# 1 Answer to the comments from Referee#1

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We thank the Reviewer#1. With his/her comments the quality and readability of the manuscript have
been strongly improved.

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The GREEN colour was used to highlight changes in the revised manuscript rose from the comments
 from Referee#1.

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# 10 General Comments.

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12 In this paper a trend analysis of PMx concentrations recorded at two different sites in NE Spain during 13 the period 2004-2014 has been performed. PM10 and PM2.5 chemical composition and PM10 source 14 contributions have also been evaluated with the aim to obtain more accurately interpretations of the 15 trends as well as of the effectiveness of the pollution control measures implemented in this period by the 16 administrations. Two different methodologies have been used. Namely, the Mann-Kendall test (MK) and 17 the Multi-Exponential fit (ME). In brief, I think that the present work shows some significant and novel 18 contributions to the global scientific community in relation with trend analysis of atmospheric pollutants 19 issues. I think that the availability of time series of PM chemical composition and of estimations of PM source contributions from receptor models, such as PMF, is nowadays a key factor for establishing 20 21 reliable source-receptor relationships and obtaining robust results from trend analysis and even 22 epidemiological analysis. However, some significant changes must be performed to clarify the usefulness of the different methodologies employed, to reduce the excess of information provided in the 23 24 manuscript that makes the reading very hard, and to justify the behaviour observed of the trends of some 25 of the time series analyzed.

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# 27 Specific Comments.

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29 1) In spite of the fact that the main results of the work (the pollution control measures effectively produced 30 a reduction of the contributions from some anthropogenic sources in such a way that the PMx levels decreased at the urban-background and the regional background sites) are rather consistent and well 31 justified, it is not clear the advantage of using simultaneously both trend analysis approaches, MK and 32 ME. In general when the ME showed a linear fit, the MK also showed a statistically significant linear 33 34 trend, but sometimes it is not statistically significant, as in the case of Zn and Na in PM10 for the 35 Montseny site. How should we interpret these different behaviours?. In the case that the MK showed a highly significant linear reduction trend and the ME showed a double exponential fit, Cd in PM10 for the 36 37 Barcelona site, what result must prevail, the MK or the ME one?. Please, try to clarify the best way to 38 take advantage of using both methodologies.

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In order to take into account the Referee's comment and to reduce the excess of information provided inthe manuscript the following changes were performed:

- 42 In the revised version of the manuscript ME and MK tests are not presented together and or MK results or ME results a) 43 are presented. Thus, when the trend is exponential (linear) the results from Mann-Kendall test (Multi-exponential 44 test) are not presented. In this way, we highlight in the manuscript the fact that the two fits are different and that the 45 MK test does not properly fit the data when these have actually an exponential trend. Similarly, there is no need to 46 use the ME test when the trend is linear. This will avoid creating confusion for the readers. In fact, when the trend is 47 exponential (single or double) the results from the MK test are always statistically significant; however data are not 48 well fitted using MK in this case. On the other side, when the trend is linear (not exponential) we do not need the ME 49 test and the trend can be or not statistically significant. Moreover, with this change we also highlight in the manuscript 50 the main difference between the MK and ME fits: when a trend is exponential it means that the observed decreasing 51 trend was not constant and gradual over time (as indicated by the MK fit).
  - b) We also removed from the manuscript (Figures and Tables) those species or source contributions showing no statistically significant trends In this way we further reduce excess information presented in the manuscript. Species or sources showing no trend are now commented in the manuscript but are removed from Tables and Figures.

Why did you decide to work with annual mean values instead of monthly mean values?. It strongly
 reduced the number of data for the trend analysis.

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We show in the manuscript that despite the rather low annual data coverage of filter data, the trends of the annual means can be reasonably studied. However, the use of monthly means could introduce undesired noise given that filter collection is not always evenly distributed along the year (i.e. because of intensive campaigns, technical problems,....). Given that the basic information (trend magnitude and statistically significance) provided using annual or monthly means is similar we would like to use the annual means in our manuscript.

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64 3) In my opinion, the MK results can be easily interpreted by the potential readers. I mean, the test provides a statistically significant downward or upward trends for a given confidence level and the value 65 66 of an estimator of the trend (%variation/year). If the test provides a non-statistically significant trend, then 67 it must be interpreted as the absence of trend. In the case of the ME results it is not clear the meaning of linear, single exponential and double-exponential fit in relation with the trends of the pollutants. For 68 the double-exponential fit cases, the values of T1 and T2 are sometimes guite different, even positive 69 70 and negative. Despite some explanations of these values are included in the text, the interpretation remains somewhat obscure. Some more information should be included in the 2.5 section about the 71 72 interpretation of the equations and coefficients representing linear, single-exponential and double-73 exponential fit. Is the multi-exponential fit statistically significant in all the cases? I suppose that the fit 74 parameters an and Tn are estimated by the program with statistically significance for a given confidence 75 level. As in the case of the coefficients of a multilineal regression analysis. Is not it?. Please, try to clarify 76 all these questions in the revised version of the manuscript.

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We agree with the reviewer that the statistical significance of the exponential fits is missing in the manuscript. In the revised version of the manuscript p-values are provided for both linear (Mann-Kendall) and exponential fits. Tables 2,3 and 5 and text were opportunely modified. Moreover, the following text was added to Paragraph 2.5 in order to help the reader to interpret the results from the exponential fit.

82 "The main difference between linear and exponential fit is that in the latter case the trend is not gradual and constant over time. For an exponential trend the absolute  $[\mu g/m^3]$  reduction per year decreases with time being 83 the highest at the beginning of the period. Conversely, for a linear fit the absolute reduction is constant over time. 84 For an exponential fit, the lower the characteristic time  $\tau$  the more rapidly the considered quantity vanish. 85 86 Deviations from single exponential fit can be taken into account introducing more exponential terms. In this work 87 for example two exponential terms were sometime used. In this case, two characteristic times are calculated by the 88 software. If the decrease of the considered quantity is very sharp at the beginning of the period (more than 89 exponential) than both  $\tau_1$  and  $\tau_2$  are positive. Conversely, an exponential term with negative  $\tau$  takes into account for possible increases of the quantity at the end of the period. Both  $\tau_n$  and  $a_n$  are calculated by the program by means of the least square method minimizing the residue  $\omega$  and the statistically significance of the exponential fit is provided by means of the p-value."

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4) Otherwise, the ME and MK methodologies provides for any case a lot of information. I mean, the trend
estimation, p-value, type of fit... These results are summarized in tables 1-5. However, the authors
decided to include most of this information again in the text. As a consequence the reading is very hard
and confusing. For the MK results it is not necessary to include the magnitude of the trend and the degree
of statistical significance in all the cases. Please, try to highlight in the manuscript only the most important
information and refer to the tables for details.

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Following the reviewer comment, in the revised manuscript the text was shortened as much as possible in order to avoid repeated information. Moreover, as already stated in the answer to the *specific comment #1*, or MK or ME results (not both) are presented in the revised manuscript, thus further helping in making the text more readable.

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107 5) I would like to underline one important fact. Sometimes it can be read in the text statements as "non-108 statistically significant decreasing trend (-1.25 %/year" (page 10, lines 297-298). I completely disagree 109 with this. If the test provides a non-statistically significant trend result, you cannot assure the existence 110 of a trend, neither downward nor upward. That is the aim of performing statistical tests. Accepting or 111 rejecting null hypothesis with statistical significance. Hence, you can neither talk about increasing or 112 decreasing trend nor show the value of the estimator of the trend in the non-statistically significant trend 113 cases. I believe that you must rewrite the manuscript and the tables, excluding the values of the estimator 114 of the trends for non-statistically significant cases.

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We agree with the reviewer. Consequently, we removed from the manuscript (Figures and Tables) those species or source contributions showing no statistically significant trends. In this way we further reduce excess information presented in the manuscript. Species or sources showing no trend are now commented briefly in the manuscript but are removed from Tables and Figures.

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6) The reason why you decide not to use the data of mineral matter and road traffic emissions from the
Barcelona site before 2009, has been repeatedly mentioned across the manuscript.

- 124
- 125 And
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127 7) The statement that "2002-2014 represented the largest period of gravimetric PM2.5 measurements
128 available at MSY station" has been also repeated unnecessarily.

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Repetitions are avoided as much as possible in the revised manuscript.

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132 8) In page 10 (lines 304-310) and page 13 (lines 402-403) you mentioned results from subsequent
133 sections. I think it should be avoided for clarity.

### 134

The sentence at page 10, lines 304-310: "We will show later (*Paragraph 4.1*) that despite the fact that the gravimetric concentrations of PM<sub>10</sub> at MSY only slightly decreased monotonically with time (-0.47 %/yr with p>0.1; cf. Table 1), the contributions from specific PM<sub>10</sub> pollutant sources from PMF model related with anthropogenic activities showed non linear (i.e. exponential) statistically significant decreasing trends. For example the *Ammonium sulfate* source contribution to PM<sub>10</sub> at MSY decreased at the rate of -2.02 %/yr with p<0.05 310 from MK test and double exponential fit of the data was needed; cf. Table 5)."

- 141
- 142 Was replaced with:

143 "Given that the trends of the considered PMx fractions were linear at both sites (NL<10%), only results from MK 144 test were reported in Table 1. However, we will show later (Paragraph 4.1) that the contributions from specific 145 PM<sub>10</sub> pollutant sources from PMF model, mainly those related with anthropogenic activities, showed non-linear 146 (i.e. exponential) decreasing trends, thus mirroring the different effectiveness of the mitigation strategies depending 147 on the source of pollutants considered."

- 148 Moreover, the sentence at page 13, lines 402-403: "Interestingly, as shown later, the PM10 149 Industrial/metallurgy source contribution at BCN also showed a DE decreasing trend", was removed from the text.
- 150

151 9) In section 3.2 you attributed the differences observed in the magnitude of the trends for different time 152 periods to meteorology variability. In section 3.3 (lines 509-511) you stated that "It is probable that 153 variations in meteorological conditions from one year to another (i.e. intensity and frequency of Saharan 154 dust outbreaks) might also explain the observed trend of mineral tracers at regional level". In section 4.1 155 (lines 656-658) you also declared that "This decreasing trend could be due to a possible decrease of the 156 emissions of anthropogenic mineral species from specific sources such as cement and concrete 157 production and construction works". These comments are highly speculative due to the fact that neither 158 meteorological variables, nor information on "intensity and frequency of Saharan dust outbreaks", nor 159 information on "cement and concrete production and construction works" have been analyzed to support 160 them. In my opinion it should be mandatory to carry out an analysis of this kind of data to confirm these 161 hypotheses.

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In section 3.2 we compared the trends of  $PM_{2.5}$  mass concentrations over different periods with the main aim of: a) being consistent with what already published at MSY station by Cusack et al. (2011) where 9 yr of data (2002-2010) were used, and b) to study the differences in the trends over short periods (9 yr to 13 yr). The fact that the trends were statistically significant over different periods confirms the effectiveness of the measures taken to improve air quality.

168 To take into account the reviewer comment, the following sentences were removed from Pragraph 3.2:

169 170 171	"However, the difference observed in the magnitude of the trends during 2004-2014 compared to the results provided by Cusack et al. (2012) suggested that meteorology (in this case a large increase in 2012; cf. Figure 2), changing from year to year, also determined the degree of comparability of trends observed over different periods."
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173	and
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175 176	"Thus, over relatively short periods (9 -11 yr), the effects of just one meteorologically different year were clearly visible.".
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178	And the following sentence was added at the end of Pragraph 3.2:
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180 181 182 183 184 185	"However, it should be noted that the statistical significance of the trends observed for the larger periods was lower compared to Cusack et al. (2012). The difference observed in the magnitude of the trends during 2004-2014 compared to the results provided by Cusack et al. (2012) was mainly due to the increase of $PM_{2.5}$ mass concentration in 2012 (cf. Figure 2). Chemical $PM_{2.5}$ speciated data revealed that this increase was partly driven by organic matter showing a mean annual concentration in 2012 higher by around 20% compared to the 2004-2014 average."
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187 188 189 190	Moreover, in Paragraph 3.3 we reported that the total reduction observed for the concentrations of mineral species was higher in $PM_{2.5}$ mass fraction compared to $PM_{10}$ , thus suggesting an anthropogenic contribution to the mass of mineral matter in $PM_{2.5}$ and that this anthropogenic contribution likely decreased with time as already observed by Cusack et al. (2012).
191	In order to better explains possible reasons for the trends observed for mineral matter the following sentence:
192 193 194 195 196 197 198 199 200 201 202	"As for Cr, Sn, Sb and Cu, the trends of mineral species (Al2O3, Ca, Fe) were studied only at MSY station. For these elements, linear (with the exception of Al2O3 in PM2.5 which was SE) and statistically significant decreasing trends (with the exception of Ca in PM2.5 with p>0.1) were detected. On average the TR were higher in the fine fraction, ranging from 50% for Ca to 66% for Al2O3, compared to PM10 (16-38% cf. Table 2) thus likely suggesting a decrease with time of the concentrations of anthropogenic mineral species from specific sources such as cement and concrete production and production works. In fact, coarse mineral matter at regional background sites is mainly of natural origin. Downward decreasing trend for mineral matter contribution in PM2.5 at MSY was also reported by Cusack et al. (2012) for the period 2002 – 2010 at the same station. It is probable that variations in meteorological conditions from one year to another (i.e. intensity and frequency of Saharan dust outbreaks) might also explain the observed trend of mineral tracers at regional level."
203	Was replaced with the following sentence:

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205 "For the mineral species (Al2O3, Ca, Fe) linear (with the exception of Al2O3 in PM2.5 which was SE) and statistically significant decreasing trends were detected at MSY. On average the TR was higher in the fine fraction, 206 207 ranging from 50% for Ca to 66% for Al2O3, compared to PM10 (6-38% cf. Table 2). Downward decreasing trend for crustal material in PM2.5 at MSY was also reported by Cusack et al. (2012) for the period 2002 - 2010 and by 208

209 Querol et al. (2014) for the period 2001 - 2012. These trends were probably driven by weather conditions associated 210 with negative NAO index (iNAO) that could be the cause for this slight reduction observed in crustal material. Pey 211 et al. (2013) found a correlation between iNAO (calculated between June and September) and the contribution of 212 Saharan dust to PM10 mass in NE of Spain showing that the more negative is the iNAO the lower is the dust 213 The iNAO was unusually negative during the period 2008 – 2012 contribution to PM. 214 (http://www.cpc.ncep.noaa.gov/products/precip/CWlink/pna/norm.nao.monthly.b5001.current.ascii) thus likely 215 contributing to explain the observed trends of crustal elements. Moreover, negative NAO can favour the presence 216 of fronts that can sweep the Iberian Peninsula from West to East causing higher wind and less stagnant conditions 217 thus favouring the dispersion of pollutants. In addition, as suggested by Cusack et al (2012), it could also be 218 hypothesised that some part of the crustal material measured at MSY is a product of the construction industry. The 219 construction industry in Spain has been especially affected by the current economic recession, and crustal material 220 produced by this industry may have contributed to the crustal load in PM2.5. For example, the number of home 221 construction works in Barcelona during 2008 - 2014 (from the beginning of the economic crisis; mean number of 222 works = 1281) reduced by around 75% compared to the period 2000 - 2007; mean number of works = 5187) 223 (http://www.bcn.cat/estadistica/castella/dades/timm/construccio/index.htm). The fact that the total reduction 224 calculated for mineral elements reported in Tables 2 and 3 was higher in PM2.5 compared to PM10 could 225 corroborate this latter hypothesis."

