Impact of aerosol particle sources on optical properties in urban, regional and remote areas in the north-western Mediterranean

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Abstract. Further research is needed to reduce the existing uncertainties on the effect that specific aerosol particle sources have on light extinction and consequently on climate. This study presents a new approach that aims to quantify the mass scattering and absorption efficiencies (MSEs and MAEs) of different aerosol sources at urban (Barcelona – BCN), regional (Montseny – MSY) and remote (Montsec – MSA) background sites in the north-western (NW) Mediterranean. An analysis of source apportionment to the measured multi-wavelength light scattering ($\sigma_{sp}$) and absorption ($\sigma_{ap}$) coefficients was performed by means of a multi-linear regression (MLR) model for the periods 2009–2014, 2010–2014 and 2011–2014 at BCN, MSY and MSA respectively. The source contributions to PM$_{10}$ mass concentration, identified by means of the positive matrix factorization (PMF) model, were used as dependent variables in the MLR model. With this approach we addressed both the effect that aerosol sources have on air quality and their potential effect on light extinction through the determination of their MSEs and MAEs. An advantage of the presented approach is that the calculated MSEs and MAEs take into account the internal mixing of atmospheric particles.

Seven aerosol sources were identified at MSA and MSY, and eight sources at BCN. Mineral, aged marine, secondary sulfate, secondary nitrate and V-Ni bearing sources were common at the three sites. Traffic, industrial/metallurgy and road dust resuspension sources were isolated at BCN, whereas mixed industrial/traffic and aged organics sources were identified at MSY and MSA. The highest MSEs were observed for secondary sulfate (4.5 and 10.7 m$^2$ g$^{-1}$, at MSY and MSA), secondary nitrate (8.8 and 7.8 m$^2$ g$^{-1}$) and V-Ni bearing source (8 and 3.5 m$^2$ g$^{-1}$). These sources dominated the scattering throughout the year with marked seasonal trends. The V-Ni bearing source, originating mainly from shipping in the area under study, simultaneously contributed to both $\sigma_{sp}$ and $\sigma_{ap}$, being the second most efficient light-absorbing source in BCN (MAE = 0.9 m$^2$ g$^{-1}$). The traffic source at BCN and the industrial/traffic at MSY exhibited the highest MAEs (1.7 and 0.9 m$^2$ g$^{-1}$). These sources were major contributors to $\sigma_{sp}$ at BCN and MSY; however at MSA, secondary nitrate exerted the highest influence on $\sigma_{ap}$ (MAE = 0.4 m$^2$ g$^{-1}$). The sources which were predominantly composed of fine and relatively dark particles, such as industrial/traffic, aged organics and V-Ni, were simultaneously characterized by low single scattering albedo (SSA) and a high scattering Ångström exponent (SAE). Conversely, mineral and aged marine showed the lowest SSA and the highest SAE, being scattering the dominant process in the light extinction. The good agreement found between modelled and measured particle optical properties allowed the reconstruction of $\sigma_{sp}$ and $\sigma_{ap}$ long-term series over the period 2004–2014 at MSY. Significant decreasing trends were found for the modelled $\sigma_{sp}$ and $\sigma_{ap}$ ($-4.6$ and $-4.1$ % yr$^{-1}$).

1 Introduction

Atmospheric aerosol particles affect the Earth’s climate through the direct scattering and absorption of solar radiation but also through indirect processes acting as cloud condensation nuclei (IPCC, 2007). Precise measurements of aerosol properties are required to reduce the current uncertainties on radiative forcing (IPCC, 2007, 2013), and further research that aims to study the existing relationship between...
aerosol optical and chemical properties is needed to better understand the air quality–climate link. However, a thorough quantification of the direct and indirect aerosol effects on the Earth’s radiative budget is difficult to achieve (Zieger et al., 2012). The high spatial and temporal variability of atmospheric aerosols along with the large differences in particle composition and size (Andrews et al., 2011; Bond et al., 2013; Haywood et al., 1999), results in a changing radiative forcing from local to global scales (Collaud Coen et al., 2013). On the global scale, atmospheric aerosols are estimated to cool the Earth system (Chen et al., 2011; IPCC, 2013). Most aerosol components (mainly sulfate, nitrate, organics and mineral matter) scatter the sunlight causing a net cooling at the top of the atmosphere (TOA); conversely other particles, such as black carbon (BC), absorb solar radiation in the whole visible spectrum, thus leading to a net warming at TOA (Jacobson, 2001a; Ramanathan and Carmichael, 2008). Assessing the role of aerosols on climate forcing often requires reducing their physicochemical properties to a set of parameters that describe their optical properties (Hand and Malm, 2007). The mass scattering and absorption efficiencies (MSEs and MAEs) are key intensive optical parameters that relate the mass concentration of specific chemical species to the particle light scattering ($\sigma_{ap}$) and absorption ($\sigma_{sp}$) coefficients. These intensive optical parameters depend on intrinsic aerosol properties, such as particle effective radius, particle mass density or refractive index, and they are very useful to better parameterize the aerosols direct radiative effect in atmospheric climate models (Seinfeld and Pandis, 1998). In fact, Obiso et al. (2017) has recently assessed the T-matrix optical code to simulate MSEs of different aerosol sources, considering the MSEs reported in the present study as reference parameters representative of the NW Mediterranean area.

Several studies have been published on the absorption efficiency of black carbon (BC) calculated as the ratio between $\sigma_{ap}$ and elemental carbon (EC) concentrations. Given that BC is the most important light-absorbing particle in the atmosphere, its MAE has been extensively studied in the literature (i.e. Bond et al., 2013; Pandolfi et al., 2014a; Reche et al., 2011, among others). In some cases, the MAE of BC has been observed to change depending on the degree of the internal mixing of BC with non-absorbing material, such as sulfate and organic compounds (Jacobson, 2001b; Moffet and Prather, 2009; Ramana et al., 2010; Zanatta et al., 2016).

Recently, the potential for organic carbon as an absorber of UV and visible light through their brown carbon (BrC) content, has been also reported in the literature (i.e. Lu et al., 2015; Updyke et al., 2012).

The MSEs of different chemical aerosol components have been extensively reported for many locations (Vrekoussis et al., 2005; Titos et al., 2012; Cheng et al., 2015 and references therein). An example is the study performed by the IMPROVE (Interagency Monitoring of Protected Visual Environments) programme, which has been considered as a reference for reporting mass extinction efficiencies depending on particle composition (Hand and Malm, 2006 and 2007). Global MSEs for dry ammonium sulfate ([NH$_4$]$_2$SO$_4$), ammonium nitrate [NH$_4$NO$_3$], organic matter (OM), soil dust and sea salt were obtained by means of a multilinear regression (MLR) model. In the IMPROVE model, $\sigma_{sp}$ measurements (from 1990 to 2007) were used as independent variable whereas the aforementioned externally mixed chemical species were used as dependent variables. In addition, the IMPROVE study demonstrated that the reconstruction of $\sigma_{sp}$ can be inversely computed by means of the calculated MSE and the mass concentration of chemical species. Revised versions of the IMPROVE algorithm have been published that aim to reduce the bias on the predicted values, which accounted for a 25 % overestimate of the measured $\sigma_{sp}$ coefficient (Ryan et al., 2005; Pitchford et al., 2007). However, none of the published studies dealing with the estimation of MSEs have considered the internal mixing state of atmospheric aerosols, given that each chemical specie was treated separately from the other.

In the present study a different approach of the MLR method is presented, where the aerosol source contributions obtained by means of the PMF (positive matrix factorization) model, instead of the single chemical species, were used as dependent variables in the MLR model. An important characteristic of the PMF factors is that these take into account the internal mixing of atmospheric particles. In fact, as evidenced by the PMF sources chemical profiles, these are constituted by some main tracers (which define the source) but are also enriched in other chemical compounds. Receptor models such as PMF are powerful and widely used techniques to design air quality mitigation strategies (i.e: Belis et al., 2013; Viana et al., 2008), thanks to the capability of these models to identify key pollutant emission sources and calculate their contributions to the measured PM mass concentration. Thus, the MLR model applied using the PMF source contributions and the measured $\sigma_{sp}$ and $\sigma_{ap}$ allows quantifying the potential of different aerosol particle sources to scatter or absorb visible light and therefore directly linking the air quality and climate effects of airborne PM.

