

1 Decadal trends (2013–2023) in PM₁₀ sources and oxidative 2 potential at a European urban supersite (Alpine Valley, 3 Grenoble, France)

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13 Abstract

14 The identification of particulate matter (PM) sources and the quantification of their contribution to the urban
15 environment is a necessary input for policymakers to reduce the air pollution impacts. The association between
16 the PM sources and the oxidative potential (OP) is also a key indicator for evaluating the ability of PM sources to
17 induce *in-vivo* oxidative stress and lead to adverse health effects, which becomes an emerging metric in the
18 Directive on ambient air quality (22024/2881/EU). Most studies in Europe have focused on PM and OP sources
19 in the short term, for only 1 or 2 years. However, the efficiency of reduction policies, trends, and epidemiological
20 impacts cannot be properly evaluated with such short-term studies due to a lack of statistical robustness. Here,
21 long-term PM₁₀ filter sampling at the Grenoble (France) urban background supersite and detailed chemical
22 analyses were used to investigate decadal trends of the main PM sources and related OP metrics. Positive matrix
23 factorization (PMF) analyses were conducted on the corresponding 11-year dataset (Jan 2013 to May 2023, n =
24 1570), enlightening the contributions of 10 PM sources: mineral dust, sulfate-rich, primary traffic, biomass
25 burning, primary biogenic, nitrate-rich, MSA-rich, aged sea salt, industrial and chloride-rich. The stability of the
26 chemical profile of these sources was validated by comparison with the profiles retrieved from shorter-term (3
27 years) successive PMF analyses. A Seasonal-Trend using LOESS decomposition was then applied to evaluate the
28 trends of these PM₁₀ sources, which revealed a substantial decrease in PM₁₀ (-0.73 µg m⁻³ yr⁻¹) as well as that of
29 many of the PM₁₀ sources. Specifically, negative trends for primary traffic and biomass burning sources are
30 detected, with a reduction of 0.30 and 0.11 µg m⁻³ yr⁻¹, respectively. The OP PM₁₀ source apportionment in 11
31 years confirmed the high redox activity of the anthropogenic sources, including biomass burning, industrial, and
32 primary traffic. Eventually, downward trends were also observed for OP_{AA} and OP_{DTT}, mainly driven by the
33 reduction of residential heating and transport emissions, respectively.

34 Keywords: PM₁₀ source apportionment, OP PM₁₀ source apportionment, long-term trend, Positive matrix
35 factorization.

36 **1. Introduction**

37 Particulate matter (PM) is the main atmospheric pollutant that significantly impacts human health, climate, and
38 the environment (Fuzzi et al., 2015; Grantz et al., 2003; Pope and Dockery, 2006), which is emitted directly or
39 formed through complex processes in the atmosphere from natural and anthropogenic gaseous precursors. The
40 identification of PM sources is important to investigate their composition, contribution, and evolution, which is
41 one necessary input for policymakers to apply strategies in reducing their impact. There are various methodologies
42 to identify these sources, where receptor models are widely used to perform source apportionment (SA) due to
43 their flexibility and performance. Positive Matrix factorization (PMF) is one of the most popular among these
44 receptor models, as it has been developed to allow SA analysis without any prior information other than the
45 measurement and uncertainty input matrices (Hopke, 2016). Scores of studies using PMF have been applied in
46 different typologies of sites over the last 15 years, with urban areas being the most common (Hopke et al., 2020;
47 Viana et al., 2008).

48 The adverse health effects of PM can be assessed through different pathways, one of which uses the concept of
49 oxidative stress within the lung (Pope and Dockery, 2006). PM has the ability to generate reactive oxygen species
50 (ROS), which can cause an imbalance with antioxidants in the lungs, eventually causing oxidative stress. This
51 capacity is evaluated as the oxidative potential (OP) of PM (Ayres et al., 2008; Li et al., 2008; Lodovici and
52 Bigagli, 2011; Mudway et al., 2020; Nelin et al., 2012; Rao et al., 2018). The redox activity of PM is mainly
53 dependent on their compositions; nevertheless, the correlation between individual components of PM and OP is
54 probably not the best approach for understanding the impact of ambient PM because of their complex mixture
55 preventing the quantification of all components of interest (Borlaza, 2021; Calas et al., 2018; Weber et al., 2018).
56 Therefore, the relationship between OP and PM sources has been investigated as a more holistic approach (Bates
57 et al., 2018; Dominutti et al., 2023; Weber et al., 2021). The implementation steps of such an approach can include,
58 first, a PM source apportionment (SA) (usually using PMF), allowing the identification of PM sources and their
59 contribution to PM. Then, the relationship between OP and PM sources is investigated by performing some
60 regression techniques, potentially including linear and non-linear ones (Ngoc Thuy et al., 2024).

61 The OP of PM is becoming an emerging metric for the European regulation on air quality, included in the new
62 European Air Quality Directive (Directive (EU) 2024/2881) as a recommended measurement at super sites in each
63 member state in order to improve the knowledge about the variability of the OP and eventually allow the
64 connections with epidemiological studies. Most previous studies have focused on PM and OP sources over a
65 relatively short period, typically less than 1 or 2 years (Borlaza et al., 2022a; Pietrodangelo et al., 2024; Weber et
66 al., 2019). Such short-term studies assess the PM and OP sources as well as their contribution, providing
67 information on the intrinsic OP of PM sources, allowing for the development of OP modeling (Daellenbach et al.,
68 2020; Vida et al., 2024) and eventually designing some public policies (Borlaza, 2021). However, long-term series
69 are needed both for evaluating the efficiency of such reduction policies in connection with the evolution of
70 contributions from sources and also for implementing epidemiological studies (Borlaza-Lacoste et al., 2024).