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227 Moreover, the following sentence in Paragraph 4.1:

228 "The statistically significant decreasing trend observed at MSY for the Mineral source was in agreement 229 with what observed at the same station by Cusack et al. (2012). This decreasing trend could be due to a possible 230 decrease of the emissions of anthropogenic mineral species from specific sources such as cement and concrete 231 production and construction works."

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233 Was replaced with the following sentence:

"Finally, the Mineral source contribution at MSY showed linear little significant decreasing trend (p<0.1) in agreement with what observed at the same station by Cusack et al. (2012). As already noted in Paragraph 3.3, this negative trend could be due to both a possible decrease of the emissions of finer anthropogenic mineral species from specific sources such as cement and concrete production and construction works and unusual weather conditions reducing Saharan dust contribution to PM and resuspension of dust."

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### 240 Minor Comments.

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# 242 10) Did you achieve a PMF source contribution study with the PM2.5 data base?. It should be 243 interesting to compare the results of the trend analysis for PM10 and PM2.5 source contributions.

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Before starting with PMF analysis on PM10 chemical speciated data, the PM10 database for Montseny station was opportunely revised. Thus, before this work, only total carbon concentration was available for the period 2004 – 2007. In order to properly apply the PMF model, the concentrations of elemental (EC) and organic (OC) carbon in PM10 were recovered from filters sampled during the period 2004 – 2007. This was not done in PM2.5. Given that the data recover implies laboratory analysis which takes rather long time we decided to apply the PMF model to the PM10 mass fraction and to present trends of chemical species for both PM mass fractions. In order to not delay the publication of this work and with the permission of the Reviewer, we would like to present PM10 source apportionment results only.

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11) Page 6, lines 188-189. "(End user's guide to multilinear engine applications from Pentti Paatero)".
What do you mean?. Is that a reference?.

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- 257 The correct reference is (Paatero, 2004). We added the following reference to the bibliography:
- 258

"Paatero P.: User's guide for positive matrix factorization programs PMF2 and PMF3, Part1: tutorial. University
 of Helsinki, Helsinki, Finland, 2004"

261

12) Page 6, line 190. You declared that rotational ambiguity of the PMF solution was handled by means
of the Fpeak parameter. However, some better tests to estimate rotational uncertainty than Fpeak are
now available in the latest version of EPA PMF (V.5.0) such as the base model displacement error
estimation and other rotational tools. Have you checked these new options?.

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267 The aim of this work was to study trends of source contributions. To do this we applied the PMF model to 268 11 yr of chemical speciated data. Thus, we supposed that the chemical profiles of the detected sources did not 269 change with time. To test this hypothesis we performed a "sliding PMF" (using groups of 3 years of data: i.e. 2004-270 2006; 2005-2007; 2006-2008 and so on) comparing the retrieved source profiles. The results suggested that the 271 error in detecting source profiles and contributions was lower using the whole (11 years) database. For example, 272 the PMF was not able to clearly detect the Industrial/Traffic and V-Ni bearing sources at MSY at the end of the 273 considered period by means of the sliding PMF. Thus, we concluded that the best option was to use the whole 274 database as input in the PMF model even if changes in the chemical profiles cannot be excluded. However, only in 275 this way, the trends of the contributions of some sources such as Industrial/Traffic and V-Ni agreed well with the 276 trends of the main tracers of these sources. Thus, given the large period used for PMF analysis, the deviation from 277 constant profiles could largely affect the displacement results and errors.

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- 13) Page 7. Section 2.4. The description of the Mann-Kendall test is very short. Some
- 280 more information should be included.
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- 282 Paragraph 2.4 was modified as it follows:

283 "The purpose of the Mann-Kendall (MK) test (Mann 1945, Kendall 1975, Gilbert 1987) is to statistically 284 assess if there is a monotonic upward or downward trend of the variable of interest over time. A monotonic upward 285 (downward) trend means that the variable consistently increases (decreases) through time. The Mann-Kendall test 286 tests the null hypothesis H0 of no trend, i.e. the observations are randomly ordered in time, against the alternative 287 hypothesis, H1, where there is an increasing or decreasing monotonic trend. The main advantage of the Mann-288 Kendall test is that data need not conform to any particular distribution and missing data are allowed. To estimate 289 the slope of the trend the Sen's method was used (Salmi et al. 2002)."

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292	14) Page 8. Lines 230-232. What does Cbeg and Cend mean?.					
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294	The following sentence was added after Equation 5:					
295	"Where C <sub>beg</sub> and C <sub>end</sub> are respectively the first and the last points of the exponential fit."					
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298 299	15) Page 9. Lines 259-261. The description of the section was made before in the previous section. This paragraph can be omitted.					
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301	The paragraph was removed from the text					
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303 304	16) Page 9. Lines 263-264. "Note that the recommended annual data coverage for trend studies is typically 75%". Where does it come from?. Can you include the source in the Reference section?.					
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306 307 308 309	Recommended annual data coverage of 75% was recently set by the Task Force on Measurements and Modeling (TFMM-CLRTAP) in order to study trends of different pollutants in EU. However, given that this limit can be considered as quite arbitrary, the sentence was removed from the manuscript. Consequently, the paragraph was modified as follow:					
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311 312 313 314 315	"Annual data coverage is an important factor to take into account in order to study trends of a given parameter. The gravimetric PM measurements, from which chemical speciated data are obtained, are typically performed with rather low frequency over one year. In our case the annual data coverage of gravimetric measurements was around 20-30% at both Barcelona and Montseny. In this section we compare the trends of PM concentrations from gravimetric and real-time optical measurements (Table 1)."					
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318	17) Page 9. Lines 276-277. The results of the comparison between simultaneous PMx					
319	chemical speciated data collected at both BCN measurement sites can be showed as					
320	Supplementary Information.					
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322 323 324	The current location of the BCN measurement station is called <i>Palau Reial (PR)</i> . The location before 2009 was called <i>IJA</i> . The 1-month simultaneous filter measurements were performed at <i>IJA</i> and at a location (called <i>Torre Girona; TG</i> ) which was very close to <i>PR</i> but likely more affected by the presence of trees and some small					

325 326 327 328 329 330 331 332 333 334 335	buildings compared to <i>PR</i> . This is why we moved again from <i>TG</i> to <i>PR</i> . The ratios between the concentrations of chemical species simultaneously measured at <i>IJA</i> and <i>TG</i> were around $0.20 \div 0.60$ (with $R^2 < 0.60$ and with <i>TG</i> strongly underestimating <i>IJA</i> ) for mineral and traffic tracers (bad correlation). The ratios were around $0.70 \div 0.90$ (with $R^2 > 0.60$ and with <i>TG</i> slightly underestimating <i>IJA</i> ) for species not related with traffic and mineral sources (good correlation). Thus, we observed rather similar concentrations between <i>TG</i> and <i>IJA</i> for those species measured at BCN and included in the manuscript. Conversely, the bad correlation observed for mineral and traffic tracers confirmed that other traffic and mineral sources were affecting <i>IJA</i> but not <i>TG</i> . The lower concentration measured at <i>TG</i> compared to <i>IJA</i> for "good" species was very probably due to the presence of small buildings around <i>TG</i> . We are confident that the ratios between <i>PR</i> and <i>IJA</i> are very close to one. In conclusion, we used the measurements performed at <i>TG</i> and <i>IJA</i> in order to certainly exclude "bad" species from the analysis. For this reason and in order to avoid confusion we would like to not present this comparison in the manuscript.
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338 339	18) Page 10. Line 317. What is the definition of the Residual Component (RC)?. How did you computed it?. This information should be included in Section 2.
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341	The following sentence was added to Section 2.5
342 343	"The relative contribution of residues (Residual Component: RC) is calculated as the standard deviation of the ratios between the residue values of the fit $\omega$ (cf. Eq. 1) and the main component of the fit."
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346 347	19) Page 20. Line 633. Querol et al. 2007, is not included in the Reference section.
348	The following reference was added:
349 350 351 352 353	"Querol, X., Viana, M., Alastuey, A., Amato, F., Moreno, T., Castillo, S., Pey, J., de la Rosa, J., Artíñano, B., Salvador, P., García Dos Santos, S., Fernández-Patier, R., Moreno-Grau, S., Negral, L., Minguillón, M.C., Monfort, E., Gil, J.I., Inza, A., Ortega, L.A., Santamaría, J.M., Zabalza, J.: Source origin of trace elements in PM from regional background, urban and industrial sites of Spain, Atm. Env., 41, 7219-7231, 2007."
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355	Technical corrections/Typing errors.
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357 358	20) Figure 2 has very low quality. It is very hard to distinguish among the different symbols. The grey lines and symbols are very diffuse. This is also true for figures 3, 4, 6 and 7.
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360	All Figures were re-edited
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362	21) Table 4 is unnecessary. This information is showed in Figure 5.
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364	Table 4 was removed.
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391	Answer to the comments from Referee#3

393 We thank the Reviewer#3. With his/her comments the quality and readability of the manuscript have 394 been strongly improved.

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The **BOLD GREEN colour** was used to highlight changes in the revised manuscript rose from the comments from Referee#3.

398

The main problem with this manuscript is that it reads like a technical report rather than a scientific publication. Therefore, although the analysis made in this paper may be scientifically sound, the outline of the manuscript requires some fundamental revisions before I can recommend accepting this paper for publication. My detailed comments in this regard are given below.

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# 405 Major comments

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1) The authors should define clearly the scientific goals/aims of this paper. Currently, the last paragraph
of section 1 merely lists what has been done in the paper without specifying what the authors aim to
solve or find out in doing all this analysis.

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411 The Section 1 was changed accordingly to the Referee's comment and the following sentences were added:

412 "Thanks to the aforementioned measures, there is clear evidence that the concentrations of PM in many European 413 countries have markedly decreased during the last decades. However, in spite of the above policy efforts, a 414 significant proportion of the urban population in Europe lives in areas exceeding the World Health Organisation 415 (WHO) air quality (AQ) standards i.e. for  $PM_{2.5}$ ,  $PM_{10}$  and ozone (EEA, 2013, 2015).

Trend analysis of the concentration of air pollutants helps in evaluating the effectiveness of specific AQ measures depending on the pollutant considered. Examining data over time also makes it possible to predict future frequencies and/or rates of occurrence making future projections. For the abovementioned reasons, it is especially attractive the feasibility of studying the trends of the contributions to PM mass from specific pollutant sources along with the trends of the chemical tracers of these sources.

For what we are concerned in the majority of studies dealing with trend analysis, linear fits were applied for example by using Mann-Kendall or Theil-Sen methods (Theil, 1950; Sen, 1968), the latter being available for example in the Openair software (Carslaw, 2012; Carslaw and Ropkins, 2012). However, linear fit of data does not always properly represent the observed trends. As we will show, different abatement strategies and periods of implementation may change from one pollutant to another thus leading to different trends for different pollutants, even over the same period. Thus, non-linear fit of the data may be at times strongly recommended.

The main aim of this work was to study the trends of source contributions to PM10 and specific chemical species in both PM10 and PM2.5 using both the consensus methodology for linear fit of the data (Mann-Kendall) and a non-linear approach. The data of Spanish national emissions and energy consumption are also evaluated to interpret the observed trends. Understanding past trends may be relevant for devising new strategies for air pollution abatement. PM chemical speciated data collected from 2004 to 2014 at regional (Montseny; NE Spain) and urban 432 (Barcelona, NE Spain) sites were used with this aim. The selected period allowed for trend analysis at these twin 433 stations over a common period. The Positive Matrix Factorization (PMF) model was used to apportion ambient 434 PM10 concentrations into pollutant sources. The PMF model, as other Receptor Models (RM), is widely used being 435 a powerful tool to help policy makers to design more targeted approaches to protecting public health. Thus, the 436 novelty of this study lies mainly in a) the opportunity to study the trends of pollutant source contributions from 437 PMF model at two twin stations representative of the urban and regional environments in the Western 438 Mediterranean, and, b) in the use of a novel non-linear approach for trend studies."

439

440 2) Section 3. and especially 3.3. like a huge number of trend values which all can be found in the tables. 441 This not only makes the text extremely unappealing to read, but also the most important findings of this 442 analysis remain hidden behind these numbers. I strongly recommend the authors i) to shorten this 443 section considerably, ii) to remove most of the trend values from the actual text, and iii) to bring up more 444 explicitly the most important findings.

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446 In order to take into account the Referee's comment and to reduce the excess of information provided in the 447 manuscript the following changes were performed:

448 a) In the revised version of the manuscript ME and MK tests are not presented together and or MK results or ME results 449 are presented. Thus, when the trend is exponential (linear) the results from Mann-Kendall test (Multi-exponential 450 test) are not presented. In this way, we highlight in the manuscript the fact that the two fits are different and that the 451 MK test does not properly fit the data when these have actually an exponential trend. Similarly, there is no need to 452 use the ME test when the trend is linear. This will avoid creating confusion for the readers. In fact, when the trend is 453 exponential (single or double) the results from the MK test are always statistically significant; however data are not 454 well fitted using MK in this case. On the other side, when the trend is linear (not exponential) we do not need the ME 455 test and the trend can be or not statistically significant. Moreover, with this change we also highlight in the manuscript 456 the main difference between the MK and ME fits: when a trend is exponential it means that the observed decreasing 457 trend was not constant and gradual over time (as indicated by the MK fit).

458 b) We also removed from the manuscript (Figures and Tables) those species or source contributions showing no 459 statistically significant trends In this way we further reduce excess information presented in the manuscript. Species 460 or sources showing no trend are now commented in the manuscript but are removed from Tables and Figures. 461

- c) The data of Spanish national emissions and energy consumption are also evaluated to interpret the observed trends.
- 462

463 3) The paragraph on line 79-97 does not fit to the introduction of a scientific paper. Some of it could be 464 part of the methods section, if needed. Even then, any citations to Wikipedia are highly questionable.

- 465
- 466 This Paragraph was removed from Section 1 and moved to section 2.5:
- 467

#### 468 "2.5 Multi-exponential (ME) fit

469 A Program aiming at studying trends of time series of air pollution in the multi-exponential form was developed 470 within the The Task Force on Measurements and Modelling (TFMM) by the Meteorological Synthesizing Centre 471 - East (MSC-E; http://www.msceast.org/) group (Shatalov et al., 2015). The TFMM together with the Task Force 472 on Emission Inventories and Projections (TFEIP), the Task Force on Integrated Assessment Modelling (TFIAM), 473 and Task Force on Hemispheric Transport of Air Pollution (TFHTAP) provide a fora for discussion and scientific 474 exchange in support of the EMEP (European Monitoring and Evaluation Programme; http://www.emep.int/) work 475 plan which is a scientifically based and policy driven programme under the Convention on Long-range 476 Transboundary Air Pollution (CLRTAP; http://www.unece.org/env/lrtap/lrtap h1.html) promoting the 477 international co-operation to solve transboundary air pollution problems. The TFMM was established in 2000 to

478 evaluate measurements and modeling and to further develop working methods and tools. In this contest, five EMEP 479 Centers are undertaking efforts in support of the EMEP work plan, namely the MSC-E, the Centre on Emission 480 Inventories and Projections (CEIP; http://www.ceip.at/), the Chemical Coordinating Centre (CCC; 481 http://www.nilu.no/projects/ccc/), the Meteorological Synthesizing Centre West (MSC-W; \_ 482 http://emep.int/mscw/index\_mscw.html), and the Centre for Integrated Assessment Modelling (CIAM; 483 http://www.iiasa.ac.at/~rains/ciam.html). In 2014, the TFMM initiated a dedicated exercise to assess the efficiency 484 of air pollution mitigation strategies over the past 20 years to assess the benefit of the CLRTAP main policy 485 instrument. Within this exercise a software was made available by EMEP/MSC-E Center aiming at studying non-486 linear trends."