With this approach we estimated the MSEs and MAEs of aerosol particle sources identified at urban, regional and remote environments in the NE of Spain. Furthermore, the computed MSEs and MAEs were used to reconstruct the particle $\sigma_{sp}$ and $\sigma_{ap}$ over an 11-year period at the MSY regional site, thus allowing trend analyses. Trend analyses of particle optical properties are extremely relevant for the detection of changes in atmospheric composition depending on changes in natural or anthropogenic emissions, atmospheric processes and sinks (Collaud Coen et al., 2013). Several studies have shown that the air quality abatement strategies adopted in the recent years have resulted in a decrease in anthropogenic pollutants in Europe (EEA, 2013; Barmpadimos et al., 2012; Querol et al., 2014; Pandolfi et al., 2016). However, the control of pollutant emissions is currently conflicted, involving
a trade-off between the impacts on environmental health and the Earth’s climate, and therefore current mitigation strategies could increase climate warming while improving air quality (Shindell et al., 2012). A relevant outcome of this new approach is the chance to study the effects that air quality mitigation strategies are having on light extinction in the area under study.

2 Methodology

2.1 Sampling sites and meteorology

The western Mediterranean Basin (WMB) is characterized by warm summers and temperate winters with irregular precipitation rates throughout the year. In winter the location of the Azores high-pressure system favours the entry of Atlantic advections that clear the atmosphere of pollutants. In summer, atmospheric dynamics coupled to local orography result in local/regional circulations with a consequent accumulation of pollutants (Millán et al., 1997). Recirculation and aging of pollutants is favoured by weak gradient atmospheric conditions, scarce precipitation and continuous exposure to solar radiation driving photochemical reactions (Rodríguez et al., 2002; Pérez et al., 2004). Additionally, large mineral dust contributions from Saharan dust events may exceed air quality standards (Escudero et al., 2007; Querol et al., 2009). The conjunction of all these processes surrounding the WMB lead to a radiative forcing among the highest in the word (Jacobson, 2001a).

PM chemical and optical measurements were performed at three sampling sites located in NE Spain (Fig. 1). The large coastal Barcelona urban area (BCN; 41°23′N, 02°6′E, 80 m a.s.l.) is one of the most populated areas in the NW Mediterranean, resulting in a very high road traffic density. Additionally, the metropolitan area is surrounded by a broad industrial area and is home to one of the major harbours in the Mediterranean Basin, with a large number of cruise ships (Pey et al., 2013). The conjunction of these emission sources contribute greatly to the air quality degradation in the area (Querol et al., 2001; Pey et al., 2008; Amato et al., 2009; Reche et al., 2011; Dall’Osto et al., 2013).

The Montseny regional background station (MSY; 41°19′N, 02°21′E, 720 m a.s.l.) is located in the Montseny natural park in a densely forested area, 50 km to the N–NE of the Barcelona urban area and 25 km from the Mediterranean coast. Despite the site being far enough from the industrialized and populated Barcelona metropolitan region, it can be affected by anthropogenic emissions transported to regional inland areas (Pérez et al., 2008).

The Montsec continental background site is a remote high altitude location (MSA; 42°3′N, 0°44′E, 1570 m a.s.l.) on the southern side of the Pre-Pyrenees at the Montsec d’Ares mountain range, located 140 km to the NW of Barcelona and 140 km to the WNW of MSY. Despite the high-altitude location and the frequent free-troposphere conditions during the cold season, the station can be slightly influenced by anthropogenic emissions during the warmer period, when it is positioned within the planetary boundary layer (PBL) (Ripoll et al., 2014).

The three sites are members of the Catalanian air quality monitoring network. Additionally, MSY and MSA are part of the ACTRIS (Aerosol, Clouds and Trace gases Research InfraStructure) and GAW (Global Atmosphere Watch) networks. Aerosol optical properties at the sites are measured following standard network protocols (WMO/GAW, 2016). Further information characterizing physical, chemical and optical properties of atmospheric aerosols detailing the prevailing atmospheric dynamics at the three stations can be found in Querol et al. (2001), Pey et al. (2009, 2010), Reche et al. (2011), Pandolfi et al. (2011, 2014a), Cusack et al. (2012) and Ealo et al. (2016).

2.2 Measurements and instrumentation

Aerosol light scattering coefficients were measured every 5 min at three wavelengths (450, 525 and 635 nm) with a LED-based integrating nephelometer (model Aurora 3000, ECOTECH Pty, Ltd, Knoxfield, Australia). \( \sigma_{sp} \) measurements were collected at MSY for the period 2010–2014 (Table S1 in the Supplement) using a PM\(_{10}\) cut-off inlet. Measurements at MSA were carried out using a PM\(_{2.5}\) cut-off inlet from 2011 to March 2014 and then replaced with a PM\(_{10}\) cut-off inlet. \( \sigma_{sp} \) measurements at BCN are not available. Calibration of the two nephelometers was performed three times per year using \( \text{CO}_2 \) as span gas, while zero adjusts were performed once per day using internally filtered particle-free air. The relative humidity (RH) threshold was set by using a processor-controlled automatic heater inside the Aurora 3000 nephelometer to ensure a sampling RH of less than 40\% (GAW report 226). \( \sigma_{sp} \) coefficients were corrected for non-ideal illumination of the light source and for truncation of the sensing volumes following the procedure described in Müller et al. (2011a).

Aerosol light absorption coefficient at 637 nm (Müller et al., 2011b) was measured at 1 min resolution with a Multi-Angle Absorption Photometer (MAAP, model 5012, Thermo) operated in the heated sampling mode and connected to a PM\(_{10}\) cut-off inlet. \( \sigma_{ap} \) measurements were collected at BCN, MSY and MSA for the periods 2009–2014, 2010–2014 and 2011–2014 respectively.

Gravimetric PM\(_{10}\) mass concentrations were determined by standard gravimetric procedures, according to UNE-EN 12341, 1999 protocol (Alastuey et al., 2011). Samples were collected every 3 to 4 days on 150 mm quartz micro-fibre filters ( Pallflex 2500 QAT-UP and Whatman QMH) using high-volume samplers (DIGITEL DH80 and/or MCV CAV-A/MSb at 30 m\(^3\)h\(^{-1}\)) for the periods 2004–2014 at BCN and MSY, and for the period 2010–2014 at MSA.
Chemical offline filter analyses were carried out at the three sites following the procedure proposed by Querol et al. (2001). A quarter of the filter was acid digested (HNO₃ : HF : HClO₄). The resulting solution was analysed by inductively coupled plasma atomic emission spectroscopy (ICP-AES; IRIS Advantage TJA Solutions THERMO) for the determination of major elements (Al, Ca, Fe, K, Na, Mg, Si, Ti, P) and by inductively coupled plasma mass spectrometry (ICP-MS; X Series II, THERMO) for the trace elements (Li, Ti, V, Cr, Mn, Co, Ni, Cu, Zn, As, Se, Rb, Sr, Cd, Sn, Sb, Ba, rare earths, Pb, Bi, Th, U). In order to examine the accuracy of the acid digestion, a few milligrams of the reference material NIST 1633b were added to a quarter of the blank filter. Another quarter of each filter was water extracted to determine soluble anions. The nitrate, sulfate and chloride concentrations were resolved by ion high-performance liquid chromatography (HPLC) using a WATERS ICpakTM anion column with a WATERS 432 conductivity detector and the ammonium concentrations with an ion selective electrode (MODEL 710 A+, THERMO Orion). Organic carbon (OC) and EC concentrations were determined by a thermal-optical carbon analyser (SUNSET) following the EUSAAR2 thermal protocol (Cavalli et al., 2010). Blank filters were analysed together with the samples, and concentrations were subtracted from those found in the samples in order to calculate the ambient concentrations.

### 2.3 Positive matrix factorization model (PMF)

The positive matrix factorization (PMF) model (PMFv5.0, EPA) was individually applied to the daily chemical speciated data collected at BCN, MSY and MSA for source identification and apportionment to PM₁₀. Source contributions obtained for BCN and MSY can be found in Pandolfi et al. (2016), whereas sources identified at MSA are presented in this study. Detailed information describing the PMF model can be found in the literature (Paatero and Tapper, 1994; Paatero, 1997; Paatero and Hopke, 2003; Paatero et al., 2005).