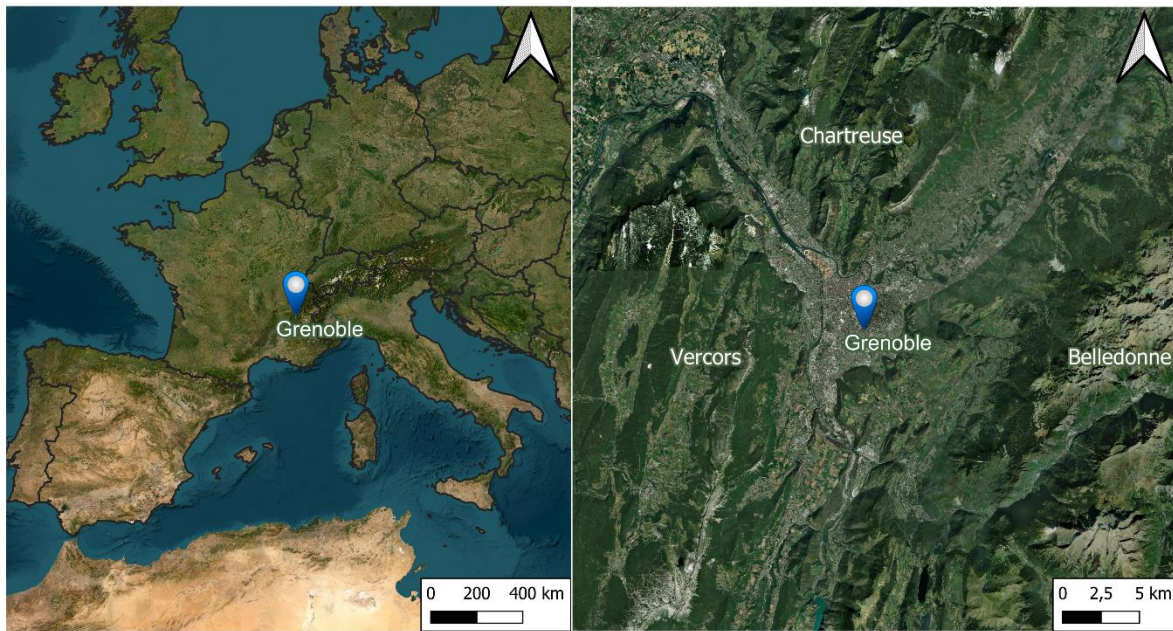
71 The present study is based on a long-term measurement program conducted in the city of Grenoble (France),
72 resulting from a sustained collaboration between the local network (Atmo AuRA), the French Reference
73 Laboratory for Air Quality Monitoring (LCSQA), and the Institute of Environmental Geosciences (IGE) to
74 investigate long-term evolution of PM₁₀ sources and OP in the PM₁₀ as well as their tendencies in the urban
75 background of the city. Here, we assessed these source contributions from daily ambient PM₁₀ samples obtained

76 from 2013 to 2023 (n= 1570) using the EPA PMF model at this site selected as one of the French urban supersites
77 for the new EU 2024/2881 Air Quality Directive. The database was augmented using imputation techniques in
78 order to fill some of the gaps in the data, relative to metallic trace elements. Since PMF is rarely applied to such
79 a long-term database, several evaluations of the validity of solutions were also implemented. The PMF-derived
80 PM₁₀ sources were then used to perform OP SA from 2013 to 2022 (n=1570). The trend of PM₁₀ concentration,
81 of the PM₁₀ sources, and the OP measurements are eventually discussed in relation to several potential factors of
82 influence.

83 **2. Methodology**

84 **2.1. Sampling site**

85 PM₁₀ daily samples were collected at an urban background site (Grenoble - Les Frênes), in the southern area of
86 Grenoble, France (45°09'41" N, 5°44'07" E). This city is known as the French Alps' capital, sprawling over 18.13
87 km² with about 154,000 inhabitants in 2023, but nearly 500,000 within the larger urbanized area (about 15 km
88 radius). With an average altitude of about 200 masl, the city sits within a complex mountainous geomorphology
89 and is surrounded by three mountain massifs: Chartreuse, Vercors, and Belledonne ([Figure 1](#)[Figure 1](#)). A pendular
90 wind regime between the three valleys of the basin regulates the ventilation of the atmosphere, with frequent
91 thermal inversion during cold periods, leading to the accumulation of pollutants. The air quality is monitored at
92 several sites in Grenoble by the regional agency (Atmo AuRA), including the urban background site of this study,
93 which is equipped with a large array of instruments. Particularly, the chemistry of PM₁₀ collected on filters has
94 been speciated at this site since 2008, within several programs, including the CARA program from the French
95 Ministry of Environment (Favez et al., 2021) and several research programs such as QAMECS (Borlaza et al.,
96 2021), or SOURCES (Weber et al., 2019). Many aspects of air quality in Grenoble were reported for this site,
97 including the characteristics of secondary anthropogenic PM fraction (Baduel et al., 2009, 2012; Favez et al.,
98 2010; Tomaz et al., 2016, 2017), of the biogenic PM components (Brighty et al., 2022; Samaké et al., 2019a, a),
99 as well as the PM OP (Borlaza, 2021; Dominutti et al., 2023; Weber et al., 2021). Several studies of one-year PM
100 sources apportionment were also performed in 2013 (Srivastava et al., 2018) and 2017-2018 (Borlaza et al., 2021).
101 Despite the difference in input data and periods of the studies, similar main sources of PM were quantified in both
102 studies, including residential heating, traffic, and secondary inorganic aerosol (SIA).



103
104 **Figure 1.** The sampling site is located in the Southeast of France (left figure), surrounded by 3 mountains massifs
105 (Vercors, Chartreuse, and Belledonne). Background map: ESRI satellites.

106 **2.2. Sampling and chemical analyses**

107 **2.2.1. PM₁₀ and their inorganic and organic composition**

108 The daily PM₁₀ sampling was performed every third day from 02/01/2013 to 28/05/2023, on 150 mm-diameter
109 quartz fibre filter (Tissu-quartz PALL QAT-UP 2500 diameter 150 mm) using high-volume samplers (Digitel
110 DA80, 30 m³ h⁻¹). A weekly PM₁₀ sampling was conducted during the same period using a low-volume sampler
111 (Partisol, 1 m³ h⁻¹) onto 47mm diameter quartz fibre filters (Tissuquartz PALL QAT-UP 2500 diameter 47 mm).
112 The processes of preparation, handling, and storing filters, in order to guarantee optimum quality for chemical
113 analyses were presented in Borlaza et al. (2021). Field blank filters were also collected (about 8-10% of the total
114 samples) to estimate the detection limits and evaluate the filter contamination during the overall handling and
115 analysis processes.

116 The daily PM₁₀ samples (n = 1570) and field blanks were analyzed for elemental carbon (EC) and organic carbon
117 (OC), major ions (Cl⁻, NO₃⁻, SO₄²⁻, Na⁺, NH₄⁺, K⁺, Mg²⁺, Ca²⁺), methanesulfonic acid (MSA), anhydrous sugar
118 and saccharides (levoglucosan, mannosan, arabitol, mannitol), and trace elements (As, Ba, Cd, Cr, Cu, Mn, Ni, Pb,
119 Sb, V, Zn). However, the concentrations of the daily trace elements were analyzed only in 3 periods, including:
120 (1) from January 2nd, 2013 to December 31st, 2013 (n = 122), (2) from February 28th, 2017 to March 13th, 2018
121 (n = 125), (3) from June 30th, 2020 to June 18th, 2021 (n=115). The weekly samples and blanks were analyzed
122 for trace metal concentrations for the whole sampling period (n = 842).