487

488 4) Section 5 is currently a single long paragraph. I would recommend the authors to organize this section
 489 better and perhaps to put different types of conclusions in separate pagraphs.

490

491 Section 5 was opportunely reorganized in order to highlight the main finding of this work:

492 *"5.0 Conclusions* 

493 PM chemical speciated data collected at two twin stations in NE of Spain (Barcelona: urban background station 494 and Montseny: regional background station) during 2004 – 2014 were used to study trends of source contributions 495 from PMF analysis and of chemical species concentrations. Despite the fact the trends of different PM fractions 496 (PM2.5 and PM10) were linear during the period under study, the trends of specific chemical elements and source 497 contributions were exponential demonstrating the different effectiveness and time of implementation of different 498 reduction strategies on specific pollutant sources. Statistically significant exponential trends (p < 0.01 or 0.001) 499 were mainly observed for the industrial tracers (Pb, Cd, As) in both PM10 and PM2.5 and at both sites. The 500 concentrations of V and Ni showed exponential trends in BCN and linear trends at MSY likely because of the higher 501 distance of the MSY station to the sources of V and Ni (shipping and, before 2008, energy production) compared 502 to BCN. Traffic tracers at MSY (Sn, Cu) showed very similar linear decreasing trends with higher magnitude of 503 the trends in the fine (PM2.5) fractions compared to PM10 likely because of possible sources of coarser Sn and 504 Cu reducing the magnitude of the trends in the PM10 mass fraction. Sb at MSY showed marked exponential 505 decreasing trends compared to other traffic tracers (Cu and Sn) which could be explained by a approgressive 506 reduction of Sb content in vehicle brakes. Secondary inorganic aerosols (SO42-, NO3- and NH4+) also showed 507 marked decreasing trends (both linear and exponential) in both fractions and at both sites. However, in general 508 the magnitude of the trends for these species and their statistical significance were higher at BCN compared to 509 MSY.

510 The PM10 source contributions that showed statistically significant downward trends at both Barcelona (BCN; 511 UB) and Montseny (MSY; RB) were from Ammonium sulfate, Ammonium nitrate, and V-Ni bearing sources. For 512 these source contributions the decreasing trends were exponential indicating that the trends were not gradual and 513 consistent over time and that the effectiveness of the control measures for these pollutants was stronger at the 514 beginning of the period under study (2004-2009 approximately) compared to the end of the period (Figs. 3 and 4). 515 Statistically significant decreasing trends were observed for the Industrial/Traffic and Mineral source (at MSY; 516 mixed road traffic and metallurgy) and the Industrial/metallurgy source at BCN. These sources were mostly linked 517 with anthropogenic activities and the observed decreasing trends confirmed the effectiveness of pollution control 518 measures implemented at EU or regional/local levels. The economic crisis which started in 2008 in Spain also 519 contributed to the observed trends. Conversely, the contributions from sources mostly linked with natural processes 520 such as Aged Marine (at both BCN and MSY) and Aged Organic (at MSY) did not show statistically significant 521 trends. The general trends observed for the calculated PMF source contributions well reflected the trends observed 522 for the chemical tracers of these pollutant sources. The decrease in the Ammonium sulfate source contribution was 523 mainly attributed to the EC Directive on Large Combustion Plants implemented from 2008 in Spain, resulting in 524 the application of fuels gas desulfurization (FGD) systems in a number of large facilities. Moreover, according to 525 the 2008 Regional AQ Plan, the use of heavy oils and petroleum coke for power generation was forbidden around 526 Barcelona from 2008 in favour of natural gas. As a consequence, a decrease of the contributions from the V-Ni 527 bearing source at both sites was also observed. The decrease observed for the contribution of the Ammonium 528 Nitrate source was mainly due to the reduction in ambient NOx concentrations. In Spain a general decrease of the 529 concentrations of NO2 at regional level was observed and it was mainly related with the lower energy consumption 530 related with the financial crisis. The decrease of nitrates concentrations and Ammonium nitrate source 531 contributions around Barcelona was also attributed to the decrease of NOx emissions from the five power 532 generation plants around the city. Moreover, a Regional AQ Plan implementing the SCRT (continuously 533 regenerating PM traps with selective catalytic reduction for NO2) and the hybridization and shift to natural gas 534 engines of the Barcelona's bus fleet may have had also an influence in NOx ambient concentrations. The 535 Industrial/Metallurgy source contribution at BCN decreased exponentially reflecting the exponential trends 536 observed for the main tracers of this pollutant source (Pb, Cd and As). The implementation of IPPC (Integrated 537 Pollution Prevention and Control) Directives together with a decrease in the emissions from industrial production 538 (smelters) at a regional scale around Barcelona explained the observed trends. Overall, the magnitude of the 539 decreasing trends of the contributions of the pollutant sources were higher at BCN compared to MSY likely because 540 of the proximity of the BCN measurement site to anthropogenic pollutant sources compared to the MSY site. The 541 results presented in this work clearly confirm the beneficial effect of the AQ measures taken in recent years in 542 Europe. However, the WHO limit values of specific pollutants, PM10 and PM2.5 among these, are still exceeded 543 especially at urban level and industrial hotspots. To meet the WHO guide levels important actions are still required 544 for the next decade and the interpretation of past air quality trends may yield relevant outcomes for planning 545 further cost-effective actions. We would like to highlight that a non-linear approach to trend studies is very 546 attractive given that some air pollutants reported in this work showed not gradual-with-time reductions. 547 Conversely, for specific pollutant source-contribution/concentration in our region, the decreasing trend was less 548 steep at the end of the period compared to the beginning thus likely indicating the attainment of a lower limit. This 549 was the case for example for the Secondary sulfate source contribution decreasing exponentially from 2004 to 2014 550 thus likely indicating a limited scope for further reduction of SO2 emissions in our region."

551

#### 552 Minor/technical comments

- 553
- 554 5) Various sections should be called sections, not paragraphs in the text.
- 555
- 556 *Paragraph was replaced with Section throughout the manuscript.*
- 557
- 558 6) Ammonium sulfate and ammonium nitrate sound a bit strange names for sources, as they are 559 compounds that originate from a number of sources as a result of atmospheric processing.
- 560
- The names *Ammonium sulfate* and *Ammonium nit*rate were replaced with *Secondary sulfate* and *Secondary sulfate*
- 563
- 564 7) Line 156: please specify EUSAAR protocol.
- 565

Details on the EUSAAR Protocol can be found in Cavalli, F., Viana, M., Yttri, K. E., Genberg, J., and Putaud, J.-P.: Toward a standardised thermal-optical protocol for measuring atmospheric organic and elemental carbon: the EUSAAR protocol, Atmos. Meas. Tech., 3, 79-89, doi:10.5194/amt-3-79-2010, 2010. This reference was added to the Bibliography. 8) The language of the paper requires some improvements here and there. Some examples:- there is something wrong the sentence on lines 76-78 - line 133: : : :will be discussed later - line 250: : : :(not shown), we concluded: :: - line 270: :: :what was observed: :: - there is something wrong the sentence on lines 280-284. The language of the manuscript was revised Trends analysis of PM source contributions and chemical tracers in NE Spain during 2004 - 2014: A multi-exponential approach. 

598 Marco Pandolfi<sup>1,\*</sup>, Andrés Alastuey<sup>1</sup>, Noemi Pérez<sup>1</sup>, Cristina Reche<sup>1</sup>, Iria Castro<sup>1</sup>, Victor Shatalov<sup>2</sup> and Xavier Querol<sup>1</sup>

- <sup>1</sup> Institute of Environmental Assessment and Water Research, c/ Jordi-Girona 18-26, 08034 Barcelona, Spain
- <sup>2</sup> Meteorological Synthesizing Centre East, 2nd Roshchinsky proezd, 8/5, 115419 Moscow, Russia
- 601 \*Corresponding author: marco.pandolfi@idaea.csic.es
- 602
- 603

#### 604 Abstract

605 In this work for the first time data from two twin stations (Barcelona, urban background, and Montseny, regional 606 background), located in NE of Spain, were used to study the trends of the concentrations of different chemical 607 species in PM<sub>10</sub> and PM<sub>2.5</sub> along with the trends of the PM<sub>10</sub> source contributions from Positive Matrix 608 Factorization (PMF) model. Eleven years of chemical data (2004–2014) were used for this study. Trends of both specie concentrations and source contributions were studied using the Mann-Kendall test for linear trends and a 609 610 new approach based on multi-exponential fit of the data. Despite the fact that different PM fractions (PM<sub>2.5</sub>, PM<sub>10</sub>) 611 showed linear decreasing trends at both stations, the contributions of specific sources of pollutants and the 612 related chemical tracers showed exponential decreasing trends. The different types of trends observed reflected 613 the different effectiveness and/or time of implementation of the measures taken to reduce the concentrations 614 of atmospheric pollutants. Moreover, the trends of the contributions from specific sources such as those related 615 with industrial activities and with primary energy consumption mirrored the effect of the financial crisis in Spain 616 from 2008. The sources that showed statistically significant downward trends at both Barcelona (BCN) and 617 Montseny (MSY) during 2004-2014 were Secondary sulfate, Secondary nitrate, and V-Ni bearing source. The 618 contributions from these sources decreased exponentially during the considered period indicating that the 619 observed decrease was not gradual and consistent over time. Conversely, the decreasing trend was less steep 620 at the end of the period compared to the beginning thus likely indicating the attainment of a lower limit. 621 Moreover, statistically significant decreasing trends were observed for the contributions to PM from the 622 Industrial/Traffic source at MSY (mixed metallurgy and road traffic) and from the Industrial (metallurgy mainly) 623 source at BCN. These sources were clearly linked with anthropogenic activities and the observed decreasing 624 trends confirmed the effectiveness of pollution control measures implemented at EU or regional/local levels. 625 Conversely, the contributions from sources mostly linked with natural processes such as Aged Marine and Aged 626 Organics did not show statistically significant trends. The general trends observed for the calculated PMF source 627 contributions well reflected the trends observed for the chemical tracers of these pollutant sources.

### 628 **1. Introduction**

Meeting the air quality (AQ) standards is one of the major environmental objectives to protect people from breathing air with high levels of pollution. Many studies have been published in these last years showing clearly that the concentrations of particulate matter (PM), and other air pollutants such as sulphur dioxide (SO<sub>2</sub>) and 632 carbon monoxide (CO), have markedly decreased during the last 15 years in many European Countries (EEA, 2013; 633 Barmpadimos et al., 2012; Cusack et al., 2012; Querol et al., 2014; Guerreiro et al., 2014 among others). Cusack 634 et al. (2012) reported the reduction in PM<sub>2.5</sub> concentrations observed at regional background (RB) stations in 635 Spain and across Europe, and, in most cases, the observed reduction was gradual and consistent over time, 636 implying the success of cleaner anthropogenic activities. Barmpadimos et al. (2012) have also shown that  $PM_{10}$ 637 concentrations decreased at a number of urban background (UB) and rural background stations in five European 638 countries. Henschel et al. (2013) reported the dramatic decrease in SO<sub>2</sub> levels across six European cities, reflecting 639 the reduction in sulphur content in fuels, as part of EU legislation, coupled with the shift towards the use of 640 cleaner fuels. EEA (2013) also reported general decreases in NO<sub>2</sub> concentrations even if lower compared to PM. 641 However, Henschel et al. (2015) showed that the NO<sub>x</sub> concentrations at traffic sites in many EU cities remained 642 unchanged underlining the need of further regulative measures to meet the air guality standards for this 643 pollutant. In fact an important proportion of the European population lives in areas exceeding the AQ standards 644 for the annual limit value of NO<sub>2</sub>, the daily limit value of PM<sub>10</sub> and the health protection objective of O<sub>3</sub> (EEA, 645 2013; 2015). PM<sub>10</sub> and NO<sub>2</sub> are still exceeded mostly in urban areas, and especially at traffic sites (Harrison et al., 646 2008; Williams and Carslaw, 2011; EEA, 2013; among others). In Spain for example it has been reported that more 647 than 90% of the NO<sub>2</sub> exceedances are attributed to road traffic emissions (Querol et al., 2012). Guerreiro et al. 648 (2014) furthermore evidenced notable reduction of ambient air concentration of SO<sub>2</sub>, CO and Pb using data 649 available in Airbase (EEA, 2013) and covering 38 European countries. Querol et al. (2014) reported trends for 73 650 measurement sites across Spain including RB, UB, traffic stations (TS) and Industrial sites (IND). They observed 651 marked downward concentration trends for PM<sub>10</sub>, PM<sub>2.5</sub>, CO and SO<sub>2</sub> at most of the RB, UB, TR and IND sites 652 considered. Similarly, Salvador et al. (2012) detected statistically significant downward trends in the 653 concentrations of SO<sub>2</sub>, NO<sub>x</sub>, CO and PM<sub>2.5</sub> at most of the urban and urban-background monitoring sites in the 654 Madrid metropolitan area during 1999-2008. Cusack et al. (2012) and Querol et al. (2014) have also shown the 655 highly statistically significant decreasing trends observed at regional level in NE Spain for many trace elements 656 since 2002 (Pb, Cu, Zn, Mn, Cd, As, Sn, V, Ni, Cr).

657 The observed reduction of air pollutants across Europe is the results of efficient emission abatement strategies 658 as for example those implemented in the Industrial Emission Directives (IPPC Integrated Pollution Prevention and 659 Control and subsequent Industrial Emission Directives 1996/61/EC and 2008/1/EC), the Large Combustion Plants 660 Directive (LCPD: 2001/80/EC), the EURO standards on road traffic emission (1998/69/EC, 2002/80/EC, 661 2007/715/EC), the IMO (International Maritime Organization) directive on sulfur content in fuel and SO<sub>x</sub> and NO<sub>x</sub> 662 emissions from ships (IMO, 2011; Directive 2005/33/EC). Additionally, the financial crisis, causing mainly a 663 reduction of the primary energy consumption from 2008-2009, contributed to the decrease of the ambient 664 concentration of pollutants observed in Spain (Querol et al., 2014).