The PMF model is a factor analytical tool based on the weighted least-squares method, which reduces the dimension of the input matrix (i.e. the daily chemical speciated data) to a limited number of factors (or sources). Calculation of individual uncertainties and detection limits were based on the approach by Escrig et al. (2009) and Amato et al. (2009), wherein both the analytical uncertainties and the standard deviations of species concentrations in the blank filters were considered in the uncertainty calculations. This procedure provides a criterion with which to separate the species which retain a significant signal from the ones dominated by noise, based on the signal-to-noise \( S/N \) ratio defined by Paatero and Hopke (2003). Species with \( S/N \) greater than 1 may often indicate a good signal, though this depends on how uncertainties are determined. In order to avoid any bias in the PMF results, the data matrix was uncensored; i.e. negative, zero and below detection limit values were included in the analyses.

The PMF was run in robust mode (Paatero, 1997) and rotational ambiguity was handled by means of the FPEAK parameter (Paatero et al., 2005). The final number of sources was selected based on several criteria: (i) investigating the variation of the objective function \( Q \) (defined as the ratio be-
tween residuals and errors in each data value) depending on the number of sources (i.e. Paatero et al., 2002), (ii) studying the physical meaningfulness of factor profiles and contributions and (iii) analysing the scaled residuals and the $G$ space plots.

### 2.4 Multilinear regression model (MLR)

Previous studies based on the IMPROVE algorithm have applied the multilinear regression (MLR) method to estimate the mass scattering and extinction efficiencies (MSEs and MAEs) of chemical species (White, 1986; Vasconcelos et al., 2001; Hand and Malm, 2007). This kind of regression model between chemical species mass concentration and aerosol particle scattering or extinction coefficients assumes an externally mixed aerosol. However, the apportionment of scattering by more than one species to the total scattering depends on the assumption of the internal or external mixing state of atmospheric aerosols, as already noted previous studies (White, 1986). The assumption of internal mixing among chemical species that form a single variable in the regression equation will reduce the possible collinearity among the dependent variables of the MLR model, making the regression coefficients less sensitive to data uncertainties at the same time (Hand and Malm, 2007). As shown in the matrix correlation in Fig. S1, a very low correlation was observed between pairs of aerosol sources identified with the PMF model at the three sites considered here.

In this study, we used the PM$_{10}$ source contributions ($\mu$g m$^{-3}$) as dependent variables in the MLR and the measured $\sigma_{sp}$ and $\sigma_{ap}$ coefficients (Mm$^{-1}$) as independent ones. Thus, the resulting regression coefficients of the model represent the MSEs and MAEs (m$^2$ g$^{-1}$) of mixed aerosol modes, given that the sources from PMF take into account the possible internal mixing among chemical species. Moreover, the MLR method assumes that all the species contributing to $\sigma_{sp}$ and $\sigma_{ap}$ are included in the equation. Thus, a better model performance is achieved here given that we used the full PM$_{10}$ chemical speciation in the PMF model for source identification and apportionment. Following Eqs. (1) and (2) (as example for MSY), the partial $\sigma_{sp}$ and $\sigma_{ap}$ contribution of each source can be computed as the product between the PM$_{10}$ source contributions and the corresponding MSEs/MAEs. Then, total aerosol light $\sigma_{sp}$ and $\sigma_{ap}$ can be modelled as the sum of the scattering or absorption source contributions.

$$
\sigma_{sp, PM_10} = \left( MSE_{\text{Secondary sulfate}} \cdot \text{[Secondary sulfate]} \right)
+ \left( MSE_{\text{Secondary nitrate}} \cdot \text{[Secondary nitrate]} \right)
+ \left( MSE_{\text{V-Ni}} \cdot \text{[V-Ni]} \right)
+ \left( MSE_{\text{Aged organics}} \cdot \text{[Aged organics]} \right)
+ \left( MSE_{\text{Mineral}} \cdot \text{[Mineral]} \right)
+ \left( MSE_{\text{Aged marine}} \cdot \text{[Aged marine]} \right)
$$

It should be considered that changes in the sampling conditions (i.e. RH or size cut-off) or differences in the chemical analysis methods used on sampled filters can affect the intensive particle optical properties (Delene and Ogren, 2002) and consequently the comparison among the computed MSEs and MAEs. In fact, the resulting efficiencies can be biased by the cut-off inlet, given that absorbing aerosols tend to be predominately in the submicron fraction (Andrews et al., 2011). In this study both $\sigma_{sp}$ and $\sigma_{ap}$ were collected using a PM$_{10}$ cut-off inlet, thus guaranteeing uniformity among the performed optical measurements. An exception occurs at MSA, where a PM$_{2.5}$ cut-off inlet was used until March 2014 and then replaced by a PM$_{10}$ inlet. Therefore, a slight over-estimation of the MSEs obtained for aged marine and mineral sources at MSA might be expected when sampling was performed through the PM$_{2.5}$ inlet, given that particles contained in these sources are mainly present in the coarse fraction and significantly contribute to PM$_{10}$ mass concentration (Ripoll et al., 2015a). However, an estimation of the influence of the inlet change on the resulting MSEs and MAEs at MSA is difficult to achieve, given the relatively short $\sigma_{sp}$ and $\sigma_{ap}$ time series available, thus preventing performing two different MLR analyses for the two fractions. Moreover, scattering RH was controlled below 40% at MSY and MSA in order to minimize the hygroscopic growth of the particles and then prevent a significant enhancement in the scattering efficiencies. An overestimation of the scattering or absorption efficiencies can also be due to the fact that the MLR method tends to give more weight to those variables that are more accurately measured (such as sulfate), and conversely, underestimates the regression coefficients for species with larger uncertainty (i.e. organic matter) (White and Macias, 1987). In the present study, a comparison between modelled and measured coefficients was performed using quantitative statistics. With this aim, the root mean square error (RMSE) and fractional bias (FB) were computed for modelling evaluation. FB is described in Eq. (3) (Ryan et al., 2005), where $\sigma_{sp, \text{sim}}$ is the modelled scattering coefficient and $\sigma_{sp}$ is the measured value for each daily data point.

$$
FB = \frac{\sigma_{sp, \text{sim}} - \sigma_{sp}}{\sigma_{sp}}
$$
A total of 303, 379 and 503 daily data points were used in the MLR analysis for source apportionment analysis of absorption at MSA, MSY and BCN respectively, whereas 222 and 307 daily data points were considered for MSE calculation at MSA and MSY.

2.5 Statistical tests for trends study

The Theil-Sen slope estimate (TS) (Theil, 1950; Sen, 1968) is a non-parametric test which was investigated for the monthly averages of light scattering and absorption in order to test for the occurrence of a non-null slope in the data series during the period 2004–2014 at MSY. The total and annual reduction of these optical parameters was investigated using bootstrap resampling for the monthly deseasonalized time series, reducing the possible influence of outliers on trend estimates and obtaining robust slope \( p \) values.

A multi-exponential fit aiming to study temporal trends in the multi-exponential form (Shatalov et al., 2015) was used to represent the decomposed modelled monthly temporal series in the main component, seasonal component and residual component. Additionally, this technique allowed us to estimate the non-linearity (NL) parameter for the trends. An NL of 10 % was used as threshold to define a linear trend (NL < 10 %).