123 All analyses were previously described in detail (Borlaza et al., 2021). In brief, EC and OC analysis was performed
124 using a Sunset Lab analyser with the EUSAAR2 thermo-optical protocol. The eight major ionic components and
125 MSA were analyzed, after aqueous extraction of the filters using orbital shacking, by ionic chromatography using
126 an ICS3000 dual-channel chromatograph (Thermo-Fisher) with a CS16 column for cation analysis and an AS11
127 HC column for anion analysis. The anhydrous-sugar and saccharides analyses were performed on the same water
128 extract by an HPLC method using PAD (Pulsed Amperometric Detection) (model Dionex DX500 + ED40) with
129 Metrosep columns (Carb 1-Guard+A Supp15-150+Carb1-150) in the period before the year 2017. From 2017 to

130 the present, the measurement with ICS 5000 with pulsed amperometric detection (HPLC-PAD) was performed
131 following the CEN method (European committee for standardization, 2024). The analysis is isocratic with 15%
132 eluent of sodium hydroxide (200 mM), sodium acetate (4 mM), and 85% water at 1 mL min⁻¹.
133 The daily and weekly metals were measured by Inductively coupled plasma mass spectroscopy (ICP-MS) (ELAN
134 6100 DRC II PerkinElmer or NEXION PerkinElmer). The measurement was performed on the mineralization of
135 a 38 mm diameter punch of each filter, using 5 mL of HNO₃ (70 %) and 1.25 mL of H₂O₂ for 30 min at 180°C in
136 a microwave.

137 **2.2.2. OP analysis**

138 Two complementary OP assays, including the two probes ascorbic acid (AA) and dithiothreitol (DTT) were
139 performed on the same filters with PM₁₀ components analysis (from 02/01/2013 to 28/05/2023, n = 1570). ~~Filter~~
140 ~~samples are extracted using a simulated lung fluid during 1h15 at 37°C, pH 7.4, as described in Calas et al. (2017),~~
141 ~~which creates a physiological environment for the extraction~~ ~~Filter samples are extracted using a simulated lung~~
142 ~~fluid which is the mixing of Gamble and DPPC (dipalmitoylphosphatidylcholine) solutions, during 1h15 at 37°C,~~
143 ~~pH 7.4, creating a physiological environment for the extraction~~ (Calas et al., 2017). The AA method quantifies
144 the consumption of ascorbic acid, an endogenous antioxidant in the lung, by PM and was described in Calas et al.
145 (2017, 2018). The reaction mixture (extract + AA) was transferred to UV-transparent 96-well plates (CELLSTAR,
146 Greiner-Bio), and the residual AA was measured at 265 nm with a TECAN Infinite M200 Pro spectrophotometer.
147 The AA consumption rate (nmol min⁻¹) reflects the capacity of PM₁₀ to catalyze electron transfer from AA to
148 molecular oxygen.

149 DTT assay relies on dithiothreitol, a chemical surrogate for cellular reducing agents, allowing for emulation of in
150 vivo interaction among PM₁₀ and biological reducing agents (for example, nicotinamide adenine dinucleotide
151 (NADH), nicotinamide adenine dinucleotide phosphate oxidase (NADPH)). After incubation of the PM
152 suspension within the lining fluid with DTT, the remaining DTT was titrated with 5,5'-dithiobis-(2-nitrobenzoic
153 acid) (DTNB) to form 5-mercaptop-2-nitrobenzoic acid (TNB). Absorbance at 412 nm (TECAN Infinite M200 Pro)
154 in 96-well plates provided the concentration of unconsumed DTT, from which the DTT consumption rate
155 (nmol min⁻¹) was calculated. The batches were standardized using common ~~external references control (lab'~~
156 ~~rooftop filter analysis for every batch)~~ to ensure consistency between batches.

157 After analysis, the OP activities were blank subtracted and then normalized using the PM₁₀ mass concentration
158 and the sampling air volumes. The mass-normalized OP (OP^m, nmol min⁻¹ µg⁻¹) represents the intrinsic OP of 1 µg
159 PM, while the volume-normalized OP (OP^v, nmol min⁻¹ m⁻³) represents PM-derived OP per m³ of air. Each sample
160 is analyzed in triplicate for AA and triplicate for DTT, respectively. Consequently, the OP values presented in the
161 study are the mean and the standard deviation of these replicates.

162 **2.2.3. Vertical temperature and other ancillary measurements**

163 Vertical temperature and humidity were measured every 30 minutes from November 2017 to May 2023 using
164 Tinytag TGP-4500 from Gemini Data Loggers. A Stevenson Type Screen protects each Tinytag loggers from
165 radiant heat (direct sunlight). Sensors are installed at a minimum of 3m from the ground. The measurements have
166 been performed at different elevations of the Bastille hill, located a few hundred meters from the city center
167 (5°43'37.0"E, 45°11'40.8"N), including z = 230, 309, 496, 916m altitudes.

168 Further, measurement of the PM₁₀ mass was conducted (hourly) using tapered element oscillating microbalances
169 equipped with filter dynamics measurement systems (TEOM-FDMS) at the same site as the filter collection. The
170 PM concentration used in this study is the 24-hour average concentration, which is associated with the days of
171 filter-based sample measurement (from 02/01/2013 to 28/05/2023).

172

173 **2.3. Multivariate imputation by chained equations (MICE)**

174 The daily concentration of metals was only accessed in some periods, with the number of samples being 362 of
175 the total of 1570 samples, which would severely limit the size of the inputs for the PMF processing. We used the
176 weekly concentration measured over the whole period to estimate the missing daily data using an imputation
177 method. The daily concentration of metals was imputed by using the MICE algorithm implemented with
178 multilinear regression (Azur et al., 2011). These values were modeled conditionally depending on the observed
179 values of the daily PM₁₀ and PM₁₀ components concentration (i.e., weekly concentration, PM₁₀, and PM₁₀
180 components concentration). These components are OC, EC, MSA, Levoglucosan, Mannosan, Polyols, NO₃⁻, SO₄²⁻
181 , Na⁺, NH₄⁺, K⁺, Mg²⁺, Cl⁻, Ca²⁺. The data preparation and imputation processes are implemented through 4 main
182 steps, as presented in S1 and Figure S1, Supplement. The validation of this imputation is shown in Table S1 and
183 Figure S2.

184 **2.4. Persistent inversions detection**

185 Thermal inversion occurs when the vertical temperature gradient between the ground-based and higher-altitude
186 stations is positive. However, this constraint is restrictive and limits thermal inversion detection, especially when
187 the calculation is on average daily temperature (Largeron and Staquet, 2016). Hence, the persistent inversion is
188 detected, as discussed in Largeron and Staquet (2016), for days with :

$$189 \text{average} \left(\frac{T_{916} - T_{230}}{\Delta z} \right)_{\text{Daily}} > \text{Mean} \left(\frac{T_{916} - T_{230}}{\Delta z} \right)_{\text{Winter}} \quad (1)$$

190 for more than 72 consecutive hours

191 with:

192 $T_{916} - T_{230}$ is the difference between temperature at ground-base station ($z = 230$ m altitude) and at high-elevation
193 station ($z = 916$ m);

194 Δz is the difference between the height of high and low elevations;

195 $\frac{T_{916} - T_{230}}{\Delta z}$: is the bulk temperature gradient between $z = 230$ and $z = 916$ m;

196 $\text{Mean} \left(\frac{T_{916} - T_{230}}{\Delta z} \right)_{\text{Winter}}$: is the mean bulk temperature gradient in wintertime (from November to March).

