dust (Sah D) and the aged 5 marine factor into Calculated sea salt (Calc ss) and Anthropogenic marine 6 sulfate of regional origin (AM Sul). Data are given in μ gm⁻³ and %. The average 7 PMF PM₁₀ concentrations are represented at the top left of each graph for each 8 9 site.

10

11