3 Results

3.1 Source profiles and contributions to PM\(_{10}\)

Seven aerosol particle sources were identified at MSA in the PM\(_{10}\) fraction by performing a PMF analysis for the period 2010–2014. The chemical profiles and source contributions to the measured PM\(_{10}\) mass are shown in Fig. 2 and Table 1. These results will be studied together with the chemical profiles (Fig. S2) and source contributions (Table 1) previously quantified by Pandolfi et al. (2016) for BCN and MSY for the period 2004–2014. The highest PM\(_{10}\) average concentration was found at the BCN urban station, followed by the regional (MSY) and remote (MSA) background sites (34.0 ± 17.1, 16.7 ± 9.3 and 9.6 ± 8.2 \( \mu g \) m\(^{-3}\) respectively), consistent with the progressive distance between the three stations and important emission sources. On average, the most abundant sources contributing to PM\(_{10}\) mass concentration at MSA were aged organics, followed, in this order, by mineral, industrial/traffic, aged marine, secondary sulfate, V-Ni bearing and secondary nitrate. Aged organics sources was mainly traced by OC and EC with maxima in summer, pointing to a large contribution from biogenic emission sources, and accounted for 2.8 ± 2.0 \( \mu g \) m\(^{-3}\) (29 %) of the PM\(_{10}\) mass concentration. The internal mixing with EC suggests a contribution from combustion sources to this source. However, the aged organics source at MSA can be considered to be dominated by secondary organic aerosols (SOAs) arising from biogenic volatile organic compounds (VOCs) due to the predominance of OC in the chemical profile. Furthermore, the higher summer VOC oxidative potential occurring in the Mediterranean should be considered, which enhances SOA concentrations due to both higher insolation and tropospheric ozone concentration (Fuzzi et al., 2006). This assertion is in agreement with previous studies deployed at MSA where SOA was found to be the foremost constituent of PM\(_1\) organic aerosols (OAs), especially in summer (90 %) (Ripoll et al., 2015a). The mineral source, traced by typical crustal elements such as Al, Ca, Mg, Fe, Ti, Rb and Sr, was related to both Saharan dust events and regional/local mineral contribution and accounted for an average PM\(_{10}\) contribution of 2.3 ± 5.2 \( \mu g \) m\(^{-3}\) (24 %). The industrial/traffic source, primarily traced by Pb, Zn, As, Sb, Cu and Ni, contributed 1.1 ± 1.0 \( \mu g \) m\(^{-3}\) (11 %). Aged marine sources, mainly traced by Na and Cl, and in a minor proportion by Mg, SO\(_4\)^{2−} and NO\(_3\)^{−}, contributed 1.1 ± 1.3 \( \mu g \) m\(^{-3}\) (11 %). Secondary sulfate, mainly traced by SO\(_4\)^{2−} and NH\(_4\)^{+}, and secondary nitrate, traced by NO\(_3\)^{−} and NH\(_4\)^{+} but also enriched in EC, contributed 0.9 ± 1.0 \( \mu g \) m\(^{-3}\) (9 %) and 0.7 ± 1.0 \( \mu g \) m\(^{-3}\) (8 %) respectively. The V-Ni bearing source, traced by V, Ni and SO\(_4\)^{2−}, represented the direct emissions from heavy oil combustion, mainly shipping in the study area, and contributed 0.8 ± 1.0 \( \mu g \) m\(^{-3}\) (8 %). In contrast to BCN and MSY, the V-Ni bearing source at MSA was not enriched in EC, possibly because of the high altitude of this station and its position far from the NW Mediterranean coastline and shipping emissions.

Common sources identified at the three stations were mineral, aged marine, secondary nitrate, secondary sulfate and V-Ni. The sources identified in BCN showed similar contributions, ranging from 10 to 17 % of the total PM\(_{10}\) mass concentration, except for the industrial source (3 %), given that most of the secondary industrial aerosols are apportioned to other secondary sources presented in this study. At BCN, sources traced by pollutants from anthropogenic activities were mostly related to fresh emissions from the Barcelona metropolitan area (i.e. traffic and road dust re-suspension), from the surrounding industrial zone (industrial) and from vessel traffic (V-Ni). However at MSY and MSA, which are representative of regional and remote backgrounds, pollutants were transported together from the urban and industrial areas of Barcelona, thus resulting in an aged aerosol mixed with local pollutants. Larger relative contributions of mineral and aged organics sources were observed at the MSA high-altitude site due to a less direct exposure to anthropogenic emissions (Fig. 3c). In agreement with previous studies (Ripoll et al., 2015b; Ealo et al., 2016), a higher relative mineral contribution was found at MSA (23 %) compared to MSY (16 %) and BCN (14 %). However, a higher absolute mineral contribution mainly originating from local sources was observed at BCN (4.6 ± 5.3 \( \mu g \) m\(^{-3}\)). The aged organics source also presented a higher relative contribution at MSA (29 %) compared to MSY (23 %). However, this source was not identified at BCN, where the traffic source...
explained the majority of the measured OC. The aged marine source in Barcelona showed higher absolute and relative contributions (5.7 ± 5.2 µg m⁻³; 17 %) due to its proximity to the coast compared to MSY (1.8 ± 1.8 µg m⁻³; 11 %) and MSA (1.1 ± 1.3 µg m⁻³; 11 %). Higher relative contributions of secondary sulfate and secondary nitrate were found at MSY (24 and 8 %) compared to MSA (9 and 7 %), likely because of the longer distance from MSA to the Mediterranean coast. Moreover, the free-troposphere conditions typically occurring in MSA during the colder months prevented the direct transport of aerosol particles from anthropogenic sources to the station. The V-Ni bearing source showed similar absolute contributions at MSY (0.7 ± 0.7 µg m⁻³; 4 %) and MSA (0.8 ± 1.0 µg m⁻³; 8 %) despite the longer distance from MSA to the Mediterranean coast, pointing to a possible influence of long-range transport affecting the mountain-top site. It should be noted that the current increasing shipping emissions contribute greatly to air quality degradation in coastal areas (Viana et al., 2014), but also in regional and remote environments as consequence of atmospheric transport processes.

Overall, the impact of the identified aerosol sources at the different background sites depended on the distance to important emission sources and on the aging and transport of aerosol particles to regional and remote inland areas driven by orography and meteorology, thus mostly explaining the differences in the chemical profiles of the sources identified at the three sites.
3.2 Seasonal variation of source contributions to PM$_{10}$

Monthly average source contributions to PM$_{10}$ obtained at the three stations are shown in Fig. 3. MSY and MSA were characterized by a marked PM$_{10}$ seasonal variation with higher concentrations in summer (June, July and August) and lower in winter (December, January and February), in agreement with previous studies (Pérez et al., 2008; Ripoll et al., 2014). The summer increase is related to the higher frequency of Saharan dust events, the recirculation of air masses that prevent air renovation, the resuspension processes due to the dryness of soils, the low precipitation and the formation of secondary aerosols (Rodriguez et al., 2002). The lower winter concentrations can be explained by the high frequency of Atlantic advections leading to a higher dispersion of pollutants and to higher precipitation rates compared to summer. Moreover, the reduced contribution from the PBL in winter due to frequent thermal inversions also contributed to the relatively low PM$_{10}$ mass concentration observed at MSY, and especially at MSA (Pandolfi et al., 2014a). The PM$_{10}$ concentration peak observed in February and March at MSY is remarkable and might be attributed to the winter regional pollution episodes typical of the WMB (Pandolfi et al., 2014b). Such scenarios are characterized by anticyclonic conditions which favour the accumulation of pollutants close to the emission sources, and the subsequent transport of pollutants towards the station with the daily increase in the PBL. Pandolfi et al. (2014b) and Pey et al. (2010) reported high nitrate concentrations during these atmospheric conditions at MSY, in agreement with the increased contributions of secondary nitrate shown in Fig. 3b during this time of the year. The relatively low PM$_{10}$ concentration observed in August at BCN and MSY could be partially explained by reduced anthropogenic activities in the Barcelona metropolitan and industrial areas as a result of the holiday period in Spain. This result is supported by the minima of absolute contributions observed in August for industrial and traffic sources at BCN (0.6 ± 0.6 and 2.7 ± 1.6 µg m$^{-3}$ respectively) and for the industrial/traffic source at MSY (0.9 ± 0.7 µg m$^{-3}$). The higher precipitation rates observed in August compared to June–July (Perez et al., 2008) might also contribute to reducing PM$_{10}$ concentrations at MSY. Conversely at MSA,
the highest PM$_{10}$ concentration was observed in August, probably due to the frequent Saharan dust events affecting the mountain top site, in accordance with the highest absolute contribution found for the mineral source in August ($4.8 \pm 4.8 \mu g \text{ m}^{-3}$).

Higher relative contributions of aged marine (23 %), mineral (18 %), secondary sulfate (16 %) and V-Ni bearing (13 %) sources were observed on average in summer at BCN. By contrast, traffic (23 %), secondary nitrate (21 %) and industrial (4 %) sources maximized in winter (Fig. 3a). The seasonal variation of secondary sulfate and secondary nitrate can be attributed to a higher SOA contribution, the favoured formation of sulfate and the nitrate gas–aerosol partitioning, leading to the thermal instability of secondary nitrate during the warmer period, as was already observed in the area under study using offline filter sampling (Pey et al., 2009; Ripoll et al., 2015b) and online measurements (Ripoll et al., 2015a). In contrast to BCN, a higher relative contribution of secondary sources, some of them related to natural processes, was observed at MSY and MSA (3b and 3c). Increased contributions of secondary sulfate were observed in summer (29 and 8 % at MSY and MSA), whereas secondary nitrate maximized in winter (17 and 11 %). Aged organics showed the highest contribution in relative terms in winter (30 and 45 % at MSY and MSA); however the highest absolute contributions were observed in summer (4.8 ± 2.8 and 4.1 ± 1.9 µg m$^{-3}$). This result is in agreement with the higher SOA formation found at MSA (Ripoll et al., 2015a) and MSY (Minguillón et al., 2015) during the warm period. The mineral source (19 and 27 % at MSY and MSA) maximized in summer, although high contributions were also observed in spring. Similarly to BCN, aged marine (14 and 13 % for MSY and MSA) and V-Ni bearing (5 and 11 %) sources showed the highest contributions in summer, whereas the industrial/traffic source maximized in winter (11 and 17 %).