Moreover, national and regional measures for AQ have been taken in many European Countries. In Spain a national AQ plan was approved in 2011 and updated in 2013 by the Council of Ministers of the Government of

- 667 Spain. Furthermore, 45 regional and 3 local (city scale) AQ plans have been implemented since 2004 in Spain.
- 668 These AQ Plans mostly focused on improving AQ at major city centers or specific industrial areas.
- 669 Thanks to the aforementioned measures, there is clear evidence that the concentrations of PM in many
- 670 European countries have markedly decreased during the last decades. However, in spite of the above policy
- 671 efforts, a significant proportion of the urban population in Europe lives in areas exceeding the World Health
- 672 Organisation (WHO) air quality (AQ) standards i.e. for PM<sub>2.5</sub>, PM<sub>10</sub> and ozone (EEA, 2013, 2015).
- 673 Trend analysis of the concentration of air pollutants helps in evaluating the effectiveness of specific AQ
- 674 measures depending on the pollutant considered. Examining data over time also makes it possible to predict
- 675 **future frequencies and/or rates of occurrence making future projections. For the abovementioned reasons, it**
- 676 is especially attractive the feasibility of studying the trends of the contributions to PM mass from specific
- 677 pollutant sources along with the trends of the chemical tracers of these sources.
- For what we are concerned in the majority of studies dealing with trend analysis, linear fits were applied for example by using Mann-Kendall or Theil-Sen methods (Theil, 1950; Sen, 1968), the latter being available for example in the Openair software (Carslaw, 2012; Carslaw and Ropkins, 2012). However, linear fit of data does not always properly represent the observed trends. As we will show, different abatement strategies and periods of implementation may change from one pollutant to another thus leading to different trends for different pollutants, even over the same period. Thus, non-linear fit of the data may be at times strongly recommended.
- 684The main aim of this work was to study the trends of source contributions to PM10 and specific chemical species685in both PM10 and PM2.5 using both the consensus methodology for linear fit of the data (Mann-Kendall) and a
- 686 non-linear approach. The data of Spanish national emissions and energy consumption are also evaluated to
- 687 interpret the observed trends. Understanding past trends may be relevant for devising new strategies for air
- 688 pollution abatement. PM chemical speciated data collected from 2004 to 2014 at regional (Montseny; NE Spain) 689 and urban (Barcelona, NE Spain) sites were used with this aim. The selected period allowed for trend analysis at 690 these twin stations over a common period. The Positive Matrix Factorization (PMF) model was used to apportion 691 ambient  $PM_{10}$  concentrations into pollutant sources. The PMF model, as other Receptor Models (RM), is widely 692 used being a powerful tool to help policy makers to design more targeted approaches to protecting public health. 693 Thus, the novelty of this study lies mainly in a) the opportunity to study the trends of pollutant source 694 contributions from PMF model at two twin stations representative of the urban and regional environments in the 695 Western Mediterranean, and, b) in the use of a novel non-linear approach for trend studies.
- 696 2. Measurement sites and Methodology

### 697 **2.1 Measurement sites**

The Montseny measurement station (MSY, 41°46′45.63″ N, 02°21′28.92″ E, 720 m a.s.l.) is a regional background
site in NE of Spain (Figure 1). The MSY station is located within a regional natural park about 50 km to the NNE of

the city of Barcelona (BCN) and 25 km from the Mediterranean coast. This site is representative of the typical regional background conditions of the Western Mediterranean Basin (WMB) characterized by severe pollution episodes affecting not only the coastal sites closest to the emission sources, but also the more elevated rural and remote areas land inwards due to thermally driven winds (i.e. Pérez et al., 2008; Pey et al., 2010; Pandolfi et al., 2011; 2014). This station is part of ACTRIS (www.actris.net) and GAW (www.wmo.int/gaw) networks, EMEP (http://www.emep.int/) and the measuring network of the Government of Catalonia.

706 The Barcelona measurement station (BCN, 41°23'24.01" N, 02°06'58.06" E, 68 m a.s.l.) is an urban background 707 measurement site influenced by vehicular emissions from one of the main avenues of the city (Diagonal Avenue) 708 located at a distance of around 300 m (cf. Fig. 1). The BCN measurement site is part of the Air Quality measuring 709 network of the Government of Catalonia. The Metropolitan Area of Barcelona (BMA), with nearly 4.5 million 710 inhabitants, covers an 8 km wide strip between the Mediterranean Sea and the coastal mountain range. Several 711 industrial zones, power plants, and highways are located in the area, making this region to one of the most 712 polluted in the WMB (i.e. Querol et al., 2008; Amato et al., 2009; Pandolfi et al., 2012; 2013; 2014). At BCN the 713 location of the measuring station changed in 2009 when it was moved by around 500 m (cf. Fig. 1). The effect of 714 this change on PM measurements performed at BCN will discuss later.

715

#### 716 2.2 Real-time and gravimetric PM measurements

Real-time PM concentrations were continuously measured at 1h resolution by optical particle counters (OPC)
using GRIMM spectrometers (GRIMM 180 at MSY, and GRIMM 1107, 1129 and 180 at BCN). Hourly PM
concentrations were corrected by comparison with 24h gravimetric mass measurements of PM<sub>x</sub> (Alastuey et al.,
2011).

For gravimetric measurements 24h PMx samples were collected at both stations every 3-4 days on 150 mm quartz micro-fiber filters (Pallflex QAT and Whatman) with a high-volume (Hi-Vol) samplers (DIGITEL DH80 and/or MCV CAV-A/MSb at 30 m<sup>3</sup>h<sup>-1</sup>). The mass of PM<sub>10</sub> and PM<sub>2.5</sub> samples collected on filters was determined using the EN 12341 and the EN14907 gravimetric procedures, respectively.

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### 728 2.2.1 PM chemical speciated data

Once the gravimetric mass was determined from filters, the samples were analyzed with different techniques including acidic digestion (½ of each filter; HNO<sub>3</sub>:HF:HCIO<sub>4</sub>), water extraction of soluble anions (¼ of each filter), and thermal-optical analysis (1.5 cm<sup>2</sup> sections). Inductively Coupled Atomic Emission Spectrometry, ICP-AES, (IRIS Advantage TJA Solutions, THERMO) was used for the determination of the major elements (AI, Ca, Fe, K, Na, Mg, S, Ti, P), and 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). Ionic Chromatography was used for the concentrations of  $NO_3^-$ ,  $SO_4^{2-}$  and  $CI^-$ , whereas  $NH_4^+$  was determined using a specific electrode MODEL 710 A+, THERMO Orion. The levels of OC and EC were determined by a thermal-optical carbon analyzer (SUNSET), using protocol EUSAAR\_2 (Cavalli et al., 2010). Other analytical details may be found in Querol et al. (2008).

Following the above procedures, PM<sub>10</sub> and PM<sub>2.5</sub> chemical speciated data were obtained at MSY for the period
2004-2014 resulting in 1093 and 794 samples, respectively. At BCN PM<sub>10</sub> and PM<sub>2.5</sub> data were obtained during
2004-2014 resulting in 1037 and 1063 samples, respectively.

741

#### 742 **2.3 Positive Matrix Factorization (PMF) model.**

743 The PMF model (PMFv5.0, EPA) was used on the collected daily speciated data for source identification and 744 apportionment in PM<sub>10</sub> at both sites. Detailed information about the PMF model can be found in literature 745 (Paatero and Tapper 1994; Paatero 1997; Paatero and Hopke 2003; Paatero et al. 2005). The PMF model is a 746 factor analytical tool reducing the dimension of the input matrix in a limited number of factors (or sources) and 747 it is based on the weighted least-squares method. Thus, most important in PMF applications is the estimation of 748 uncertainties of the chemical species included in the input matrix. In the present study, individual uncertainties 749 and detection limits were calculated as in Escript et al. (2009) and Amato et al. (2009). Thus, both the analytical 750 uncertainties and the standard deviations of species concentrations in the blank filters were considered in the 751 uncertainties calculations. The signal-to-noise ratio (S/N) was estimated starting from the calculated uncertainties 752 and used as a criteria (S/N >2) for selecting the species used within the PMF model. In order to avoid any bias in 753 the PMF results, the data matrix was uncensored (Paatero 2004). The PMF was run in robust mode (Paatero 754 1997), and rotational ambiguity was handled by means of the F<sub>PEAK</sub> parameter (Paatero et al. 2005). The optimal 755 number of sources was selected by inspecting the variation of the objective function Q (defined as the ratio 756 between residuals and errors in each data value) with varying number of sources (i.e. Paatero et al., 2002) and by 757 studying the physical meaningfulness of the calculated factors.

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- 759

#### 760 2.4 Mann-Kendall (MK) fit

The purpose of the Mann-Kendall (MK) test (Mann 1945, Kendall 1975, Gilbert 1987) is to statistically assess if there is a monotonic upward or downward trend of the variable of interest over time. A monotonic upward (downward) trend means that the variable consistently increases (decreases) through time. The Mann-Kendall test tests the null hypothesis H<sub>0</sub> of no trend, i.e. the observations are randomly ordered in time, against the

765	alternative hypothesis, H1, where there is an increasing or decreasing monotonic trend. The main advantage of
766	the Mann-Kendall test is that data need not conform to any particular distribution and missing data are allowed
767	To estimate the slope of the trend the Sen's method was used (Salmi et al. 2002).

- 768
- 769 2.5 Multi-exponential (ME) fit

770 A Program aiming at studying trends of time series of air pollution in the multi-exponential form was developed 771 within the The Task Force on Measurements and Modelling (TFMM) by the Meteorological Synthesizing Centre - East (MSC-E; http://www.msceast.org/) group (Shatalov et al., 2015). The TFMM together with the Task Force 772 773 on Emission Inventories and Projections (TFEIP), the Task Force on Integrated Assessment Modelling (TFIAM), 774 and Task Force on Hemispheric Transport of Air Pollution (TFHTAP) provide a fora for discussion and scientific 775 exchange in support of the EMEP (European Monitoring and Evaluation Programme; http://www.emep.int/) 776 work plan which is a scientifically based and policy driven programme under the Convention on Long-range Transboundary Air Pollution (CLRTAP; http://www.unece.org/env/lrtap/lrtap\_h1.html) promoting the 777 778 international co-operation to solve transboundary air pollution problems. The TFMM was established in 2000 779 to evaluate measurements and modeling and to further develop working methods and tools. In this contest, 780 five EMEP Centers are undertaking efforts in support of the EMEP work plan, namely the MSC-E, the Centre on 781 Emission Inventories and Projections (CEIP; http://www.ceip.at/), the Chemical Coordinating Centre (CCC; 782 http://www.nilu.no/projects/ccc/), the Meteorological Synthesizing Centre – West (MSC-W; 783 http://emep.int/mscw/index\_mscw.html), and the Centre for Integrated Assessment Modelling (CIAM; http://www.iiasa.ac.at/~rains/ciam.html). In 2014, the TFMM initiated a dedicated exercise to assess the 784 785 efficiency of air pollution mitigation strategies over the past 20 years to assess the benefit of the CLRTAP main 786 policy instrument. Within this exercise a software was made available by EMEP/MSC-E Center aiming at 787 studying non-linear trends. Annual, monthly and daily resolution data can be analyzed with the help of this 788 program. Since in this paper we will apply the program to annual averages of specie concentrations and source 789 contributions, we restrict the description of the multi-exponential approximations for this case. In particular, 790 seasonal variations are not included into consideration. The basic equations solved by the program for this 791 particular case (annual averages) are reported below:

792

793 
$$C_t = a_1 \cdot \exp\left(-\frac{t}{\tau_1}\right) + a_2 \cdot \exp\left(-\frac{t}{\tau_2}\right) + \dots + a_n \cdot \exp\left(-\frac{t}{\tau_n}\right) + \omega_t$$
(1)

794

Where,  $C_t$  are the values of the considered time series, with t = 1, ..., N, N being the length of the series (years),  $\tau_n$  are the characteristic times of the considered exponential,  $a_n$  are constants and  $\omega_t$  are the residue values. In the case of single exponential decay (n=1) the characteristic time  $\tau$  is the time at which the pollutant

798	concentration is reduced to 1/e (= 0.3678) times its initial value. The main difference between linear and
799	exponential fit is that in the latter case the trend is not gradual and constant over time. For an exponential trend
800	the absolute $[\mu g/m^3]$ reduction per year decreases with time being the highest at the beginning of the period.
801	Conversely, for a linear fit the absolute reduction is constant over time. For an exponential fit, the lower the
802	characteristic time $ au$ the more rapidly the considered quantity vanish. Deviations from single exponential fit can
803	be taken into account introducing more exponential terms. In this work for example two exponential terms were
804	sometime used. In this case, two characteristic times are calculated by the software. If the decrease of the
805	considered quantity is very sharp at the beginning of the period (more than exponential) than both $ au_1$ and $ au_2$ are
806	positive. Conversely, an exponential term with negative $ au$ takes into account for possible increases of the quantity
807	at the end of the period. Both $\tau_n$ and $a_n$ are calculated by the program by means of the least square method
808	minimizing the residue $\omega$ and the statistically significance of the exponential fit is provided by means of the p-
809	value. The number of exponential terms that should be included into the approximation can be evaluated using
810	F-statistics (i.e. Smith, 2002). For example, the F-statistics for the evaluation of the statistical significance of the
811	second term in equation (1) for n = 2 can be calculated as:

813 
$$F = \frac{(SS_1 - SS_2)}{2 \cdot S}$$
 (2)

815 where SS<sub>1</sub> and SS<sub>2</sub> are sums of squares of residual component for approximations with one and two exponential 816 terms, respectively, and s is the estimate of standard deviation of residual component. This statistics follows 817 approximately the Fisher distribution with 2 and N – 2 degrees of freedom. Second exponential is considered to 818 be significant if *F* exceeds the corresponding threshold value at the chosen significance level.

819 The following parameters can be calculated from equation (1):

820 - Total Reduction (TR): 
$$TR = \frac{(C_{beg} - C_{end})}{C_{beg}} = 1 - \frac{C_{end}}{C_{beg}}$$
 (3)

821 - Annual reduction for year *i*: 
$$R_i = \frac{\Delta C_i}{C_i} = 1 - \frac{C_{i+1}}{C_i}$$
 (4)  
822 Average appual reduction:  $R_i = 1 - \frac{\left(C_{end}\right)^{\frac{1}{N-1}}}{C_i}$  (5)

Where 
$$C_{beg}$$
 and  $C_{end}$  are respectively the first and the last points of the exponential fit. The formula for calculation  
of average annual reduction takes into account that the ratio  $C_{i+1} / C_i$  is a multiplicative quantity, so that  
geometrical mean of ratios should be used. The relative contribution of residues (Residual Component: RC) is

calculated as the standard deviation of the ratios between the residue values of the fit  $\omega$  (cf. Eq. 1) and the main 826 827 component of the fit.

so that

828 The MSC-E also proposed a statistic which measures the deviation of the obtained trend from the linear one (Non-829 Linearity parameter: NL). A trend is defined as linear if the NL parameter is lower than 10%, indicating a small 830 difference between ME and MK fits (Shatalov et al., 2015). In the following, the reported trends were analysed

using the MK test for NL<10% and the ME test for NL>10%. More detailed description of the multi-exponential
 approach is available in the TFMM wiki and in the MSC-E Technical report 2015 (Shatalov et al., 2015).

833

#### 834 3. Results

835 Results are presented and discussed in the following order: In Section 3.1, we compare the trends at both stations 836 of PM<sub>x</sub> concentrations from optical counters (OPC; annual data coverage around 90%) and from 24h gravimetric 837 samples (filters; annual data coverage around 20-30%). This comparison will demonstrate the feasibility of 838 studying trends of chemical species concentrations from filters despite the relatively low annual data coverage. 839 In Section 3.2, we compare the magnitude of the trend of PM<sub>2.5</sub> concentrations at MSY during 2004-2014 (period 840 selected for this study) with the magnitude of trends calculated at the same station over different periods, namely 2002-2010 (the period used in Cusack et al., 2012) and 2002-2014 (representing the largest period of gravimetric 841 PM<sub>2.5</sub> measurements available at the time of writing at MSY station). This comparison was performed in order to 842 843 study the differences in the trends over short periods (9 yr to 13 yr). The gravimetric concentrations of PM<sub>2.5</sub> 844 measured at MSY were used with this aim. Then (Section 3.3), we present and discuss the trends at both stations 845 of chemical species in both PM<sub>10</sub> and PM<sub>2.5</sub> from 24h filter analyses. In the Section 4.0 we discuss the sources of 846 pollutants identified by PMF model in PM<sub>10</sub> at both sites. Finally, we present and discuss the trends of PM<sub>10</sub> source 847 contributions at BCN and MSY (Section 4.1) providing possible explanations for the observed trends. Some 848 conclusions are reported in Section 5.