197 **2.5. Positive Matrix Factorisation (PMF)**

198 **2.5.1. PMF input**

199 EPA PMF 5.0 (Gary Norris et al., 2014) was used to identify and quantify the PM₁₀ sources based on the observed
200 concentrations and their related uncertainties. The concept of PMF is to use the weighted least square fit method
201 and apply a non-negative constraint with the weight calculated based on analysis uncertainties (Paatero and
202 Tappert, 1994) (Eq. (S1), Supplement S2). In this study, the input matrix of the PMF comprises 25 chemical
203 species, including PM₁₀ (set as the total variable), carbonaceous fractions (OC*, EC), ions (Cl⁻, NO₃⁻, SO₄²⁻, Na⁺,

204 NH_4^+ , K^+ , Mg^{2+} , Ca^{2+}), organic tracers (MSA, levoglucosan, mannosan, polyols) and trace metals (As, Ba, Cd, 205 Cr, Cu, Ni, Pb, Sb, V, Zn). The trace metals were the daily measured metals in some periods (2013, 2017-2018, 206 2020-2021) and the daily imputed metals. The OC^* (=OC minus the sum of the carbon mass from the organic 207 tracers used in the input variables) was used to avoid considering twice the mass of C atoms in organic markers. 208 Polyols were calculated as the sum of arabitol and mannitol, supposing that their origin is similar (Samaké et al., 209 2019a). The input uncertainties were calculated based on the concentrations and the uncertainties in the analysis 210 (Gianini et al., 2012; Waked et al., 2014). Details on the calculation of OC^* and uncertainties of PMF input are 211 presented in Section S3, Supplement. The selection of the input variables is guided by our previous yearly PMF 212 studies at this site (Borlaza et al., 2021; Srivastava et al., 2018; Weber et al., 2019).

213 **2.5.2. Set of constraints**

214 The application of PMF constraints is recommended in the European guide on air pollution source apportionment 215 with receptor models (Belis et al., 2014) to avoid mixing between some factors and reduce the uncertainty of the 216 rotational ambiguity. The constraints used in this study are also based on the previous PMF studies in Grenoble 217 (Borlaza et al., 2021; Srivastava et al., 2018; Weber et al., 2019) and are detailed in Table S3.

218 **2.5.3. Choice of the final PMF solution**

219 Several solutions, including those from 4 to 11 factors, were investigated to determine the optimal output. This 220 selection is based on the ratio of $Q_{\text{true}} / Q_{\text{robust}}$ (evaluating the outlier's effect), the clarity of the chemical profile, 221 the contribution of factors to PM_{10} , the correlation between measured and predicted concentration, and the stability 222 of the solution. This stability was evaluated using the bootstrapping (BS) and displacement (DISP) methods. BS 223 analysis randomly resamples the data observation matrix and uses it to run a new PMF. The base-run and boot- 224 run factors are matched if their correlation exceeds the threshold (generally chosen at 0.6). DISP estimates each 225 species' uncertainty in the factor profile by fitting the model many times until this variable turns displaced (upper 226 or lower) from its fitted value. The details of the set criteria for validation are presented in S4.

227 To evaluate the stability of the PMF solution over time (including possible changes in the chemical profiles of the 228 sources), we also implemented separated PMF SA for every successive period of 3 years (2013-2016, 2017-2020, 229 2021-2023) and then we investigated the homogeneity of the chemical profiles by using the Pearson distance (PD) 230 and standardized identity distance (SID) metrics (Belis et al., 2015). The chemical profiles of PMF solutions every 231 3 years and 11 years, and those published in Borlaza et al. (2021) are compared to assess the homogeneity of the 232 chemical profiles.

233 **2.6. Regression techniques for PM_{10} OP SA**

234 The regression technique is applied to apportion OP^v (AA, DTT) and PMF-derived PM_{10} sources' contribution, as 235 expressed in Eq.2. Principally, OP^v ($\text{nmol min}^{-1} \text{m}^{-3}$) is treated as a dependent variable, and PMF-derived PM_{10} 236 sources' contribution ($\mu\text{g m}^{-3}$) are independent variables. The OP SA methodology in this study follows the 237 methodology reported by Ngoc Thuy et al. (2024).

$$238 \quad OP_v = \sum_p^{i=1} OP_m^i * PM^i + e \quad (2)$$

239 Where:

240 OP_v is the volume-normalized OP (nmol min⁻¹ m⁻³)
 241 p is the number of PMF-derived PM₁₀ sources
 242 OP_m^i is the regression slope, denoted as the intrinsic OP of source i (nmol min⁻¹ µg⁻¹)
 243 PM^i is the contribution of source i to PM₁₀ (µg m⁻³)
 244 e is the residual of the regression technique (nmol min⁻¹ m⁻³)
 245 The appropriate regression tool is selected based on the collinearity among independent variables and the variance
 246 of regression residuals (Ngoc Thuy et al., 2024). The collinearity among PMF-derived sources was tested using
 247 the variance inflation factor (VIF), which is calculated using Eq. (S3) in Supplement S2 (Craney and Surles, 2002;
 248 O'Brien, 2007; Rosenblad, 2011). The variance of the regression residual is assessed using the Goldfeld-Quandt
 249 test (Goldfeld and Quandt, 1965) to investigate if the regression residual varies by the value of the dependent
 250 variable (OP^v). The most appropriate regression method is selected among a wide choice of possible tools
 251 (including ordinary least square, weighted least square, positive least square, Ridge, Lasso, random forest, and
 252 multiple layer perceptron), following the methodology developed by Ngoc Thuy et al. (2024). It is performed with
 253 considering the characteristics of the data and comparing the accuracy metrics (R-square, root mean square error,
 254 and mean absolute error) for each of them. For instance, if the regression residual is constant (homoscedasticity),
 255 the model ordinary least square (OLS) and Positive least square (PLS) are satisfactory. On the other hand, if the
 256 regression residual varied with the dependent variable (heteroscedasticity), the models incorporating some sort of
 257 weighting are chosen (including weighted least squares (WLS) and weighted positive least squares (wPLS)),
 258 where the weighting is the standard deviation of replicated OP analyses.
 259 The most appropriate model was trained by randomly choosing 80% of the dataset and validated with the
 260 remaining 20%. The model was run 500 times to ensure the robustness of the results, especially considering the
 261 remarkable seasonality of many components in the dataset. The contribution to OP of each source is calculated
 262 by multiplying its contribution to PM₁₀ with the arithmetic mean intrinsic OP (or regression slope) of the 500
 263 iterations.