### 3.3 Mass scattering and absorption efficiencies of aerosol sources

Source-dependent mass scattering (at 450, 525 and 635 nm) and absorption (at 637 nm) efficiencies obtained at the different sites are shown in Table 2. The MSEs and MAEs for some of the sources reported in this study cannot be directly compared to MSEs and MAEs published in the literature for specific chemical species, given that the sources identified from PMF take into account the possible particle internal mixing. Similar MSEs were observed for secondary nitrate at MSY and MSA ($8.8 \pm 0.4$ and $7.8 \pm 0.8$ m$^2$ g$^{-1}$ respectively at 525 nm). These values are in the upper range when compared to MSEs reported in the literature for the ammonium nitrate specie calculated through stoichiometry. Hand and Malm (2007) determined MSEs of $3.2 \pm 1.2$ m$^2$ g$^{-1}$ for dry PM$_{2.5}$ ammonium nitrate, Cheng et al. (2015) obtained values of $4.3 \pm 0.6$ m$^2$ g$^{-1}$ under high mass loading in Shanghai, Tao et al. (2014) found MSEs ranging from $1.7 \pm 0.8$
in summer to 6.7 ± 1.8 m² g⁻¹ in winter in Chengdu (China) and Titos et al. (2012) observed a coefficient of 5 ± 2 m² g⁻¹ for nitrate ion in an urban area in southern Spain. MSES for secondary sulfate were quite different between MSY and MSA (4.5 ± 0.2 and 10.7 ± 0.5 m² g⁻¹), probably due to differences in the source origin and the related particle size. Hand and Malm (2007) published lower values for the total mode of dry ammonium sulfate ranging between 0.8 and 2.4 m² g⁻¹, whereas a MSE of 3.5 ± 0.5 m² g⁻¹ was found by Cheng et al. (2015) in a polluted environment. Tao et al. (2014) showed MSES of 4.4 ± 0.7 and 5.7 ± 0.2 m² g⁻¹ in winter and summer respectively for the PM₂.₅ fraction in Chengdu. MSES for non-sea-salt (nss) sulfate ion were calculated at Finokalia and Erdemli from the slope between total scattering and nss sulfate concentration, showing values of 5.9 ± 1.8 and 5.7 ± 1.4 m² g⁻¹ (Vrekoussis et al., 2005). Higher MSES were found in an urban background in the south of Spain (Titos et al., 2012) and in the Negev desert (Formenti et al., 2001) at 7 ± 1 and 7 ± 2 m² g⁻¹. Given that in our study sulfate concentrations were mainly explained by secondary sulfate and V-Ni sources, significant differences were also observed for the MSE of the V-Ni bearing source at MSY and MSA (8.0 ± 1.5 and 3.5 ± 0.5 m² g⁻¹). The V-Ni bearing source at MSY originated mainly from shipping emissions at regional (vessel traffic in the Mediterranean) and local (Barcelona harbour) scales. Conversely at MSA, located at a higher altitude, this source might also be influenced by continental transboundary transport and then internally mixed with different chemical species. In fact, as shown in Fig. 2a for MSA and in Fig. S2 for MSY, the V-Ni bearing source profile at MSA is enriched in OC, which is not observed at MSY. The aged marine source at MSA showed negative MSES at 525 and 635 nm. This is likely due to the larger distance from the coast of MSA, thus preventing a strong signal from the aged marine source at this site, and/or due to the PM₂.₅ cut-off inlet used at the beginning of the measurement period which prevented the sampling of coarse particles. However, MSES for the aged marine source at MSY (1.2 ± 0.3 m² g⁻¹) exhibited values within the same range as those reported by Hand and Malm (2007) for coarse-mode sea salt (1.0 m² g⁻¹). The MSE for the mineral source (1.3 ± 0.1 and 1.1 ± 0.1 m² g⁻¹) was similar at MSY and MSA. This similarity could be explained by the low reactivity of mineral dust particles, which were mostly externally mixed with other chemical species. Thus, less chemical transformation can be expected for mineral particles during the transport towards the stations. Lower MSES were found for mineral matter by Hand and Malm (2007) (0.7 ± 0.2 m² g⁻¹), by Titos et al. (2012) in the urban background of Granada (0.2 ± 0.3 m² g⁻¹) and by Vrekoussis et al. (2005) in Erdemli (0.2 m² g⁻¹). Similar coefficients were obtained by Vrekoussis et al. (2005) in Finokalia (1 m² g⁻¹), and by Pereira et al. (2008) and Wagner et al. (2009) for mineral dust in Portugal, 1 ± 0.1 and 0.9 m² g⁻¹ respectively. The MSE for the aged organics source (1.4 ± 0.2 and 1.3 ± 0.3 m² g⁻¹) was also quite similar at MSY and MSA, probably due to similarities in the processes that govern the OA formation at both sites, which originated mainly from local/regional biogenic emissions and SOA formation (Minguillon et al., 2015; Ripoll et al., 2015a). A similar MSE (1.4 m² g⁻¹) was reported by Hand and Malm (2007) for the total mode of primary organic matter (POM). Higher MSES were found by Cheng et al. (2015) during a pollution episode (4.5 ± 0.7 m² g⁻¹) and by Tao et al. (2014) in China (4.8 ± 0.8 and 6.5 ± 0.5 m² g⁻¹ in summer and winter). The industrial/traffic source showed similar MSES at MSY (2.1 ± 0.8 m² g⁻¹) and MSA (2.3 ± 0.5 m² g⁻¹). This similarity was related to the common origin of this source at both sites (i.e. emission from the traffic and industrial activities). It is remarkable that MSES for some of the sources identified in this work which contribute greatly to air quality degradation, such as industrial/traffic or V-Ni, are not available in the literature.