849

## 850 **3.1 Trends of PM: Comparison between gravimetric and real-time optical measurements**

851	Annual data coverage is an important factor to take into account in order to study trends of a given parameter.
852	The gravimetric PM measurements, from which chemical speciated data are obtained, are typically performed
853	with rather low frequency over one year. In our case the annual data coverage of gravimetric measurements was
854	around 20-30% at both Barcelona and Montseny. In this section we compare the trends of PM concentrations
855	from gravimetric and real-time optical measurements (Table 1). Given that the trends of the considered $PM_x$
856	fractions were linear at both sites (NL<10%), only results from MK test were reported in Table 1. However, we
857	will show later ( <i>Section 4.1</i> ) that the contributions from specific $PM_{10}$ pollutant sources from PMF model, mainly
858	those related with anthropogenic activities, showed non-linear (i.e. exponential) decreasing trends, thus
859	mirroring the different effectiveness of the mitigation strategies depending on the source of pollutants
860	considered.

As reported in Table 1, statistically significant decreasing trends were observed for the considered PM size fractions at BCN (-2.20  $\mu$ gm<sup>-3</sup>/yr with p<0.001 for PM<sub>10</sub> and -1.55  $\mu$ gm<sup>-3</sup>/yr with p<0.01 for PM<sub>2.5</sub> from OPC 863 measurements), whereas at Montseny only the PM<sub>2.5</sub> fraction showed a little significant decreasing trend (-0.26 864  $\mu$ gm<sup>-3</sup>/yr; p<0.1 from OPC measurements). Total reductions (TR) ranged between 50.4% (OPC PM<sub>10</sub> at BCN) to 865 7.8% (OPC PM<sub>10</sub> at MSY) and residual component (RC) was lower than 18% reflecting the goodness of the linear 866 (MK) fit used. It must be noted that the higher p-values, magnitude of the trends and TR observed at BCN 867 compared to MSY was likely due to the change of the measuring station in 2009 in BCN (cf. Fig. 1). Based on the 868 comparison between simultaneous PM<sub>x</sub> chemical speciated data collected at both BCN measurement sites during 869 1 month (not shown) we concluded that after 2009 the BCN measuring site was less affected by mineral matter 870 and, to a lesser extent, by road traffic emissions both being important sources of PM in Barcelona. In Figure 1 we 871 highlighted the proximity of the BCN measuring station before 2009 to an unpaved parking and different 872 construction works. The effect of the change of the station in BCN in 2009 on PM<sub>10</sub> gravimetric measurements 873 was reported in Supporting Information (Figure SI-1). However, despite the change of the station, the comparison 874 between BCN and MSY for specific chemical species and pollutant sources not linked with mineral matter and 875 road traffic emissions was possible.

876 Table 1 shows that the p-values calculated using gravimetric and OPC measurements was the same despite the 877 different annual data coverage. The differences in the magnitude of the trends were 22% and 24% between 878 gravimetric and OPC PM<sub>10</sub> and PM<sub>2.5</sub> measurements, respectively, at BCN and 24% and 21%, respectively, at 879 MSY. Relative differences of total reductions (TR) ranged between 24% between gravimetric and OPC PM<sub>10</sub> 880 measurements at MSY and 15% for PM<sub>10</sub> at BCN. Thus, despite the different data coverage the magnitude of the trends and TR calculated from OPC and gravimetric measurements were rather similar. Other PM mass 881 882 fractions (PM<sub>1-10</sub> and PM<sub>2.5-10</sub>) and PM ratios (PM<sub>1</sub>/PM<sub>10</sub> and PM<sub>2.5</sub>/PM<sub>10</sub>) at MSY showed non-statistically 883 significant trends.

884

#### 885 3.2 Trends of PM: Comparison among different periods

886 In this study we used the period 2004-2014 for trends analysis given that gravimetric PM<sub>2.5</sub> measurements at BCN 887 were available since 2004. Conversely, at MSY PM<sub>2.5</sub> gravimetric measurements started in 2002. Figure 2 shows 888 the trends of PM<sub>2.5</sub> concentrations at MSY calculated using the MK test for the three different periods. ME test 889 was not used here given that the observed trends were linear (NL<10%). The period 2002-2010 was the period 890 considered in the paper from Cusack et al. (2012) presenting the trends of PM<sub>2.5</sub> gravimetric mass and chemical 891 species at MSY. The period 2002-2014 is the largest period with  $PM_{2.5}$  filter measurements available at the time 892 of writing. The trend observed at MSY for the PM<sub>2.5</sub> fraction during 2004-2014 confirmed what already observed 893 by Cusack et al. (2012) at the same station for the period 2002 – 2010. In Cusack et al. (2012) the MK test provided 894 a decreasing trend of around -0.66 µgm<sup>-3</sup>/yr at 0.01 significance level (TR = 35%). During the periods 2004 – 2014 895 and 2002 - 2014 decreasing trends of -0.33  $\mu$ gm<sup>-3</sup>/yr (p<0.1; TR = 26%) and -0.37  $\mu$ gm<sup>-3</sup>/yr (p<0.05; TR = 31%), 896 respectively, were observed. Thus, a statistically significant trend for PM<sub>2.5</sub> mass at regional level can be confirmed 897 even considering different periods thus confirming the effectiveness of mitigation measures together with the

898	effect of the economic crisis in Spain from 2008. However, it should be noted that the statistical significance of
899	the trends observed for the larger periods was lower compared to Cusack et al. (2012). The difference observed
900	in the magnitude of the trends during 2004-2014 compared to the results provided by Cusack et al. (2012) was
901	mainly due to the increase of PM <sub>2.5</sub> mass concentration in 2012 (cf. Figure 2). Chemical PM <sub>2.5</sub> speciated data
902	revealed that this increase was partly driven by organic matter showing a mean annual concentration in 2012
903	higher by around 20% compared to the 2004-2014 average.

#### 905 3.3 Trends of chemical species

906 The trends of the **annual mean** concentrations of chemical species at BCN and MSY are reported in Table 2 (for 907

908 PM<sub>10</sub>. In Tables 2 and 3 and Figures 3 and 4 only the species having statistically significant trends were reported.

PM<sub>10</sub>) and Table 3 (for PM<sub>2.5</sub>). Figure 3 (for BCN) and Figure 4 (for MSY) show the trends of chemical species in

909 As already noted, we assume that the change of the station in BCN in 2009 affected the trends of the 910 concentrations of **OC, EC,** Cu, Sn, Sb and Zn (mainly traffic tracers), Al<sub>2</sub>O<sub>3</sub>, Ca, Mg, Ti, Rb, Sr (crustal elements 911 related with both natural and anthropogenic sources) and Fe (traffic and crustal tracer). These chemical species 912 at BCN were removed from Tables 2 and 3 and from Figure 3.

Other species with less local character measured in BCN were instead included in the analysis. These are SO<sub>4</sub><sup>2-</sup>, 913 914  $NH_{4^{+}}$ , V, Ni (related with heavy oil combustion in the study area according to source apportionment results, cf. 915 Par. 4), Pb, Cd, and As (related with industrial/metallurgy activities), Na and Cl (sea spray), and  $NO_3^-$ . Although nitrate particles in Barcelona were mainly from traffic, the concentrations of these particles were not strongly 916 917 affected by the change of the station due to their secondary origin. The MSY station will be considered as 918 reference station given that no location change occurred at this monitoring site during the study period.

919 Statistically significant exponential trends (p < 0.01 or 0.001) were mainly observed for the industrial tracers 920 (Pb, Cd, As) in both PM<sub>10</sub> and PM<sub>2.5</sub>. For these elements TR was high and around 50-80% in PM<sub>10</sub> and 67-81% in 921 PM<sub>2.5</sub>. The RCs were lower than 20% thus suggesting the goodness of the exponential fits used to study the 922 trends of these species. Exponential fits were on average needed indicating that the trends were not gradual and consistent over time and that the effectiveness of the control measures for these pollutants was stronger 923 924 at the beginning of the period under study (2004-2009 approximately) compared to the end of the period (Figs. 3 and 4). This is also evident by comparing the linear MK fit (dashed black line) with the ME fit (red line) in Figs. 925 926 3 and 4. In PM<sub>10</sub> the magnitudes of the trends ranged between -0.00222  $\mu$ gm<sup>-3</sup>/yr (Pb; p<0.001) to -3.10E-5 927 μgm<sup>-3</sup>/yr (Cd; p<0.001) at BCN and from -0.00031 μgm<sup>-3</sup>/yr (Pb; p<0.01) to -1.12E-5 μgm<sup>-3</sup>/yr (Cd; p<0.01) at 928 MSY. In PM<sub>2.5</sub> the magnitude of the trends were similar and ranged between -0.00163  $\mu$ gm<sup>-3</sup>/yr (Pb; p<0.001) 929 and -3.11E-5 μgm<sup>-3</sup>/yr (Cd; p<0.001) at BCN and between -0.00049 μgm<sup>-3</sup>/yr (Pb; p<0.001) and -1.35E-5 μgm<sup>-1</sup> 930 <sup>3</sup>/yr (Cd; p<0.001) at MSY. Similar magnitude of the trends for these species in both PM fractions at both sites 931 confirmed the common origin of these elements and the impact at regional scale of industrial sources. For Pb and

Cd the characteristic time ( $\tau$ ) of the exponential trends was similar at both sites, whereas for As it was higher due to the slightly less intense exponential downward trend observed for As compared to Cd and Pb. Note that the PMF analysis **(cf. Section 4)** revealed that the concentrations of As were explained by multiple sources (especially at BCN) whereas the *Industrial/metallurgy* source alone explained more than around 70% of Pb and Cd concentrations (not shown). The implementation of the IPPC Directive in 2008 in Spain is the most probable cause for this downward trend. The decrease observed for Pb, Cd and As may be also attributed to a decrease in the emissions from industrial production (smelters, Querol et al., 2007) at a regional scale around Barcelona.

939 The concentrations of V and Ni in Barcelona in both PM<sub>10</sub> and PM<sub>2.5</sub> fractions showed very similar exponential 940 decreasing trends. Similar characteristic times (around 10-11 yr), TR (around 59-63%) and RC (15-17%) in both 941 fractions suggested the common and mainly fine origin of these two elements. At MSY, V and Ni showed linear 942 trends likely because of the higher distance of the MSY station to the sources of V and Ni (shipping and, before 943 2008, energy production) compared to BCN. Note also that the NL parameter for BCN V and Ni was around 10-944 12%, indicating that in this case the exponential fit did not differ very much from the linear one. Total reduction 945 for V and Ni at MSY was around 59-64% and 42-43%, respectively, and RCs were lower than 24%.

946 Sn and Cu in PM<sub>10</sub> at MSY showed very similar behavior decreasing linearly with time with TR around 36-39% and 947 RC around 16-20%. Decreasing rates of -3.65E-5  $\mu$ gm<sup>-3</sup>/yr (p<0.05) and -0.00014  $\mu$ gm<sup>-3</sup>/yr (p<0.05) were observed 948 in PM<sub>10</sub> for Sn and Cu, respectively. In PM<sub>2.5</sub>, the concentrations of Sn and Cu decreased markedly compared to 949  $PM_{10}$  at the rate of -0.00084  $\mu$ gm<sup>-3</sup>/yr (p<0.001) and -0.00026  $\mu$ gm<sup>-3</sup>/yr (p<0.01), respectively. This difference 950 could be explained by possible sources of coarser Sn and Cu which reduced the magnitude of the trends in PM<sub>10</sub> 951 mass fraction. Sb showed marked decreasing trends in both PM mass fractions compared to Sn and Cu with TR 952 around 62-70%. The magnitude of the trends for Sb were similar in both fractions and around -3.57 ÷ -3.86E-5 953  $\mu$ gm<sup>-3</sup>/yr. Sb concentrations were better fitted with exponential curves (SE with p<0.01 in PM<sub>10</sub> and DE with 954 p<0.01 in PM<sub>2.5</sub>). The DE fit for Sb in PM<sub>2.5</sub> had one positive and one negative characteristic time, the latter 955 needed to explain the slight increase in Sb concentrations at the end of the considered period. The marked 956 decreasing trend observed for Sb compared to other traffic tracers could be explained by a progressive reduction 957 of Sb contained in the vehicle brakes. Cr did not show a statistically significant trend in both PM fractions.

958 Sulfate (SO<sub>4</sub><sup>2</sup>) and ammonium (NH<sub>4</sub><sup>+</sup>) particles concentrations showed very similar behavior in PM<sub>2.5</sub> and PM<sub>10</sub> 959 size fractions due to their fine nature. In BCN the magnitude of the trends were -0.37868  $\mu$ gm<sup>-3</sup>/yr (p<0.001) and 960 -0.11095  $\mu$ gm<sup>-3</sup>/yr (p<0.001) for SO<sub>4</sub><sup>2-</sup> and NH<sub>4</sub><sup>+</sup>, respectively, in PM<sub>10</sub> and -0.32778  $\mu$ gm<sup>-3</sup>/yr (p<0.001) and -961 0.12701 µgm<sup>-3</sup>/yr (p<0.001), respectively, in PM<sub>2.5</sub>. The trends were SE with very similar characteristic times (9.64-962 9.81 yr in PM<sub>10</sub> and 9.69-10.53 yr in PM<sub>2.5</sub>), TR (64-65% in PM<sub>10</sub> and 61-64% in PM<sub>2.5</sub>) and RC (12-14% in PM<sub>10</sub> and 963 9-15% in PM<sub>2.5</sub>). At MSY the magnitude of the trends of SO<sub>4</sub><sup>2-</sup> and NH<sub>4</sub><sup>+</sup> and their statistically significance were 964 lower compared to BCN in both fractions. Moreover, at MSY the trends were linear for SO<sub>4</sub><sup>2-</sup> in both fractions (as 965 for V and Ni). These differences could be explained by the distance of MSY to direct specific sources of sulfate, 966 such as shipping, compared to BCN, thus slightly reducing the magnitude and the statistically significance of the

trend of  $SO_4^{2^{-}}$  at regional level. It is also interesting to note the similitude between the characteristic times of the exponential fits for V and Ni and  $SO_4^{2^{-}}$  in both PM fractions at BCN suggesting the main common origin of these chemical species. Possible reasons for the observed reduction in the concentrations of ambient sulfate in and around Barcelona will be discussed later.

Fine NO<sub>3</sub><sup>-</sup> (Table 3) showed statistically significant SE trends similar at both sites with p<0.001, TR around 73-82%, RC around 16-21% and characteristic times around 5.8-7.6 yr. In PM<sub>10</sub> the TR were lower and around 54-64% and the fits were linear at MSY and SE at BCN. The SE fit at BCN in PM<sub>10</sub> provided a characteristic time around 9.8 yr, higher compared to  $\tau$  obtained for the fine mode because fine NO<sub>3</sub><sup>-</sup> had a more pronounced downward trend compared to PM<sub>10</sub> NO<sub>3</sub><sup>-</sup>.