264 **2.7. Seasonal-trend using LOESS decomposition**

265 Seasonal-trend decomposition using LOESS (SLT) was developed by RB Cleveland et al. (1990) and is a robust
 266 method for decomposing time series into trends, seasonality, and residuals. This method uses LOESS, a method
 267 for estimating the non-linear relationships to decompose a time series. In our case, we used monthly average
 268 concentration as input data in order to have a more robust data set, smoothing high variability noise. The trend
 269 component is first calculated by applying a convolution filter to the data. Then, this trend is removed from the
 270 series. Finally, the average of this detrended in each period is the seasonal component. The residuals can be
 271 explained neither by trend nor by season. The STL is an iterative model that uses LOESS to smooth seasonal and
 272 trend components to obtain the minimum residuals. Further, in STL decomposition, the outliers in the data are
 273 given less weight in the estimation of trend and season. The STL model is described in the equation below:

$$274 \quad y_t = S_t + T_t + R_t \quad (t = 1, 2, \dots, n) \quad (3)$$

275 where, in our case, y_t is the monthly contribution of PMF-derived sources, S_t is the seasonal component, T_t is the
 276 trend component, and R_t denotes the residual component. The seasonal frequency was chosen 6 months before
 277 and 6 months after the evaluated month (seasonal frequency = 13 months) to estimate the yearly trend cycle.

278 Hence, the first and last 6 months of the decomposition time series were removed from the results to prevent edge
279 effects.

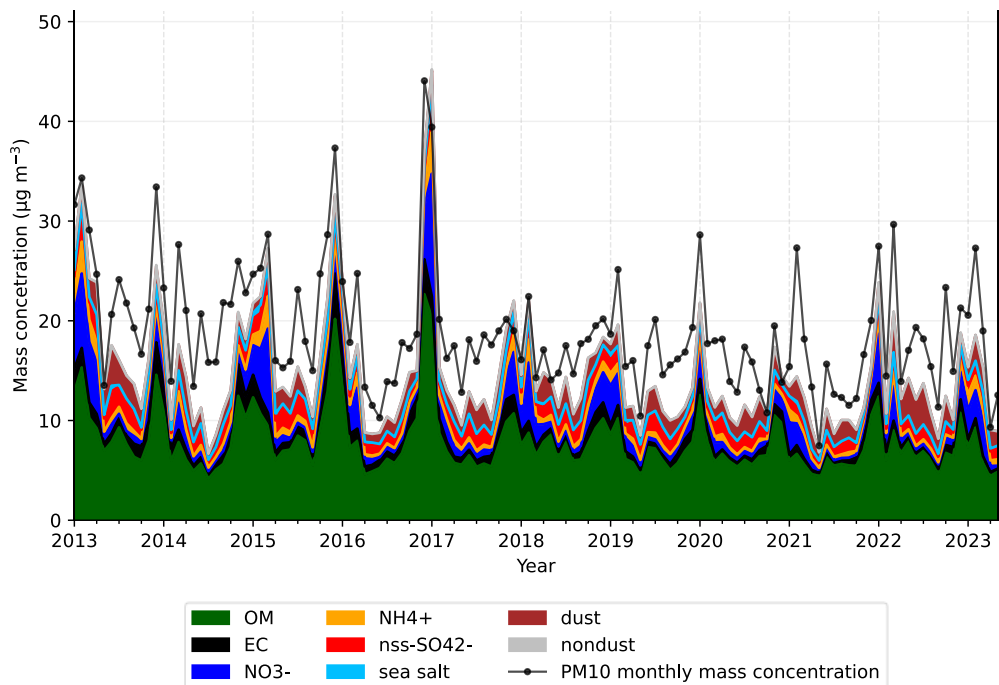
280 The long-term trend of PM_{10} sources was accessed by applying the STL model to the monthly contribution of
281 sources to PM_{10} (output of PMF). The fit line of the trend was assessed by using ordinary least squares linear
282 (OLS). The annual rate change of the trend is the slope of the fit line multiplied by 12 months ($\mu\text{g m}^{-3} \text{yr}^{-1} / \text{nmol}$
283 $\text{min}^{-1} \mu\text{g}^{-1} \text{yr}^{-1}$). The STL decomposition and the fit line of the trend were performed in Python 3.9 using the
284 package "statsmodels" (Seabold and Perktold, 2010).

285 **3. Results and discussion**

286 **3.1. Evolution of PM_{10} concentration and chemical components**

287 The annual average concentration of PM_{10} , considering all available daily measurements, is $19.0 \pm 10.6 \mu\text{g m}^{-3}$ for
288 the whole studied period (2013-2023). The highest annual concentration is observed in 2013 ($24.4 \pm 13.7 \mu\text{g m}^{-3}$),
289 and the lowest is in 2021 ($15.3 \pm 9.8 \mu\text{g m}^{-3}$). The number of days with concentrations surpassing the European
290 standard daily thresholds ($40 \mu\text{g m}^{-3}$) is 176 days in 11 years, representing 4.6% of the total observed days, which
291 are principally found in the cold season (Nov, Dec, Jan, Feb, Mar).

292 The PM_{10} main components are organic matter (assuming $\text{OM} = 1.8 * \text{OC}$ (Favez et al., 2010)), representing on
293 average over the overall period $41.3 \pm 8.0\%$ of PM_{10} mass concentration, followed by dust ($9.6 \pm 4.4\%$), nitrate
294 (NO_3^- , $7.5 \pm 6.2\%$), non-sea salt sulfate (nss-SO_4^{2-} , $7.4 \pm 2.4\%$), elemental carbon (EC, $5.5 \pm 2.5\%$), ammonium
295 (NH_4^+ , $3.9 \pm 2.0\%$), sea salt (Na^+ and Cl^- , $1.7 \pm 0.8\%$) and other non-dust elements (Cu, Pb, V, Zn, representing
296 $0.2 \pm 0.1\%$). These main composition fractions are estimated using the formula as shown in S2, Eq. (S4). The
297 monthly evolutions of PM_{10} and its main chemical components for the whole period are shown in [Figure 2](#).
298 The maximum concentration of PM_{10} was observed in winter months (December, January, and February),
299 corresponding to the highest concentration of OM and EC ($7.82 \pm 3.11 \mu\text{g m}^{-3}$ and $1.09 \pm 0.74 \mu\text{g m}^{-3}$, respectively).
300 Nitrate concentrations are higher in the middle of winter and the early spring, corresponding also with the high
301 concentrations of ammonium (1.63 ± 1.87 and $0.78 \pm 0.62 \mu\text{g m}^{-3}$). The agricultural activities (especially manure
302 spreading) could explain this high contribution in spring under humidity and temperature conditions favoring the
303 condensation of ammonium nitrate in the particulate phase. Nss-sulfate concentrations are more abundant in the
304 warmer season (summer), where the photochemical production is favorable. No clear seasonal pattern could be
305 observed for other components (sea salt, dust, non-dust, estimated as described in section S2), suggesting that the
306 emissions of these components are stable for the whole year. At first glance, decreasing trends appear visible for
307 PM_{10} and OM, EC, NO_3^- , NH_4^+ , and non-dust components, while sea salt, dust, and nss-SO_4^{2-} do not seem to
308 present significant trends. With chemical components coming from several emission sources, an advanced
309 analysis, including a PMF model followed by an STL decomposition, was performed to assess the trend of PM_{10}
310 sources. The result of the PMF model is presented in section 3.2, and the tendencies of PM_{10} sources and OP are
311 shown in sections 3.3 and 3.4, respectively.