Prior studies dealing with the absorption efficiency of aerosol particles referred mainly to BC particles and to the possible effect of coating with non-absorbing material (Bond et al., 2013; Ramana et al., 2010). Other studies have reported the MAE of mineral matter (Linke et al., 2006) and OA (Lu et al., 2015; Updyke et al., 2012) due to the significant contribution of BrC to UV light absorption. However, to the author’s knowledge, this is the first time that absorption efficiencies, as well as scattering efficiencies, are computed for aerosol particle sources. MAE values at 637 nm for the three sites are summarized in Table 2. The highest absorption efficiencies were observed for the traffic source identified at BCN (1.672 ± 0.050 m² g⁻¹) and for the industrial/traffic source at MSY (0.867 ± 0.047 m² g⁻¹) and MSA (0.206 ± 0.02 m² g⁻¹) due to their content in BC particles from fossil fuel combustion. The V-Ni bearing source, which contributed greatly to light scattering, also exhibited high MAE at BCN (0.928 ± 0.058 m² g⁻¹) with decreasing coefficients at MSY (0.526 ± 0.065 m² g⁻¹) and MSA (0.165 ± 0.017 m² g⁻¹). We have shown here that the V-Ni bearing source, which is progressively becoming more relevant for air quality degradation due to the increased shipping emissions in recent years (Viana et al., 2014), also has an important effect on light absorption as consequence of the internal mixing with combustion aerosols. The large MAE observed for secondary nitrate at MSA (0.364 ± 0.023 m² g⁻¹) was due to the fact that this source explained around 20% of the measured EC concentration (Fig. 2a). Recently, Ripoll et al. (2015b) have shown the increased concentration of nitrate, ammonium, EC and traffic/industrial tracers at MSA under European scenarios. Such scenarios are characterized by the transport of polluted air masses at high altitude from central and eastern Europe to the MSA site. This fact may explain the internal mixing of BC particles in the chemical profile of secondary nitrate, and consequently the high MAE values found for this source at MSA. Lower MAEs for the secondary nitrate source were observed at BCN and
MSY (0.28 ± 0.040 and 0.234 ± 0.028 m² g⁻¹) compared to MSA. Lower MAEs were observed for secondary sulfate (0.359 ± 0.035, 0.122 ± 0.010 and 0.173 ± 0.021 m² g⁻¹) at BCN, MSY and MSA. Overall, higher absorption efficiencies were observed for the main anthropogenic sources at BCN, where fresh primary pollutants, mostly composed of darker particles, are emitted within the metropolitan, industrial and harbour areas. However, lower MAEs were found for the same pollutant sources at MSY and MSA. This result points to a decrease in the absorption efficiency towards inland areas, as a consequence of the different mixing and aging of pollutants during the transport towards the stations. Aerosol sources dominated by natural contributions, such as aged marine and mineral sources, showed the lowest MAE at MSY and MSA. The road dust resuspension source, which was partially composed of mineral matter, exhibited the lowest MAE at BCN (0.062 ± 0.084 m² g⁻¹). The mineral source presented MAE values of 0.09 ± 0.05, 0.005 ± 0.007 and 0.03 ± 0.003 m² g⁻¹ at BCN, MSY and MSA. Coefficients in the same order of magnitude at 660 nm were found for the Sahara–Sahel and Gobi deserts, ranging between 0.01 and 0.02 m² g⁻¹ (Alfaro et al., 2004), and for El Cairo and Morocco, 0.02 ± 0.004 and 0.06 ± 0.014 m² g⁻¹ (Linke et al., 2006). Aged marine also exhibited low absorption efficiencies at BCN, MSY and MSA (0.108 ± 0.021, 0.027 ± 0.018 and 0.015 ± 0.010 m² g⁻¹), but was higher at BCN due to a possible mixing with darker particles in the urban area. Similarly to the results observed for MSEs, aged organics showed similar MAEs at MSY and MSA (0.169 ± 0.011 and 0.140 ± 0.009 m² g⁻¹) due to the local/regional origin of this source with a similar composition at both sites. The absorption efficiency of this latter source was mainly explained by the EC contained within the source chemical profile, but also might be partially due to the presence of light-absorbing material detected as OC, such as BrC (Putaud et al., 2014).

3.3.1 SAE and SSA of aerosol sources

The source-specific scattering Ångström exponents (SAEs) were calculated as a linear fit of 3λ MSEs in the 450–635 nm range (Table 2). The MSE values used for computing SAE are shown in Table 2. The SAE parameter provides information on the size of the particles; generally a SAE lower than 1 or higher than 2 indicates that the scattering is dominated by large or fine particles respectively (Schuster et al., 2006). Aged organics and V-Ni bearing sources showed the highest SAE at MSY (2.2 and 2.4 respectively) and MSA (3.6 and 2.2), pointing to a predominance of fine particles within these sources. Previous studies have demonstrated the strong contribution from shipping emissions to fine aerosols (Viana et al., 2009), and especially to ultrafine particles (Saxe and Larsen, 2004). As reported in Table 2 for both MSY and MSA, the SAE of secondary sulfate (1.9 and 1.8) and secondary nitrate (1.2 and 1.4) sources was lower compared to the SAE of the aged organics source. This was probably due to the contribution of very fine primary organic aerosols (POAs) to the aged organic source, whereas both SOAs and secondary inorganic aerosols are expected to strongly contribute to the accumulation mode (Sun et al., 2016). The lowest SAE was observed for mineral (~0.9 at MSY and MSA) and aged marine (0.01 at MSY) sources, which primarily consist of coarse-mode particles. A relatively low SAE was found for the industrial/traffic source (0.6 and 0.8 at MSY and MSA respectively), which could be related to specific industrial processes in the area under study that include handling of dusty materials.

The single scattering albedo (SSA) coefficients obtained for each source at MSY are summarized in Table 2 and provide information on the relative importance of scattering or absorption in the light extinction process. The corresponding SSA to each source was computed as the ratio between the source-specific MSE and the sum of MSE and MAE (Table 2). As expected, the sources that internally mixed with combustion particles, such as industrial/traffic, aged organics and V-Ni, exhibited lower SSAs of 0.74, 0.84 and 0.9. Conversely, aged marine and mineral sources showed the highest coefficients of 1 and 0.98, leading to a scattering dominance in the light extinction process. Accordingly to studies in the literature, the mineral source showed a SSA close to 1. Linke et al. (2006) observed values around 0.98–0.99 at 532 nm, and lower coefficients were found by Müller et al. (2011c) for mineral dust (0.96) and marine (0.95) aerosols at 530 nm. Note that equivalent wavelengths should be considered when comparing SSA with coefficients in the literature due to the strong wavelength dependence of mineral dust particles.

3.4 Seasonal variation of source contributions to scattering and absorption

Monthly source contributions to the total scattering and absorption coefficients are shown in Fig. 3. The partial \( \sigma_{sp} \) and \( \sigma_{ap} \) apportioned to each source was calculated as the product between the aerosol source contribution and the corresponding MSEs or MAEs (Eqs. 1 and 2). According to the scattering efficiencies previously reported in Table 2, average scattering for the whole period was mainly dominated by secondary sulfate (35 and 33 % at MSY and MSA) and secondary nitrate (24 and 21 %) (Fig. 3g and h). The annual cycle of secondary sulfate and secondary nitrate scattering coefficients followed those of the PM10 mass concentration, with maxima in summer (46 and 35 % at MSY and MSA) and winter (42 and 29 %) respectively. The scattering contribution from aged organics accounted for 11 and 16 % of the total \( \sigma_{sp} \) at MSY and MSA. The V-Ni bearing source exhibited substantial contribution to \( \sigma_{sp} \) in summer (16 %), despite the relatively low contribution to PM10 mass concentration (5 % and 10 % at MSY and MSA). Less relevant were the scattering contributions from industrial/traffic (6 and 11 % at
MSY and MSA) and mineral (7 %) sources, peaking in winter and summer respectively. The traffic source at BCN and the industrial/traffic source at MSY clearly exerted a major influence on light absorption contributing 54 and 41 % to \( \sigma_{ap} \) respectively, despite the relatively low PM\(_{10} \) contributions (16 and 10 %). Maxima contributions were observed in winter at BCN for the traffic source (65 %) and in October–January at MSY for the industrial/traffic source (46 %), whereas a lower influence of industrial/traffic was observed on average at MSA (18 %) (Fig. 3 d, e, f). Interestingly, the V-Ni bearing source also played an important role in light absorption, especially in summer as a consequence of the increased vessel traffic in the Mediterranean but also because of the more intense sea breeze circulations, transporting pollutants to inland regions. Average contributions to \( \sigma_{ap} \) in summer were 31 % at BCN, 17 % at MSY and 16 % at MSA. Therefore, traffic, industrial/traffic and V-Ni bearing sources, which highly influenced air quality, also significantly contributed to \( \sigma_{ap} \), especially in those sites closer to the emission sources. Aged organics became a relevant source in the absorption process at the Regional and remote background sites, contributing on average 20 and 32 % due to both its large contribution to PM\(_{10} \) and its relatively large MAE compared to other sources. Secondary sulfate contributed on average 10, 16 and 12 % to the total \( \sigma_{sp} \) at BCN, MSY and MSA, whereas secondary nitrate showed increasing contributions to \( \sigma_{ap} \) towards inland areas (8, 10 and 21 %), markedly maximizing during the colder months.