976 For the mineral species (Al<sub>2</sub>O<sub>3</sub>, Ca, Fe) linear (with the exception of Al<sub>2</sub>O<sub>3</sub> in PM<sub>2.5</sub> which was SE) and statistically 977 significant decreasing trends were detected at MSY. On average the TR was higher in the fine fraction, ranging 978 from 50% for Ca to 66% for  $Al_2O_3$ , compared to  $PM_{10}$  (6-38% cf. Table 2). Downward decreasing trend for crustal 979 material in PM<sub>2.5</sub> at MSY was also reported by Cusack et al. (2012) for the period 2002 – 2010 and by Querol et 980 al. (2014) for the period 2001 – 2012. These trends were probably driven by weather conditions associated with 981 negative NAO index (iNAO) that could be the cause for this slight reduction observed in crustal material. Pey et 982 al. (2013) found a correlation between iNAO (calculated between June and September) and the contribution of 983 Saharan dust to PM<sub>10</sub> mass in NE of Spain showing that the more negative is the iNAO the lower is the dust 984 contribution to PM. The iNAO was unusually negative during the period 2008 - 2012 985 (http://www.cpc.ncep.noaa.gov/products/precip/CWlink/pna/norm.nao.monthly.b5001.current.ascii) thus 986 likely contributing to explain the observed trends of crustal elements. Moreover, negative NAO can favour the 987 presence of fronts that can sweep the Iberian Peninsula from West to East causing higher wind and less stagnant 988 conditions thus favouring the dispersion of pollutants. In addition, as suggested by Cusack et al (2012), it could 989 also be hypothesised that some part of the crustal material measured at MSY is a product of the construction 990 industry. The construction industry in Spain has been especially affected by the current economic recession, and 991 crustal material produced by this industry may have contributed to the crustal load in PM<sub>2.5</sub>. For example, the 992 number of home construction works in Barcelona during 2008 – 2014 (from the beginning of the economic crisis; 993 mean number of works = 1281) reduced by around 75% compared to the period 2000 – 2007; mean number of 994 works = 5187) (http://www.bcn.cat/estadistica/castella/dades/timm/construccio/index.htm). The fact that the 995 total reduction calculated for mineral elements reported in Tables 2 and 3 was higher in PM<sub>2.5</sub> compared to  $PM_{10}$ 996 could corroborate this latter hypothesis.

Finally, Na concentrations showed linear decreasing trends at both sites, with the exception of  $PM_{10}$  Na at MSY. Other species at MSY such as OC and EC did not show statistically significant trends. Consider that the concentrations of EC at MSY are very low and around at 0.2-0.3  $\mu$ g/m<sup>3</sup> as annual mean. Both anthropogenic activity and biomass burning were expected to contribute to this chemical specie. Concerning OC the lack of trend was probably due to the contribution from biogenic sources to the concentration of this specie at regional level.

# 1003 4. PMF source profiles and contributions

Eight and seven sources were detected at BCN and MSY, respectively, in PM<sub>10</sub> from PMF model. The absolute and relative contributions of these sources to the measured PM<sub>10</sub> mass are reported in Figure 5. The chemical profiles of the detected sources were reported in Supporting Information (Figure SI-2).

1007 Some of these sources were common at both BCN and MSY. These are: Secondary Sulfate (secondary inorganic 1008 source traced by  $SO_4^2$  and  $NH_4^+$  and contributing 3.95 µg/m<sup>3</sup> (23.7%) and 4.67 µg/m<sup>3</sup> (13.7%) at MSY and BCN, 1009 respectively), Secondary nitrate (secondary inorganic source traced by NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup> and contributing 1.31  $\mu$ g/m<sup>3</sup> 1010 (7.9%) and 4.45 µg/m<sup>3</sup> (13.1%) at MSY and BCN, respectively), V-Ni bearing source (traced mainly by V, Ni and 1011  $SO_4^{2-}$  it represents the direct emissions from heavy oil combustion and contributed 0.71  $\mu$ g/m<sup>3</sup> (4.3%) and 3.32 1012 µg/m<sup>3</sup> (9.8%) at MSY and BCN, respectively), *Mineral* (traced by typical crustal elements such as AI, Ca, Ti, Rb, Sr 1013 and contributing 2.70 µg/m<sup>3</sup> (16.2%) and 4.61 µg/m<sup>3</sup> (13.6%) at MSY and BCN, respectively), Aged marine (traced 1014 by Na and CI mainly with contributions from  $SO_4^{2}$  and  $NO_3^{-1}$  and contributing 1.76  $\mu$ g/m<sup>3</sup> (10.6%) and 5.73  $\mu$ g/m<sup>3</sup> 1015 (16.9%) at MSY and BCN, respectively). Sources detected at MSY but not at BCN were: Industrial/Traffic source 1016 (traced by EC, OC, Cr, Cu, Zn, As, Cd, Sn, Sb and Pb it includes mixed contributions from anthropogenic sources 1017 such as road traffic and metallurgic industries and contributed 1.43 µg/m<sup>3</sup> (8.6%)) and Aged organics (traced 1018 mainly by OC and EC with maxima in summer indicating mainly a biogenic origin and contributing 3.78  $\mu$ g/m<sup>3</sup> 1019 (22.7%)). The ratio OC:EC in the Industrial/Traffic and Aged organic source profiles at MSY were 4.2 and 11.7, 1020 respectively, thus indicating a strong influence of aged particles in the latter source with the former source being 1021 more fresh. The statistic of the OC:EC ratio based on chemical data at MSY is reported in Supporting Information 1022 (Figure SI-3). Mean and median values of OC:EC ratio at MSY were 9.1 and 7.8, respectively.

Finally, some sources were detected at BCN but not at MSY: *traffic* (traced by  $C_{nm}$ , Cr, Cu, Sb and Fe mainly and contributing 5.14 µg/m<sup>3</sup> (15.1%)), *road/work resuspension* (traced by both crustal elements, mainly Ca, and traffic tracers such as Sb, Cu and Sn and contributing 4.25 µg/m<sup>3</sup> (12.5%)) and *Industrial/metallurgy* (traced by Pb, Cd, As and Zn and contributing 0.96 µg/m<sup>3</sup> (2.8%).

1027 A sensitivity study was performed in order to better interpret the PMF sources at BCN. In fact, for the period 2007 1028 - 2014 separate OC and EC concentration measurements were available and a PMF was performed. The 1029 comparison between the PMF source contributions obtained using the period 2007-2014 (separate OC and EC 1030 measurements) and the whole period (2004-2014; Cnm (non-mineral carbon) available) is reported in Supporting 1031 Information (Figure SI-4). As reported in Figure SI-4 the differences in source contribution and R<sup>2</sup> ranged between 1032 -3% (Mineral source) and +20% (Industrial source) and 0.894 to 0.997, respectively, thus confirming the correct 1033 interpretation of the 2004-2014 PMF sources where C<sub>nm</sub> was used. The OC:EC ratio in the Traffic source from 1034 2007-2014 PMF was 1.70 (cf. Figure SI-5) whereas the mean and median OC:EC ratio from chemistry data were 1035 2.5 and 2.3, respectively, thus being in agreement with the contribution of fresh particles from *Traffic* source at

1036 BCN.

#### 1037

#### 1038 **4.1 Trends of annual PM**<sub>10</sub> source contributions

1039 Figures 6 and 7 and Table 4 show the results from MK or ME test applied to the annual averages of PM<sub>10</sub> source 1040 contributions at BCN and MSY. As already noted we cannot study trends for Traffic, Road/work resuspension and 1041 *Mineral* source contributions at BCN because of the change of the station location in 2009. The contributions that 1042 showed statistically significant downward trends at both stations were from **Secondary** sulfate, **Secondary** 1043 *nitrate*, and *V-Ni bearing* sources (p<0.001 or p<0.01). Moreover, statistically significant decreasing trends were observed for the Industrial/Traffic (p<0.01) and Mineral (p<0.1) source contributions at MSY and the 1044 1045 Industrial/metallurgy source (p<0.001) at BCN. These sources were mostly linked with anthropogenic activities 1046 and the observed decreasing trends confirmed the effectiveness of pollution control measures together with the possible effect of the economic crisis in Spain from 2008. Conversely, the contributions from sources mostly 1047 1048 linked with natural processes such as Aged Marine (at both BCN and MSY) and Aged Organic (at MSY) did not 1049 show statistically significant trends.

1050 The trends of the **Secondary** sulfate source contributions were DE and SE at BCN and MSY, respectively, thus the 1051 decrease over time of this source contribution was not gradual and monotonic. Overall the observed decreasing 1052 trends at both stations may be attributed to the legislation that came into force in 2007-2008 in Spain, the EC 1053 Directive on Large Combustion Plants, which resulted in the application of flue gas desulfurization (FGD) systems 1054 in a number of large facilities in 2007-2008 in Spain. Figure 8 shows the sharp decreases after 2007 observed for 1055 the national SO<sub>2</sub> and NO<sub>x</sub> emissions mostly from power generation (MAGRAMA, 2013; Querol et al., 2014). In BCN 1056 the two characteristic times (one low and the other high, cf. Table 4) of the DE fit indicated a strong decrease of 1057 the **Secondary** sulfate source contribution at the beginning of the period. Moreover, this decrease was sharper 1058 compared to MSY where SE fit was used. This difference was mostly due to the ban of heavy oils and petroleum 1059 coke for power generation around Barcelona from 2007. The effects of this AQ Regional Plan were likely more 1060 effective in BCN compared to MSY thus explaining the two different exponential fits used. Overall, for the 1061 Secondary sulfate source contributions the TRs were rather high around 53% at MSY and 67% at BCN with RC 1062 ranging from 16% (BCN) to 21% (MSY). The fact that the trend of the Secondary sulfate source contribution was 1063 exponential likely suggested the attainment of a lower limit and indicated a limited scope for further reduction 1064 of SO<sub>2</sub> emissions in our region. In fact, it has been estimated that the maximum in EU will be a further 20% 1065 reduction through measures in industry, residential and commercial heating and reduced agricultural waste 1066 burning (UNECE, 2016). Conversely, in Eastern European countries the scope for reduction is much greater and 1067 around 60% (UNECE, 2016).

The trends of the **Secondary** nitrate source contributions were SE at both stations with very similar  $\tau$  (8.96 yr – 1068 1069 8.59 yr), TR (67-69 %) and RC (13-17%). The decrease observed for the contribution from the *Secondary* Nitrate 1070 source was related to the reduction in ambient NO<sub>x</sub> concentrations (Figures 8 and 9). Figure 9 shows the levels of 1071 tropospheric NO<sub>2</sub> column from 2005 to 2014 in South Europe from NASA NO<sub>2</sub> OMI level3 plotted using the 1072 Giovanni online data system (Acker and Leptoukh, 2007). In Spain it can be observed a general decrease of the 1073 concentrations of columnar NO<sub>2</sub> at regional level. Overall, the implementation of European directives affecting 1074 industrial and power generation emissions as well as the increase of the proportion of energy produced from 1075 renewable sources (cf. Figure 10 for Spain), among others, produced a significant reduction of  $SO_2$  and  $NO_x$ 1076 emissions. Around Barcelona the observed decreases were also attributed to the decrease of NO<sub>x</sub> emissions mainly from the five power generation plants around the city. Moreover, the implementation during the 2008-1077 1078 2012 Regional AQ Plan of SCRT (continuously regenerating PM traps with selective catalytic reduction for NO<sub>2</sub>) 1079 and the hybridization and shift to natural gas engines of the Barcelona's bus fleet may have had an influence 1080 in the observed reductions.

The decreasing trends (p<0.01) of the V-Ni bearing source contributions were SE and L at BCN and MSY, respectively, reflecting the trends observed at both stations for the concentrations of V and Ni (cf. Table 2). At BCN the characteristic times (τ) was very similar to the characteristic times calculated for PM<sub>10</sub> V and Ni (cf. Table 2) which were the main tracers of this source. TRs were around 61% at BCN and 64% at MSY and RCs were similar (19-25 %). The observed decrease in the V-Ni bearing source contribution was mainly attributed to the ban of the use of heavy oils and petroleum coke for power generation from 2008 in Spain.

1087 The Industrial/Metallurgy source contribution at BCN decreased exponentially (SE) at the rate of -0.10 µgm<sup>-</sup> <sup>3</sup>/yr (p<0.001) reflecting the SE decreasing trends observed for the main tracers of this pollutant source (Pb, Cd 1088 1089 and As; cf. Table 2). The decrease of industrial emissions was mainly attributed to the implementation of IPPC 1090 (Integrated Pollution Prevention and Control) Directives. Moreover, the observed decrease may be attributed to a decrease in the emissions from industrial production (smelters, Querol et al., 2007) at a regional scale 1091 1092 around Barcelona. Also, the financial crisis, whose impact on industrial production and use of fuels is evident since October 2008 also contributed to the observed trend. TR and RC for the Industrial source contributions 1093 1094 at BCN were 65% and 16%, respectively. As for the contributions from Secondary sulfate and nitrate sources, 1095 the exponential trend observed for the Industrial/Metallurgy source contribution suggested the attainment of a lower limit. As evidenced in Fig. 6 the contribution from this source from 2010 was guite low and rather 1096 1097 constant.

The contribution of the *Industrial/Traffic* source at MSY showed similar magnitude of the trend (-0.11 μgm<sup>-3</sup>/yr with p<0.01) compared to the BCN *Industrial* contribution trend, being both sources traced mostly by the same industrial tracers. The trend of this source at MSY was linear with TR and RC of 56% and 13%, respectively, similar to those calculated for *Industrial* source contribution at BCN.

1102	Finally, the <i>Mineral</i> source contribution at MSY showed linear little significant decreasing trend (p<0.1) in
1103	agreement with what observed at the same station by Cusack et al. (2012). As already noted in Section 3.3, this
1104	negative trend could be due to both a possible decrease of the emissions of finer anthropogenic mineral species
1105	from specific sources such as cement and concrete production and construction works and unusual weather
1106	conditions reducing Saharan dust contribution to PM and resuspension of dust.
1107	In order to further interpret the observed trends, annual data on the annual National Energy Consumption (NECo)

- from different energy sources (MINETUR, 2013) were also evaluated (Figure 10). Overall, the primary energy 1108 1109 consumption in Spain (NECo statistical data for Spain-MINETUR, 2013) increased from 2004 to 2007 and decreased from 2007 with marked decrease in 2009. Since 2009, the energy consumption indicator remained 1110 rather low and constant until 2012 when an additional decrease in 2013 and 2014 was observed. Oil 1111 consumption was fairly constant during 2004–2007 showing an important decrease during 2008–2014. This 1112 trend was probably governed by the fuel consumption for traffic road. Coal consumption remained constantly 1113 1114 high from 2004 to 2007 whereas, as for the emissions of SO<sub>2</sub> (Fig. 8), a sharp decrease occurred from 2007. However, in the period 2011-2014 there was an important increase of coal consumption leading to an average 1115 1116 consumption similar to the year 2008. However, the implementation of FGD systems contributed to maintain SO<sub>2</sub> at low concentrations, even in the coal production regions in Spain (cf. Querol et al., 2014). The 1117 1118 hydroelectric generation was rather specular to coal consumption. For example, the increase in 2010 of 1119 hydroelectric consumption, due to high rainfall rate, mirrored the decrease in the coal consumption observed the same year. Finally, renewable energy consumption increased by 440% from 2004 to 2014, with a gradual 1120 1121 growth in the NECo.
- 1122
- 1123

# 1124 5.0 Conclusions

PM chemical speciated data collected at two twin stations in NE of Spain (Barcelona: urban background station 1125 and Montseny: regional background station) during 2004 – 2014 were used to study trends of source 1126 1127 contributions from PMF analysis and of chemical species concentrations. Despite the fact the trends of different 1128 PM fractions (PM<sub>2.5</sub> and PM<sub>10</sub>) were linear during the period under study, the trends of specific chemical elements and source contributions were exponential demonstrating the different effectiveness and time of implementation 1129 1130 of different reduction strategies on specific pollutant sources. Statistically significant exponential trends (p < 1131 0.01 or 0.001) were mainly observed for the industrial tracers (Pb, Cd, As) in both PM<sub>10</sub> and PM<sub>2.5</sub> and at both 1132 sites. The concentrations of V and Ni showed exponential trends in BCN and linear trends at MSY likely because 1133 of the higher distance of the MSY station to the sources of V and Ni (shipping and, before 2008, energy production) compared to BCN. Traffic tracers at MSY (Sn, Cu) showed very similar linear decreasing trends with 1134 higher magnitude of the trends in the fine (PM<sub>2.5</sub>) fractions compared to PM<sub>10</sub> likely because of possible sources 1135

of coarser Sn and Cu reducing the magnitude of the trends in the PM<sub>10</sub> mass fraction. Sb at MSY showed marked
 exponential decreasing trends compared to other traffic tracers (Cu and Sn) which could be explained by a
 possible progressive reduction of Sb content in vehicle brakes. Secondary inorganic aerosols (SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup> and
 NH<sub>4</sub><sup>+</sup>) also showed marked decreasing trends (both linear and exponential) in both fractions and at both sites.
 However, in general the magnitude of the trends for these species and their statistical significance were higher
 at BCN compared to MSY.