312
313 **Figure 2. The average monthly evolution of PM₁₀ and its main components from 2013 to 2023. The line represents the**
314 **monthly average concentration of PM₁₀ measured by TEOM-FDMS.**

315 **3.2. PM₁₀ sources apportionment**

316 **3.2.1. PMF chemical profiles**

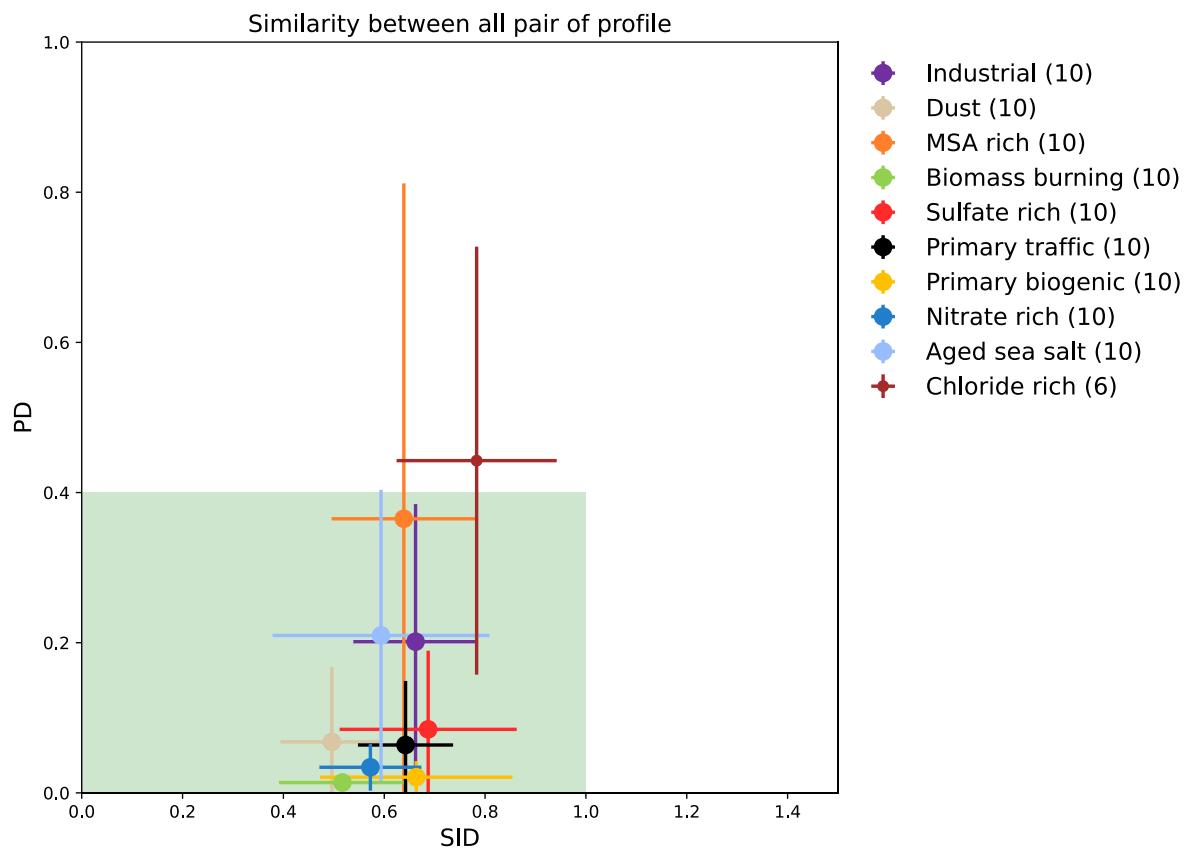
317 Using a unique chemical profile for each of the sources for such a long-term period can potentially limit the
318 assessment of its evolution (Borlaza et al., 2022a). To evaluate such a phenomenon in our case, we investigated
319 the chemical profile and contribution of PM₁₀ sources for three distinct periods (2013-2016, 2017-20240, 20212-
320 2023) and compared the results with those for the full 11-year period, as well as to the results presented in (Borlaza
321 et al. (2021) for the year 2017. Particularly, we checked the similarity of the chemical profiles of these PMF
322 solutions using PD and SID metrics (Belis et al., 2015).

323 For each SA, the PMF solution was tested from 4 to 11 factors and validated by the criteria presented in section
324 S4. The results of these validations (Q_{true}/Q_{robust} , bootstrap run, displacement run, and statistical validation) are
325 presented in S5, Tables S4, S5 and S6. The runs of 4 to 9 factors returned at least one merging factor, and the
326 solution with 11 factors led to a factor without geochemical identity. Finally, for each PMF tested (11 years, 2013-
327 2016, 2017-2021, 2022-2023), the best solution includes 10 PM₁₀ sources, with mineral dust, sulfate-rich, primary
328 traffic, biomass burning, primary biogenic, nitrate-rich, MSA-rich, aged sea salt, industrial, and chloride-rich.