### 3.5 Reconstruction of scattering and absorption coefficients

Scattering (\( \sigma_{sp} \)) and absorption (\( \sigma_{ap} \)) time series were reconstructed by means of the partial sum of the particle scattering and absorption contributions determined for each source (Eqs. 1 and 2). Strong correlations were found between the measured and modelled extensive optical parameters at the three sites (Fig. 4). Results showed good agreement for \( \sigma_{ap} \) at 525 nm at MSY \((R^2 = 0.88)\) and MSA \((R^2 = 0.92)\). \( \sigma_{ap} \) at 637 nm also exhibited a good correlation when comparing measured and predicted coefficients at BCN \((R^2 = 0.81)\), MSY \((R^2 = 0.80)\) and MSA \((R^2 = 0.93)\). Slopes were close to one in all cases and ranged between 0.96 and 0.98. These results are consistent with the good agreement obtained in the MLR model for MSE and MAE calculation. A \( R^2 \) of 0.96 was obtained for all the cases ensuring the accuracy of the regression coefficients computed for each site. The root mean square error (RMSE) was calculated for the observed–modelled data sets, showing low dispersion and high accuracy in the modelled values. Scattering and absorption coefficients were well reproduced by the model, showing RMSE values of 8.76 and 6.06 Mm\(^{-1}\) for \( \sigma_{sp} \) at MSY and MSA, and values of 2.61, 0.55 and 0.23 Mm\(^{-1}\) for \( \sigma_{ap} \) at BCN, MSY and MSA. The fractional bias (FB) between measured and predicted coefficients was calculated for each sampling site following Eq. (3). Results are shown in Fig. 5, where the FB is broken down by quintile from lowest to highest \( \sigma_{sp} \) and \( \sigma_{ap} \) values. According to published results (Ryan et al., 2005 and references therein), a consistent overestimation was observed for all the modelled coefficients in the lower ranges of \( \sigma_{sp} \) and \( \sigma_{ap} \), showing the highest bias in the 1st quintile. Biases were substantially reduced in the median range values, whereas a minor underestimation was observed for the highest \( \sigma_{sp} \) and \( \sigma_{ap} \) values, 4th and 5th quintiles, with negative FB. On average, a 3.8 and 5.6 % overprediction was obtained for the modelled \( \sigma_{ap} \) coefficients at MSY and BCN, using 503 and 375 daily data points in the analysis. A \( \sigma_{sp} \) overprediction at MSY pointed to 4 % using 307 daily points. However at MSA, \( \sigma_{sp} \) and \( \sigma_{ap} \) coefficients biased the observed values by 30.9 and 19.9 %, considering 220 and 303 data points in the analysis. A larger overestimation of the measured coefficients at MSA might be mainly explained by the lower number of daily chemical data used in the PMF model for the quantification of source contributions, but also because of the lower number of scattering and absorption data points available for the MLR analysis.

An independent subset of the study period was considered in order to further evaluate the PMF–MLR technique and the accuracy of the method to simulate optical properties when chemical source contributions were available. Therefore, a new PMF was performed in order to obtain the source contributions for the period 2004–2015 at MSY. With this aim, the simulation of \( \sigma_{sp} \) and \( \sigma_{ap} \) coefficients for the period January–December 2015 was carried out by means of the source-specific MSEs and MAEs previously obtained in the MLR analysis for the period 2010–2014. Good agreement was found between modelled and measured \( \sigma_{ap} \) \((R^2 = 0.85)\) and \( \sigma_{ap} \) \((R^2 = 0.76)\) coefficients, at 525 and 637 nm respectively, showing slopes close to one for the year 2015 at MSY (Fig. 6). This analysis confirms the confidence of the PMF–MLR technique to accurately estimate \( \sigma_{sp} \) and \( \sigma_{ap} \) coefficients when chemical data are available.

As a result of the aforementioned sensitivity test, long-term time series of \( \sigma_{sp} \) and \( \sigma_{ap} \) were satisfactorily reconstructed for the period 2004–2014 at BCN and MSY and for the period 2011–2014 at MSA (Fig. 7), when PM\(_{10} \) chemical speciated data were available.

### 3.6 Long-term trends in scattering and absorption coefficients at MSY

Long-term trends of \( \sigma_{sp} \) and \( \sigma_{ap} \) and their relationship with the trends of PM\(_{10} \) source contributions were investigated for an 11-year period at MSY (2004–2014). The trend of \( \sigma_{ap} \) at BCN was not studied due to the change in the location of the BCN sampling station in 2009 (Pandolfo et al., 2016), which affected mainly the contribution from the traffic source. The short time series available for chemical species concentration at MSA made the analysis of \( \sigma_{sp} \) and \( \sigma_{ap} \) trends unfeasible at this station.
Figure 4. Relationship between modelled and measured optical parameters: absorption at 637 nm for (a) MSA, (b) MSY and (c) BCN and scattering at 525 nm (d and e for MSA and MSY).

Figure 5. Average fractional bias (FB) calculated for the observed-modelled data pairs of scattering (Sc) and absorption (Abs) coefficients at BCN, MSY and MSA broken down by quintile from the lowest to highest scattering and absorption coefficient values. n accounts for the number of daily data points used in the FB calculation.

Temporal trends of the deseasonalized monthly averages for the modelled $\sigma_{sp}$ and $\sigma_{ap}$ at MSY for the period 2004–2014 are shown in Table 3. The multi-exponential (ME) approach allowed $\sigma_{sp}$ and $\sigma_{ap}$ time series to decompose in main, seasonal and residual components (Fig. 8). Linear trends were identified for $\sigma_{sp}$ and $\sigma_{ap}$, given that the non-linearity (NL) parameter was less than 10% (Shatalov et al., 2015). Statistically significant decreasing trends were found for both $\sigma_{sp}$ and $\sigma_{ap}$ at MSY (Table 3). $\sigma_{ap}$ decreased by $-4.1$ yr$^{-1}$ ($-0.16$ Mm$^{-1}$ yr$^{-1}$), whereas a reduction of $-4.6$ yr$^{-1}$ ($-2.14$ Mm$^{-1}$ yr$^{-1}$) was obtained for $\sigma_{ap}$ at 635 nm. Very similar trends were observed for $\sigma_{sp}$ at 450 ($-4.4$ yr$^{-1}$) and 525 nm ($-4.5$ yr$^{-1}$).

According to these results, decreasing trends were also observed for the majority of the PM$_{10}$ source contributions identified at MSY for the period 2004–2014, except for aged organics and aged marine sources (Pandolfi et al., 2016). A reduction in the absorption coefficient was directly related to the significant decreasing trends found by Pandolfi et al. (2016) for strong light-absorbing sources, such as industrial/traffic ($-5.09$ yr$^{-1}$) and V-Ni bearing source ($-5.82$ yr$^{-1}$). The observed scattering decreasing trend could be mainly associated with a reduction in the contributions from those sources which scattered light more efficiently, i.e. secondary nitrate and secondary sulfate. In Pandolfi et al. (2016) these sources showed reduction rates of $-6.27$ and $-4.82$ yr$^{-1}$ respectively. A marked decline has also been observed for nitrate and sulfate particles in other European monitoring sites since 1990, as outlined in the EMEP report 1/2016 (Colette et al., 2016). Other studies have been published in recent years, clearly showing that the concentrations of PM and other air pollutants, such as SO$_2$ and NO$_2$, have markedly decreased during the last 15 years.
Figure 6. Relationship between modelled and measured (a) scattering at 525 nm and (b) absorption at 637 nm at MSY for the period January 2015–December 2015.

Table 3. Theil-Sen (TS) trends at a 95 % confidence level for deseasonalized monthly averages of scattering absorption time series at MSY during the period 2004–2014. AR (Mm⁻¹; %) is average reduction; TR (%) is total reduction; the significance of the trends (p value trend) was obtained by means of TS method using monthly averages: *** (p value < 0.001), ** (p value < 0.01), * (p value < 0.05). The non-linearity parameter (%) was calculated by means of the multi-exponential (ME) test.

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in many European countries (EEA, 2013; Barmpadimos et al., 2012; Cusack et al., 2012; Querol et al., 2014; among others). Querol et al. (2014) and Pandolfi et al. (2016) investigated trends of PM chemical components and aerosol sources at MSY, providing further explanation on the causes leading to the reduction of the atmospheric pollutants in the area. The financial crisis affecting Spain from 2008 contributed a reduction in the ambient PM concentrations. A decrease in secondary nitrate can be explained by the reduction of ambient NOₓ and NH₃ concentrations (Querol et al., 2014). The decreasing trend of the secondary sulfate source may be supported by the reduction of sulfate particles, mainly attributed to the gas desulfurization at several facilities (Pandolfi et al., 2016). A decrease in secondary sulfate may be also explained by the 75 % reduction of SO₂ concentration in the Barcelona harbour, supported by the regulation of sulfur content in shipping emissions in EU harbours from 2010 (Schembari et al., 2012). This regulation together with the 2007 ban around Barcelona on the use of heavy oils and petroleum coke for power generation, which contributed to a drastic decrease in V and Ni concentrations (Querol et al., 2014), were the main causes of the observed reduction of the contribution of the V-Ni bearing source. This source was enriched in sulfate and combustion particles as shown in the source chemical profile, and, as was already observed, contributed simultaneously to both σsp and σap. Thus, the abatement strategies adopted in the recent years might have caused changes in the internal mixture of particles emitted from the V-Ni bearing source, and consequently in the contribution of this source to light extinction.