1142 **The PM10 source contributions** that showed statistically significant downward trends at both Barcelona (BCN; 1143 UB) and Montseny (MSY; RB) were from **Secondary** sulfate, **Secondary** nitrate, and V-Ni bearing sources. For 1144 these source contributions the decreasing trends were exponential indicating that the trends were not gradual 1145 and consistent over time and that the effectiveness of the control measures for these pollutants was stronger at 1146 the beginning of the period under study (2004-2009 approximately) compared to the end of the period (Figs. 3 1147 and 4). Statistically significant decreasing trends were observed for the Industrial/Traffic and Mineral sources at 1148 MSY and the Industrial/metallurgy source at BCN. These sources were mostly linked with anthropogenic activities 1149 and the observed decreasing trends confirmed the effectiveness of pollution control measures implemented at 1150 EU or regional/local levels. The economic crisis which started in 2008 in Spain also contributed to the observed trends. Conversely, the contributions from sources mostly linked with natural processes such as Aged Marine 1151 1152 (at both BCN and MSY) and Aged Organic (at MSY) did not show statistically significant trends. The general 1153 trends observed for the calculated PMF source contributions well reflected the trends observed for the chemical 1154 tracers of these pollutant sources. The decrease in the **Secondary** sulfate source contribution was mainly attributed to the EC Directive on Large Combustion Plants implemented from 2008 in Spain, resulting in the 1155 1156 application of fuels gas desulfurization (FGD) systems in a number of large facilities. Moreover, according to the 1157 2008 Regional AQ Plan, the use of heavy oils and petroleum coke for power generation was forbidden around 1158 Barcelona from 2008 in favour of natural gas. As a consequence, a decrease of the contributions from the V-Ni 1159 bearing source at both sites was also observed. The decrease observed for the contribution of the Secondary 1160 *Nitrate* source was mainly due to the reduction in ambient NO<sub>x</sub> concentrations. In Spain a general decrease of the 1161 concentrations of  $NO_2$  at regional level was observed and it was mainly related with the lower energy 1162 consumption related with the financial crisis. The decrease of nitrates concentrations and **Secondary** nitrate 1163 source contributions around Barcelona was also attributed to the decrease of NO<sub>x</sub> emissions from the five power 1164 generation plants around the city. Moreover, a Regional AQ Plan implementing the SCRT (continuously 1165 regenerating PM traps with selective catalytic reduction for NO<sub>2</sub>) and the hybridization and shift to natural gas 1166 engines of the Barcelona's bus fleet may have had also an influence in NO<sub>x</sub> ambient concentrations. The 1167 Industrial/Metallurgy source contribution at BCN decreased exponentially reflecting the exponential trends observed for the main tracers of this pollutant source (Pb, Cd and As). The implementation of IPPC (Integrated 1168 1169 Pollution Prevention and Control) Directives together with a decrease in the emissions from industrial production (smelters) at a regional scale around Barcelona explained the observed trends. Overall, the 1170 1171 magnitude of the decreasing trends of the contributions of the pollutant sources were higher at BCN compared

1172	to MSY likely because of the proximity of the BCN measurement site to anthropogenic pollutant sources
1173	compared to the MSY site. The results presented in this work clearly confirm the beneficial effect of the AQ
1174	measures taken in recent years in Europe. However, the WHO limit values of specific pollutants, PM $_{ m 10}$ and PM $_{ m 2.5}$
1175	among these, are still exceeded especially at urban level and industrial hotspots. To meet the WHO guide levels
1176	important actions are still required for the next decade and the interpretation of past air quality trends may
1177	yield relevant outcomes for planning further cost-effective actions. We would like to highlight that a non-linear
1178	approach to trend studies is very attractive given that some air pollutants reported in this work showed not
1179	gradual-with-time reductions. Conversely, for specific pollutant source-contribution/concentration in our
1180	region, the decreasing trend was less steep at the end of the period compared to the beginning thus likely
1181	indicating the attainment of a lower limit. This was the case for example for the Secondary sulfate source
1182	contribution decreasing exponentially from 2004 to 2014 thus likely indicating a limited scope for further
1183	reduction of SO₂ emissions in our region.

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1320	Table 1: Trends of different PM mass fractions from gravimetry (grav) and optical (OPC) measurements at BCN (bold italic) and MSY (2004-
1321	2014). TR (%) = Total Reduction; RC (%) = Residual Component. Significance of the trends following the Mann-Kendall test: *** (p-value <
1322	0.001), ** (p-value < 0.01), * (p-value < 0.05), + (p-value < 0.1).

PMx	PMx		Mann-Kendall fit			
	Conc.	Conc.		Trend	TR	RC [%]
Fraction	2004	2014	р-	[µgm <sup>-3</sup> /yr]	[%]	
	(µgm <sup>-3</sup> )	(µgm <sup>-3</sup> )	value			
PM <sub>10</sub> (grav.)	41.1	19.2	***	<mark>-2.83</mark>	59.2	8.5
	19.2	13.9		<mark>-0.17</mark>	10.5	17.6
PM <sub>25</sub> (gray.)	31.6	13.2	***	<mark>-2.03</mark>	60.1	7.9
(grath)	16.2	9.8	+	-0.33	25.6	17.3
PM <sub>10</sub> (OPC)	39.1	19.8	***	<mark>-2.20</mark>	50.4	10.0

	18.6	12.3		<mark>-0.13</mark>	7.8	16.9
PM25 (OPC)	27.1	12.9	**	<mark>-1.55</mark>	49.6	9.8
	16.5	9.3	+	<mark>-0.26</mark>	21.2	17.5

1337Table 2: Mann-Kendall and Multi-exponential trends of different chemical species in PM10 at BCN (bold italic) and MSY. Type of trend:1338linear (L), single-exponential (SE), double exponential (DE); a ( $\mu$ gm<sup>-3</sup>) and  $\tau$  (yr) are the constants and the characteristic times, respectively,1339of the exponential data fittings; NL (%) = Non-Linearity; TR (%) = Total Reduction; RC (%) = Residual Component; ns = not statistically1340significant; ni = not included. Significance of the trends: \*\*\* (p-value < 0.001), \*\* (p-value < 0.01), \* (p-value < 0.05), + (p-value < 0.1).</td>

	PN ( <i>BCN</i>	Л <sub>10</sub> ;MSY)				Mann- Kendall fit	Multi-exponential fit				
Spacia	Concentration	Concentration	Fit	NL	p-	Trend	а	τ	Trend	TR	RC
specie	2004 (µgm <sup>-3</sup> )	2014 (µgm <sup>-3</sup> )	type	(%)	value	[µgm <sup>-3</sup> /yr]	(µgm-3)	(yr)	[µgm <sup>-3</sup> /yr]	(%)	(%)
Ph	0.02685	0.00694	<mark>SE</mark>	27	***		<mark>0.03246</mark>	<mark>6.12</mark>	<mark>- 0.00222</mark>	<mark>80</mark>	<mark>17</mark>
	0.00481	0.00190	SE	11	**		0.00553	10.22	-0.00031	62	13
Cd	0.00043	0.00015	<mark>SE</mark>	<mark>19</mark>	***		<mark>0.00048</mark>	<mark>7.59</mark>	<mark>-3.10e-5</mark>	<mark>73</mark>	<mark>17</mark>
ou	0.00017	0.00006	SE	18	**		0.00018	7.92	-1.12E-5	72	16
As	0.00094	0.00036	SE	14	***		0.00118	9.11	-7.07E-5	67	11
7.5	0.00029	0.00017	L	<10	***	<mark>-1.29E-5</mark>				<mark>50</mark>	<mark>9</mark>
v	0.01116	0.00454	SE	12	**		0.01502	10.04	<mark>-0.00086</mark>	63	17
-	0.00328	0.00175	L	<10	**	<mark>-0.00022</mark>				<mark>59</mark>	<mark>15</mark>
Ni	0.00531	0.00284	SE	11	***		0.00678	10.61	-0.00037	61	16
	0.00155	0.00100	L	<10	**	<mark>-7.10E-5</mark>				<mark>43</mark>	<mark>20</mark>
Sn	ni	ni									
511	0.00127	0.00057	L	<10	*	<mark>-3.65E-5</mark>				<mark>39</mark>	<mark>16</mark>
Cu	ni	ni									
54	0.00420	0.00216	L	<10	*	-0.00014				<mark>36</mark>	20
Sb	ni	ni									

SQr <sup>1</sup> 5.74436         2.28596         SE         12         11         6.56033         9.87         40.37868         64           NO3:         5.07816         1.72401         SE         12         14         6.49990         9.83         0.37868         64           NO3:         5.07816         1.72401         SE         12         14         6.49990         9.83         0.37868         64           NH:         1.9262         0.57008         SE         12         14         6.49990         9.83         0.37868         64           NH:         1.92662         0.57008         SE         12         1.00         1.90645         9.64         0.1095         65           1.14268         0.40135         SE         13         1.28868         9.26         0.07640         66           Al:03         ni         ni         1         1.28868         9.26         0.07640         66         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102         1.02102	SD,2         5.74436         2.28596         SE         12         11         0.11056         0         037561         6.4         12           NO1         5.07816         1.7712         L         <10         **         011056         0         037641         6.4         10           NO1         5.07816         1.77407         SE         12         11         6.49890         9.83         10.37641         64         1           NO1         1.80724         0.67419         L         <10         **         010591          6         5         1           NH         1.92662         0.57008         SE         12         12         1         1.90645         9.64         01052         65         1           NH         1.22557         0.40382         L         <10         *         010281          1         2866         9.26         00764         66         1           Ca         nl         nl          1         10         10         10         10         10         10         10         10         10         10         10         10         10         10         10         10 <th></th>												
3.44         2.84849         1.67712         L         <10         ***         0.11836           6.49890         9.83         0.37484         64           NO <sub>3</sub> *         5.07816         1.72401         SE         12         1         6.49890         9.83         0.37484         64           NO <sub>3</sub> *         1.80724         0.67419         L         <10	3.41         2.84649         1.67712         L         <10         **         0.11834         6.49890         9.83         10.3244         6.4           1.80724         0.67419         L         <10	\$0.2	5.74436	2.28596	SE	12	***		6.56033	9.81	<mark>-0.37868</mark>	64	
NO;:         5.07816         1.72401         SE         12         1         6.49890         9.83         0.37484         64           1.80724         0.67419         L         <10	NO1         5.07816         1.72401         SE         12 <th12< th=""> <th12< th="">         12</th12<></th12<>	304-	2.84849	1.67712	L	<10	**	-0.11836				<mark>42</mark>	
INO3         1.80724         0.67419         L         <10         **         0.10592          64         0.17095         65           NH4*         1.92662         0.57008         SE         12         1         1.90645         9.64         0.17095         65           Al <sub>2</sub> O3         ni         ni          1.28868         9.26         0.07840         66           Al <sub>2</sub> O3         ni         ni           0.02383              O.72357         0.46382         L         <10	Image         1.80724         0.67419         L         <10         **         0.10593	NOs	5.07816	1.72401	SE	12	**		6.49890	9.83	-0.37484	64	
NH4*         1.92062         0.57008         SE         12         14         1.90645         9.64         0.11095         65           ALO3         ni         ni <t< td=""><td>NH.*         1.92062         0.57008         SE         12         14         1.90645         9.64         0.11095         65           AlcO3         II         III         IIII         IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII</td><td>1403</td><td>1.80724</td><td>0.67419</td><td>L</td><td>&lt;10</td><td>**</td><td>-0.10593</td><td></td><td></td><td></td><td><mark>5</mark>4</td><td></td></t<>	NH.*         1.92062         0.57008         SE         12         14         1.90645         9.64         0.11095         65           AlcO3         II         III         IIII         IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	1403	1.80724	0.67419	L	<10	**	-0.10593				<mark>5</mark> 4	
Initial         Initial <t< td=""><td>Initial         Initial         <t< td=""><td>NH4<sup>+</sup></td><td>1.92062</td><td>0.57008</td><td>SE</td><td>12</td><td>***</td><td></td><td>1.90645</td><td>9.64</td><td>-0.11095</td><td>65</td><td></td></t<></td></t<>	Initial         Initial <t< td=""><td>NH4<sup>+</sup></td><td>1.92062</td><td>0.57008</td><td>SE</td><td>12</td><td>***</td><td></td><td>1.90645</td><td>9.64</td><td>-0.11095</td><td>65</td><td></td></t<>	NH4 <sup>+</sup>	1.92062	0.57008	SE	12	***		1.90645	9.64	-0.11095	65	
ni         ni<	ni         ni<	1 1 1 4	1.14268	0.40135	SE	13	<b>*</b>		1.28868	9.26	-0.07640	66	
0.72357         0.46382         L         <10         *         002383         Image: Comparison of the comparison	Ca         0.72357         0.46382         L         <10         *         0.02282         I         <10         I <thi< th="">         I         <thi< th=""> <thi< th=""></thi<></thi<></thi<>	Al <sub>2</sub> O <sub>2</sub>	ni	ni									
Ca         ni         ni	Ca         ni         ni<	1.203	0.72357	0.46382	L	<10	*	-0.02383				<mark>34</mark>	
Image: Normal System         0.42703         0.28279         L         <10         *         000038         Image: Normal System	Image: Normal System         Image: Output System         Image: Ou	Са	ni	ni									
ni         ni <td>Fe         ni           0.000994</td> <td></td> <td>0.42703</td> <td>0.28279</td> <td>L</td> <td>&lt;10</td> <td>*</td> <td><mark>-0.01638</mark></td> <td></td> <td></td> <td></td> <td><mark>38</mark></td> <td></td>	Fe         ni           0.000994		0.42703	0.28279	L	<10	*	<mark>-0.01638</mark>				<mark>38</mark>	
0.22371         0.14895         L         <10         +         1000593         I         I         I           Na         1.02188         0.77408         L         <10	0.22371         0.14895         L         <10         +         000593         -         -         -         8           Na         1.02188         0.77408         L         <10	Fe	ni	ni									
1.02188         0.77408         L         <10         *         0.03943         Image: Comparison of the second se	Na         1.02188         0.77408         L         <10         *         0.03943         Image: Comparison of the second sec		0.22371	0.14895	L	<10	+	-0.00593				<mark>6</mark>	
		Na	1.02188	0.77408	L	<10	*	<mark>-0.03943</mark>				<mark>34</mark>	
		INC					ns						