329 The similarity of the chemical profiles is presented in [Figure 3](#). Most of the factors (i.e., aged sea salt,
330 mineral dust, primary biogenic, biomass burning, primary traffic, industrial, nitrate-rich, and sulfate-rich) present
331 quite homogenous chemical profiles over the 3 successive periods, indicating that these source profiles are quite
332 stable during the full 11-year period and similar compared to sources reported in Borlaza et al. (2021). The MSA-
333 rich and chloride-rich sources are the most divergent but are still within the limit of the accepted PD and SID
334 range; however, their standard deviations for PD are slightly higher than for the other sources ([Figure 3](#)).
335 This is due to differences in the contributions of SO₄²⁻ in the chemical profile of MSA-rich, which varied from 6
336 % to 17%, and that of Cl⁻ (73% - 83%) in the chloride-rich factor. In a previous study, Weber et al. (2019) also

337 reported that the proportion of SO_4^{2-} in the MSA-rich source can significantly vary across French sites, from 6%
 338 to 24%. The chloride-rich source in our study (previously named sea/roadsalt in Borlaza et al. (2021) is essentially
 339 composed of a high proportion of Cl^- , with less than 10% of Na^+ and some metals (Cu, Mn, Ni, V). This source is
 340 detected in other alpine valley environments (Glojek et al., 2024), with a similar temporal evolution as here. Since
 341 chloride depletion from the particulate phase can greatly depend on solar radiation, relative humidity, and
 342 temperature, the chemical profile of this factor can vary on different time scales. This source was also observed
 343 to be heterogeneous in the three neighboring sites investigated within 15 km in the previous study in Grenoble
 344 (Borlaza et al., 2021). Nevertheless, it should be noted that it represents only a very minor fraction of the PM_{10}
 345 total mass (about 1%).

346 With these stabilities of the chemical profiles over the years, the solution for the 11-year SA is considered suitable
 347 for further data analyses in this paper. In the next section (3.3.2), we investigate how the contribution of these
 348 sources to total PM_{10} loadings changed over time.

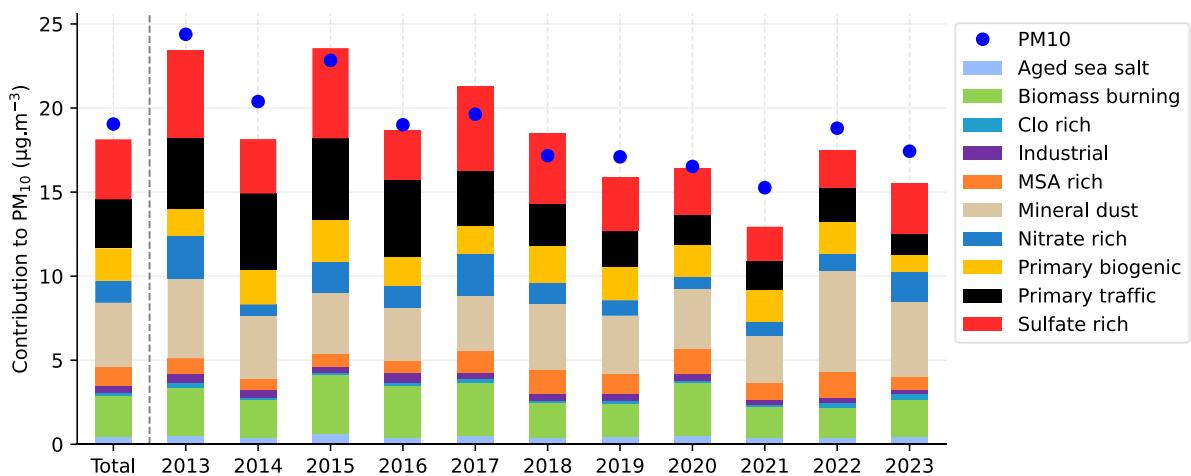


349
 350 **Figure 3.** Similarity plots of the chemical profiles of the solution for the 11-year SA against the 3 SA solutions every 3
 351 years, and those presented by Borlaza et al. (2021). The shaded area (in green) shows the limit of the homogeneous
 352 chemical profile. For each point, the error bars represent the standard deviation when comparing all pairs of SA
 353 solutions (number of pairs in parentheses in the legend).

354 **3.2.2. Variations of the source's contribution in the 11-year PMF SA**

355 As presented in [Figure 4](#), the optimal PMF solution for the 11-year time series identifies 10 PM_{10} sources,
 356 with the contributions of mineral dust (20.9%), sulfate-rich (19.7%), traffic (16.0%), biomass burning (13.5%),
 357 primary biogenic (10.7%), nitrate-rich (7.2%), MSA-rich (6.2%), industrial (2.2%), aged sea salt (2.5 %), and
 358 chloride-rich (1.0%). The chemical profile and contribution of each source are shown in Figures S3 and S4,
 359 respectively. Even though the chemical profiles are homogenous, the contributions of these sources show minor

360 differences from those reported for this same site by Borlaza et al. (2021) and Srivastava et al. (2018), partly
 361 because of the differences in the respective periods of the studies. However, the main sources are similar, i.e., SIA
 362 (nitrate and sulfate-rich), mineral dust, biomass burning, and primary traffic. Similar general results are also
 363 presented for Swiss Alpine (Ducret-stich and Tsai, 2013), French Alpine (Weber et al., 2018), and Slovenian
 364 Alpine areas (Glojek et al., 2024), showing biomass burning and secondary inorganic aerosols being the most
 365 abundant contributions to PM mass. Primary biogenic and MSA-rich sources are the biogenic sources rarely
 366 reported in the literature; however, they account together for 17% of total PM₁₀ mass on average in our study,
 367 which is in line with those reported in urban background sites in France (Samaké et al., 2019b; Weber et al., 2019).
 368 The absolute PM₁₀ source contributions are also compared to the average annual concentration of PM₁₀ mass to
 369 demonstrate the ability of the PMF model to reconstruct the PM₁₀ mass. The difference between observed and
 370 reconstructed PM₁₀ concentrations on the 11-year average is about 1 $\mu\text{g m}^{-3}$ (5 %), with no more than 2 $\mu\text{g m}^{-3}$ for
 371 any single year, demonstrating that the PMF model performs well at reconstructing the PM₁₀ concentrations.



372
 373 **Figure 4. The absolute average contribution of sources to PM₁₀ for every year and the 11 years (total), and the**
 374 **concentration of PM₁₀ (blue circle).**

375 Significant trends in source contributions over this 11-year period are detected (and discussed in section 3.3);
 376 nevertheless, the main contributors to the total PM₁₀ mass do not change, with mineral dust, biomass burning,
 377 sulphate-rich, nitrate-rich, and primary traffic being the main contributors to PM₁₀. The highest PM₁₀
 378 concentrations (observed in winter/spring 2013 and 2015) are associated with the highest contribution of SIA and
 379 biomass burning sources. On the other hand, the relative contribution of SIA and biomass burning showed a
 380 negligible difference (varied from 0.3 to 4%) between these years compared to 2014 and 2016 (Figure S5). The
 381 lowest PM₁₀ annual concentration was detected in 2021, notably when the third COVID-19 pandemic lockdown
 382 restrictions applied in France. In addition, the relative contributions (see Figure S5) showed only small changes
 383 compared to those in other years, with an increasing contribution of primary biogenic sources in 2021 (4%
 384 compared to 2020), and only a very light decrease in the anthropogenic sources.
 385 The decrease in PM₁₀ annual average concentrations observed since 2017 is associated with decreases in the
 386 contribution of some of the anthropogenic PM₁₀ sources. However, using yearly averages for trend analysis may
 387 prevent a proper understanding of the variation in time and of the estimation of the trends based on monthly
 388 averages, which might be more informative, as discussed in section 3.3.