Only a few studies have been published in Europe aiming to study trends of particle optical properties. Statistically significant downward trends of PM mass concentration, σsp, σap and SSA were found in the Po valley (Italy) for the period 2004–2010 (Putaud et al., 2014). A higher decreasing rate was observed for σsp (−2.8 % yr⁻¹) compared to σap (−1.1 % yr⁻¹), likely due to the increasing contribution of light-absorbing organic matter to light absorption during cold months in the Po Valley (Putaud et al., 2014). In the present study, smaller differences between σsp and σap were observed at MSY, accounting for the total reduction trends of −50 and −45 % respectively. This fact might be explained by the different background sites considered, whereas the Po Valley is a highly polluted area, MSY is representative of a cleaner environment where biomass burning emissions, which contribute greatly to light absorption, are considerably lower (Minguillón et al., 2015; Ealo et al., 2016).

Further research on light scattering and absorption long-term trends and their relation to changes in atmospheric composition is needed to better understand the role of aerosols
Figure 7. Time series of the daily average modelled and measured extensive optical coefficients (scattering at 525 nm and absorption at 637 nm) for (a) BCN and (b) MSY during the period 2004–2014, and for (c) MSA during the period 2010–2014.
on optical properties and on the climate system. Based on the published studies and the present results, further efforts focusing on the reduction of atmospheric pollutants containing BC particles (mainly emitted from fossil fuel combustion and biomass burning sources) need to be addressed. Given the toxicity of their chemical tracers, as well as their large contribution to light absorption, industrial/traffic and V-Ni bearing sources must be reduced through the implementation of win–win policies which aim to improve air quality and public health and mitigate climate warming.

4 Summary and conclusions

Mass scattering and absorption efficiencies (MSEs and MAEs) of different aerosol particle sources were investigated at urban, regional and remote backgrounds in the NW Mediterranean, using unique large data sets of PM$_{10}$ chemical speciation and particle optical properties. For this purpose, a new approach was presented that aimed to apportion the PM$_{10}$ source contributions arising from a PMF model to the measured particle $\sigma_{sp}$ and $\sigma_{ap}$ coefficients.

Seven aerosol sources were identified at the Montsec (MSA) mountain-top site, where aged organics (29 %) was the foremost constituent of PM$_{10}$, followed by mineral (24 %), industrial/traffic (11 %), aged marine (16 %), secondary sulfate (9 %), V-Ni bearing (8 %) and secondary nitrate (7 %). The same sources were found at the regional background of Montseny (MSY), showing that relative PM$_{10}$ contributions of secondary aerosol sources were higher at the background sites than at the urban station in Barcelona (BCN). Aged organics was not identified at BCN; however specific pollutant sources related to the direct anthropogenic emissions were isolated (traffic, industrial/metallurgy and road dust resuspension). The impact of aerosol sources and the different chemical profiles obtained at the three sites depended on the distance and transport of pollutants to inland areas, driven by orography and meteorology.

The highest absorption efficiencies were attributed to aerosol sources internally mixed with BC particles. The traffic source at BCN (MAE = 1.7 m$^2$ g$^{-1}$) and the mixed industrial/traffic source at MSY (MAE = 0.87 m$^2$ g$^{-1}$) exerted a major influence on light absorption and reached the highest contributions during the colder period (65 and 46 %). The V-Ni bearing source was the second most efficient light-absorbing source in BCN (MAE = 0.93 m$^2$ g$^{-1}$), showing also a notable absorption efficiency at MSY and MSA (0.53 and 0.16 m$^2$ g$^{-1}$). This source contributed greatly in summer to both $\sigma_{sp}$ (16 % at MSY and MSA) and $\sigma_{ap}$ (31, 17 and 16 %, at BCN, MSY and MSA) due to the internal mixing of sulfate and combustion aerosols. These combustion sources were relevant but not dominant at MSY and MSA, where secondary aerosol sources (secondary sulfate, secondary nitrate and aged organics) gained relative importance in the light extinction process. A high spatial variability of MAE was observed for most of the anthropogenic sources, from high values at the BCN site to decreasing coefficients at the background stations, pointing to the aging and mixing state of aerosols as key factors influencing light absorption.
The highest scattering efficiencies were observed for secondary sulfate (4.5 and 10.7 m$^2$ g$^{-1}$ at MSY and MSA), secondary nitrate (8.8 and 7.8 m$^2$ g$^{-1}$) and V-Ni bearing (8 and 3.5 m$^2$ g$^{-1}$) sources, dominating the scattering throughout the year with marked seasonal cycles. Secondary nitrate contributed greatly in winter (42 and 29 % at MSY and MSA), whereas in summer secondary sulfate (46 and 35 %) was the main contributor to scattering.

Sources internally mixed with relatively dark and fine particles and greatly contributing to light absorption, such as industrial/traffic, aged organics and V-Ni, were simultaneously characterized with low single scattering albedo (SSA) and a high scattering Angström exponent (SAE). Conversely, mineral and aged marine showed the lowest SAE and the highest SSA, scattering being the dominant process in the light extinction. These findings for the intensive parameters were consistent at MSY and MSA. The observed variability in the intensive optical properties of aerosol sources provides valuable constraints for future simulations of aerosol parameters.

Significant decreasing trends were observed for the modelled scattering (−4.6 % yr$^{-1}$) and absorption (−4.1 % yr$^{-1}$) series at MSY for the period 2004–2014. The scattering reduction was mainly attributed to the decrease in the contributions from secondary nitrate, secondary sulfate and V-Ni bearing sources, whereas the absorption decreasing trend was mainly related to the decrease in industrial/traffic and V-Ni bearing sources. Given the toxicity of their chemical tracers, as well as the large contribution to light absorption, further efforts need to be addressed to reduce aerosol sources containing combustion particles, such as industrial/traffic and V-Ni bearing sources. However, further studies focusing on the study of long-term trends of optical parameters and their relationship with changes in atmospheric composition are needed to assess future win–win mitigation strategies.

Findings from the PMF–MLR technique are summarized as follows:

- The apportionment of PM source contributions to scattering and absorption allows the determination of MSEs and MAEs of atmospheric aerosol sources, taking into account the particle-internal mixing.

- Knowledge of both MSEs and MAEs makes it possible to study the existing relationship between the sources contributing to air quality degradation and their potential to absorb and scatter visible light. Anthropogenic sources such as secondary sulfate, secondary nitrate, traffic, industrial/traffic and V-Ni bearing source, which contribute greatly to air quality degradation, also revealed a substantial contribution to light extinction in the NW Mediterranean.

- For the first time, to the author’s knowledge, this work quantifies the absorption efficiency exerted by the different aerosol sources constituting the PM$_{10}$ mass concentration, in contrast to previous studies where light absorption was entirely attributed to BC particles. Interestingly, secondary sulfate, secondary nitrate and organic aerosols, for which light-absorbing properties are poorly represented in current climate models, significantly contributed to light absorption due to internal mixing with BC or BrC particles, and especially in regional and remote areas.

- The proposed approach allowed a satisfactory reconstruction of $\sigma_{ap}$ and $\sigma_{sp}$ compared to previous studies, given that the sum of the source contributions used in the MLR model reached around 100 % of the measured PM$_{10}$ mass concentration. Correlation coefficients are higher than 0.8 with slopes close to 1.0 between modelled and measured $\sigma_{ap}$ and $\sigma_{sp}$.

- Statistically significant decreasing trends were observed for the modelled $\sigma_{sp}$ and $\sigma_{ap}$ series, mirroring the effectiveness of the mitigation strategies adopted to improve air quality. The simultaneous analysis of the trends of climate relevant aerosols parameters ($\sigma_{sp}$ and $\sigma_{ap}$) together with the trends of PM source contributions allowed us to study the effects that the abatement strategies implemented in the last years are having on atmospheric composition and light extinction.

Data availability. The Montseny and Montsec data sets used for this publication are accessible online on the WDC (World Data Centre for Aerosols) web page: http://ebas.nilu.no (IDA-A-CSIC, 2017). The Barcelona data sets were collected within different national and regional projects and/or agreements and are available upon request.

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

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