1351<br/>1352**Table 3**: Mann-Kendall and Multi-exponential trends of different chemical species in PM2.5 at BCN (bold italic) and MSY. Type of trend:<br/>linear (L), single-exponential (SE), double exponential (DE); **a** ( $\mu$ gm<sup>-3</sup>) and  $\tau$  (yr) are the constants and the characteristic times, respectively,<br/>of the exponential data fittings; NL (%) = Non-Linearity; TR (%) = Total Reduction; RC (%) = Residual Component; ns = not statistically<br/>significant; ni = not included. Significance of the trends: \*\*\* (p-value < 0.001), \*\* (p-value < 0.01), \* (p-value < 0.05), + (p-value < 0.1).</th>

	PN ( <i>BCN</i>	<b>Л</b> 2.5 (:MSY)				Mann-Kendall fit	Multi-exponential fit				
Creale	Concentration	Concentration	Fit	NL	n yalya	Trend	а	τ	Trend	TR	RC
specie	2004 (µgm <sup>-3</sup> )	2014 (µgm <sup>-3</sup> )	type	(%)	p-value	[µgm <sup>-3</sup> /yr]	(µgm-3)	(yr)	[µgm <sup>-3</sup> /yr]	(%)	(%)
Ph	0.02117	0.00500	SE	27	***		0.02390	6.24	-0.00163	80	13
10	0.00642	0.00149	SE	28	***		0.00716	6.08	-0.00049	81	18
Cd	0.00041	0.00011	SE	23	***		0.00047	6.81	<mark>-3.11E-5</mark>	77	13
cu	0.00020	0.00005	SE	23	***		0.00020	6.77	-1.35E-5	77	18
٨٥	0.00069	0.00027	SE	14	***		0.00091	9.00	<mark>-5.43E-5</mark>	67	11
AS	0.00029	0.00013	SE	15	**		0.00033	8.56	-2.04E-5	69	19
v	0.00823	0.00368	<mark>SE</mark>	<mark>11</mark>	**		<mark>0.01121</mark>	<mark>11.13</mark>	-0.00061	59	16
v	0.00271	0.00130	L	<10	**	<mark>-0.00017</mark>				<mark>64</mark>	<mark>24</mark>
Ni	0.00402	0.00185	<mark>SE</mark>	<mark>10</mark>	**		0.00498	<mark>11.23</mark>	-0.00027	59	15
NI INI	0.00189	0.00080	SE	13	**		0.00205	9.36	-0.00012	42	21
Sn	ni	ni									
511	0.00157	0.00043	L	<10	***	-0.00084				<mark>61</mark>	<mark>12</mark>
Cu	ni	ni									
Cu	0.00394	0.00113	SE	14	**		0.00426	8.99	-0.00026	67	13
	ni	ni									
Sb	0.00053	0.00015	DE	48	**		0.00069 1.3E-6	4.52 -2.50	-3.86E-5	70	16

4.86564	1.92388	SE	12	***		5.64582	9.69	<u>-0.32778</u>	64	9
2.98922	1.43381	L	<10	**	<mark>-0.16222</mark>				<mark>54</mark>	<mark>15</mark>
3.45513	0.86002	SE	19	***		4.14459	7.61	-0.26753	73	16
1.66095	0.29452	SE	30	***		1.96014	5.81	- <mark>0.13550</mark>	82	21
2.19735	0.68393	SE	11	***		2.27813	10.53	<mark>-0.12701</mark>	61	15
1.39366	0.48049	SE	18	**		1.62588	7.94	- <mark>0.10266</mark>	72	14
ni	ni									
0.30245	0.10153	SE	13	*		0.26678	9.36	- <mark>0.01574</mark>	66	35
ni	ni									
0.11478	0.06540	L	<10	+	-0.00494				<mark>50</mark>	<mark>33</mark>
ni	ni									
0.09679	0.03716	L	<10	*	<mark>-0.00504</mark>				<mark>61</mark>	<mark>31</mark>
0.27476	0.17863	L	<10	+	<mark>-0.01247</mark>				<mark>41</mark>	<mark>15</mark>
0.13091	0.07252	L	<10	*	-0.00584				<mark>45</mark>	<mark>18</mark>
	A.86564           2.98922           3.45513           1.66095           2.19735           1.39366           ni           0.30245           ni           0.11478           ni           0.09679           0.27476           0.13091	4.86564         1.92388           2.98922         1.43381           3.45513         0.86002           1.66095         0.29452           2.19735         0.68393           1.39366         0.48049           ni         ni           0.30245         0.10153           ni         ni           0.11478         0.06540           ni         ni           0.09679         0.03716           0.27476         0.17863           0.13091         0.07252	4.86564         7.92388         SE           2.98922         1.43381         L           3.45513         0.86002         SE           1.66095         0.29452         SE           2.19735         0.68393         SE           1.39366         0.48049         SE           ni         ni         0           0.30245         0.10153         SE           ni         ni         0           0.11478         0.06540         L           ni         ni         1           0.09679         0.03716         L           0.13091         0.07252         L	4.86564         1.92388         SE         12           2.98922         1.43381         L         <10	4.86564       1.92388       SE       12       14         2.98922       1.43381       L       <10	4.86564       1.92388       SE       12       14         2.98922       1.43381       L       <10	4.86564       1.92388       SE       12       1       5.64582         2.98922       1.43381       L       <10	4.86564       1.92388       SE       72       1       1       5.64582       9.69         2.98922       1.43381       L       <10	4.86564       1.92388       SE       72       1       1       5.64582       9.69       4.3278         2.98922       1.43381       L       <10	4.86564       1.92388       SE       12       12       12       12       5.64582       9.69       0.32778       64         2.98922       1.43381       L       <10

**Table 4**: Mann-Kendall and Multi-exponential trends of source contributions in  $PM_{10}$  from PMF at BCN (bold italic) and MSY. Type: linear1365(L), single-exponential (SE), double exponential (DE); **a** ( $\mu$ gm<sup>-3</sup>) and  $\tau$  (yr) are the constants and the characteristic times, respectively, of1366the exponential data fittings; NL (%) = Non-Linearity; TR (%) = Total Reduction; RC (%) = Residual Component; **ni = not included**. Significance1367of the trends following the Mann-Kendall test: \*\*\* (p-value < 0.001), \*\* (p-value < 0.01), \* (p-value < 0.05), + (p-value < 0.1).</td>

	PN ( <i>BCN</i>	<b>/1</b> 10 (;MSY)					Mann- Multi-exponential fit				
Source	Contribution 2004 (µgm <sup>-3</sup> )	Contribution 2014 (µgm <sup>-3</sup> )	Fit type	NL (%)	p-value	Trend [µgm <sup>-3</sup> /yr]	a (µgm <sup>-3</sup> )	τ (yr)	Trend [µgm <sup>-3</sup> /yr]	TR (%)	RC (%)
<mark>Secondary</mark> sulfate	10.27	3.38	DE	45	**		12.33 3.82	1.65 105.80	<mark>-0.7</mark> 1	67	16
	6.57	3.07	SE	<mark>12</mark>	**		<mark>5.99</mark>	<mark>13.22</mark>	<mark>-0.32</mark>	53	<mark>21</mark>
Secondary	6.99	1.96	SE	14	***		8.54	8.96	<mark>-0.51</mark>	67	13
nitrate	2.03	0.47	SE	15	**		2.44	8.59	<mark>-0.15</mark>	69	17
V Ni booring	4.23	1.84	SE	11	**		5.66	10.59	<mark>-0.3</mark> 2	61	19
v-ni bearing	0.79	0.44	L	8	**	<mark>-0.07</mark>				<mark>64</mark>	<mark>25</mark>
Industrial/Metall urgy (BCN)	1.64	0.71	SE	21	***		1.76	<mark>9.56</mark>	-0.10	<mark>65</mark>	<mark>16</mark>
Mineral	ni	ni									
	3.46	2.32	L	5	+	<mark>-0.10</mark>				<mark>30</mark>	<mark>21</mark>
Industrial/Traffic (MSY)	2.08	1.01	L	7	**	<mark>-0.11</mark>				56	13

- **Figure Captions:** Figure 1: Location of the Barcelona (BCN) and Montseny (MSY) measuring stations. Red full circle highlights the location of the BCN measuring station before 2009. Green full circle highlights the new location of the BCN (from 2009) and MSY measuring stations. Figure 2: Mann-Kendall fit of PM2.5 trends at MSY station for the periods 2002-2010 (as in Cusack et al., 2012), 2004 – 2014 (this work),
- and 2002 2014 (largest period available in the time of writing). Reported are: magnitude of the trends [µgm<sup>-3</sup>/yr]; p-value; Total Reduction (TR) and Residual Component (RC). Significance of the trends following the Mann-Kendall test: \*\*\* (p-value < 0.001), \*\* (p-value < 0.01),
- 1391 \* (p-value < 0.05), + (p-value < 0.1).
- Figure 3: Mann-Kendall (MK) and Multi-exponential (ME) trends for chemical species at BCN in PM<sub>10</sub>. Measured concentration (green line);
   Multi-exponential trend (red line); Multi-exponential residuals (blue line); Mann-Kendall trend (black line); Mann-Kendall residuals (grey
   Iine). Trend type: linear (L), single-exponential (SE), double exponential (DE).
- Figure 4: Mann-Kendall (MK) and Multi-exponential (ME) trends for chemical species at MSY in PM<sub>10</sub>. Measured concentration (green line);
   Multi-exponential trend (red line); Multi-exponential residuals (blue line); Mann-Kendall trend (black line); Mann-Kendall residuals (grey
   Iine). Trend type: linear (L), single-exponential (SE), double exponential (DE).

- Figure 5: Source contributions from PMF model in PM<sub>10</sub> at Montseny (MSY) and Barcelona (BCN). Mean values during 2004-2014. Values
   reported are: Source; μg/m<sup>3</sup>; %.
- 1400 Figure 6: Mann-Kendall and Multi-exponential trends for source contributions in PM<sub>10</sub> at BCN. Measured concentration (green line); Multi-
- exponential trend (red line); Multi-exponential residuals (blue line); Mann-Kendall trend (black line); Mann-Kendall residuals (grey line).
   Trend type: linear (L), single-exponential (SE), double exponential (DE). Highlighted with yellow colour the source contributions at BCN
- 1403 from *Mineral, Traffic* and *Road/work resuspension* were excluded from the trend discussion.
- 1404 Figure 7: Mann-Kendall and Multi-exponential trends for source contributions in PM<sub>10</sub> at MSY. Measured concentration (green line); Multi-
- 1405 exponential trend (red line); Multi-exponential residuals (blue line); Mann-Kendall trend (black line); Mann-Kendall residuals (grey line).
   1406 Trend type: linear (L), single-exponential (SE), double exponential (DE).
- 1407 **Figure 8**: Spanish national emission of SO<sub>2</sub> and NO<sub>X</sub> (normalized to year 2004).
- Figure 9: NASA OMI level 3 tropospheric NO<sub>2</sub> column plotted using the Giovanni online data system, developed and maintained by theNASA GES DISC.
- Figure 10: Annual (2004–2014) energy consumption for Spain (normalized to year 2004). Data from the Spanish Ministry of Industry(MINETUR, 2013).
- 1412
- 1413
- 1414









1426 Figure 2



- 1436 Figure 3



1442 Figure 4





1464 Figure 6



- 1467 Figure 7









- 1482 Figure 9



- 1488 Figure 10

# 1508 Supporting Information

- **1)** Effect of the change of the location of the measuring station in BCN in 2009



Figure SI-1: Trends of PM<sub>10</sub> concentrations from gravimetric measurements at BCN and MSY. Red rectangle highlights the decrease of PM<sub>10</sub> concentration at BCN due to the change of the location of the BCN measuring station in 2009.

2) PMF source profiles at BCN and MSY





Source profiles from PMF analysis at BCN and MSY. Common sources at both BCN and MSY were: Secondary Sulfate (secondary inorganic source traced by SO<sub>4</sub><sup>2-</sup>, NH<sub>4</sub><sup>+</sup> with contribution from OC), Secondary nitrate (secondary inorganic source traced by NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup>), V-Ni bearing (traced mainly by V, Ni and SO<sub>4</sub><sup>2-</sup> it represents the direct emissions from heavy oil combustion), mineral (traced by typical crustal elements such as Al, Ca, Ti, Rb, Sr), aged marine (traced by Na and CI mainly with contributions from  $SO_4^{2-}$  and  $NO_3^{-}$ ). Non common sources at MSY were: Industrial/Traffic source (Anthropogenic source traced by EC, OC, Cr, Cu, Zn, As, Cd, Sn, Sb and Pb includes contributions from anthropogenic sources such as traffic and metallurgic) and Aged organics (traced by OC and EC mainly with maxima in summer indicating a biogenic origin). Non common sources at BCN were: Traffic (traced by Cnm, Cr, Cu, Sb and Fe mainly and contributing 5.14 µg/m<sup>3</sup> (15.1%)), Road resuspension (traced by both crustal elements, mainly Ca, and traffic tracers such as Sb, Cu and Sn and contributing 4.25 µg/m<sup>3</sup> (12.5%)) and Industrial (traced by Pb, Cd, As and Zn and contributing 0.96  $\mu$ g/m<sup>3</sup> (2.8%). 

1557 3) OC:EC ratio statistic at Montseny (MSY)



1558

Figure SI-3: OC and EC scatterplot (a) and frequency distribution of the OC:EC ratio (b) at Montseny (MSY)station.

- 1561
- 1562 Mean and median values of the OC:EC ratio at MSY were 9.1 and 7.8, respectively.
- 1563

#### y = 1.0333x + 0.1166 1.2044x + 0.0578 y = 1.0464x - 0.1002 35 $R^2 = 0.9962$ 25 $R^2 = 0.988$ $R^2 = 0.8943$ 30 20 25 whole 20 15 15 10 10 Amm Sul Industria Nit 2007-2014 2007-2014 35 40 <sup>2</sup>200<sup>3</sup>7-201<sup>4</sup> 35 y = 1.0013x - 0.2388 y = 0.9992x + 0.188 y = 0.9954x - 0.0059 30 30 R<sup>2</sup> = 0.9963 R<sup>2</sup> = 0.99 25 = 0.9843 25 25 20 whole 20 20 15 15 15 10 10 Road Resusp Marine Traffic <sup>10</sup> <sup>15</sup> <sup>20</sup> <sup>25</sup> 2007-2014 ${\overset{{}_{10}}{2007}}{\overset{{}_{15}}{-2014}}$ 35 30 35 2007-2014 80 y = 0.9722x - 0.5228 y = 1.1085x + 0.0875 R<sup>2</sup> = 0.9972 18 70 R<sup>2</sup> = 0.9925 16 14 60 50 whole 12 10 40 30 20 10 Minera V-Ni 6 8 10 12 14 16 18 20 2007-2014 20 30 40 50 60 2007-2014 10 80

# 1564 4) PMF Barcelona: 2007-2014 (with OC and EC) vs. 2004 – 2014 (with Cnm)

- 1565
- Figure SI-4: Comparison between PMF results at BCN obtained using the period 2007-2014 (separate OC and EC
   measurements available) and using the whole period 2004-2014 (Cnm was used in PMF).
- 1568
- 1569 5) OC:EC ratio statistic at Barcelona (BCN)
- 1570



1572 Figure SI-5: OC and EC scatterplot (a) and frequency distribution of the OC:EC ratio (b) at Barcelona (BCN)1573 station.