389 **3.3. Trends in sources' contributions**

390 **3.3.1. Mean rate change in the contribution of PM₁₀ sources**

391 The source contribution trend analysis was achieved through STL deconvolution (see section 2.6). These trends
392 for all sources over the full period of the study are presented in Table 1~~Table 1~~. In this table, the part labeled
393 "Rest" represents the difference between the total PM₁₀ measured mass and the sum of the mass of all PMF-
394 derived factors in order to assess any trend of the unresolved part of PM₁₀ within our SA study.

395 PM₁₀ concentrations present a downward trend from 2013 to 2023, with an average diminution of 0.73 $\mu\text{g m}^{-3} \text{yr}^{-1}$ (3.9%) (S6, Figure S6). Such a downward trend of PM₁₀ in Grenoble is in line with that observed in other urban
396 sites in Europe (Aas et al., 2024; Borlaza et al., 2022a; Caporale et al., 2021; Colette et al., 2021; Gama et al.,
397 2018; Li et al., 2018; Pandolfi et al., 2016). ~~The reduction of PM₁₀ in Grenoble during this period is significantly
398 larger than that in 30 rural sites of the European Monitoring and Evaluation Programme (EMEP) from 2000 to
400 2017, which show reductions of PM₁₀ from -0.008 to -0.58 $\mu\text{g m}^{-3}$ (-1.5% to -2.5%). However, the results of our
401 study are highly coherent with results from Aas et al., presenting a reduction of PM₁₀ in 2 rural sites in France
402 (La Tardière and Revin) of -3.5% yr^{-1} between 2005 and 2019. Indeed, France is amongst the EU countries with
403 the highest reduction trend, as presented by Aas et al..~~

404

405 The reduction of PM₁₀ in Grenoble during this period is significantly larger than that in 30 rural sites of the
406 European Monitoring and Evaluation Programme (EMEP) from 2000 to 2017, which show reductions of PM₁₀
407 from -1.5% to -2.5% (-0.008 to -0.58 $\mu\text{g m}^{-3}$) (Colette et al., 2021). However, the results of our study are highly
408 coherent with results from Aas et al. (2024), presenting a reduction of PM₁₀ in 2 rural sites in France (La Tardière
409 and Revin) of -3.5% yr^{-1} between 2005 and 2019. The reduction of PM in this Grenoble site, as an urban site,
410 being higher than those at the rural sites, is due to the changes in specific emission activities at the site. While in
411 the rural sites, the PM emission are influenced by long range transport activities, the PM at the urban site is usually
412 largely impacted by different local activities (Borlaza et al., 2022b). Further, France is amongst the EU countries
413 with the highest reduction trend, as presented by Aas et al. (2024).

414 ~~The anthropogenic sources, such as primary traffic, sulfate rich, and biomass burning, display the highest decrease~~
415 ~~between 2013 and 2023 in Grenoble, with a reduction of 0.37, 0.25, and 0.13 $\mu\text{g m}^{-3} \text{yr}^{-1}$ (12.9, 6.9, and 5.5%),~~
416 ~~respectively. The other anthropogenic sources also have a significant decreasing trend; however, they are much~~
417 ~~lower (nitrate rich: -0.11 $\mu\text{g m}^{-3} \text{yr}^{-1}$, industrial: -0.02 $\mu\text{g m}^{-3} \text{yr}^{-1}$). The downward trends of these anthropogenic~~
418 ~~sources (mainly traffic, SIA, and industrial) were also underlined for other European urban sites (Colette et al.,~~
419 ~~2021; Diapouli et al., 2017; Pandolfi et al., 2016). For instance, a similar approach was followed by Pandolfi et~~
420 ~~al. (2016), investigating the Mann Kendall trend of PMF-derived sources, and reported an almost equivalent~~
421 ~~downward trend of the sulfate rich factor of -0.32 $\mu\text{g m}^{-3} \text{yr}^{-1}$ between 2004 and 2014 in Spain. The decreasing~~
422 ~~trends of primary traffic, domestic biomass burning, and industrial emissions are potentially influenced by the~~
423 ~~reduction in primary emissions due to various abatement strategies (as discussed in the following subsections,~~
424 ~~notably in 3.3.3 and 3.3.4).~~

425 The anthropogenic sources, such as primary traffic, sulfate-rich, and biomass burning, display the highest decrease
426 between 2013 and 2023 in Grenoble, with a reduction of 12.9, 6.9, and 5.5% (0.37, 0.25, and 0.13 $\mu\text{g m}^{-3} \text{yr}^{-1}$),
427 respectively. The other anthropogenic sources also present significant decreasing trends; however, these trends
428 are much lower (nitrate-rich: -0.11 $\mu\text{g m}^{-3} \text{yr}^{-1}$, industrial: -0.02 $\mu\text{g m}^{-3} \text{yr}^{-1}$). The downward trends of these

429 anthropogenic sources (mainly traffic, SIA, and industrial) were also underlined for other European urban sites
 430 (Colette et al., 2021; Diapouli et al., 2017; Pandolfi et al., 2016) with various approaches. For instance, a similar
 431 approach using PMF (albeit without organic markers) was followed by Pandolfi et al. (2016), investigating the
 432 Mann-Kendall trend of PMF-derived sources, and reported an almost equivalent downward trend of the sulfate-
 433 rich factor of 53% (i.e., 0.53 yr^{-1}) between 2004 and 2014 in Spain. The decreasing trends of primary traffic,
 434 domestic biomass burning, and industrial emissions are potentially influenced by the reduction in primary
 435 emissions due to various abatement strategies (as discussed in the following subsections, notably in 3.3.3 and
 436 3.3.4).

437 Conversely, natural sources such as mineral dust and chloride-rich factors do not show any significant trend or
 438 follow a very weak one (aged sea salt, primary biogenic). MSA-rich is the only source that displays a significant
 439 upward trend, with an increase of $0.08 \mu\text{g m}^{-3} \text{ yr}^{-1}$; further studies would be needed to relate this last increase to
 440 changes in precursor emissions or reactivity during transport. Finally, the low evolutions in the contributions of
 441 the natural sources demonstrate that the reduction in PM_{10} in Grenoble is essentially related to the reduction of
 442 anthropogenic activities, especially sources related to traffic and domestic biomass burning activities.

443
 444 **Table 1. Trend of PM_{10} sources and PM_{10} (in $\mu\text{g m}^{-3} \text{ yr}^{-1}$ and $\% \text{ yr}^{-1}$).**

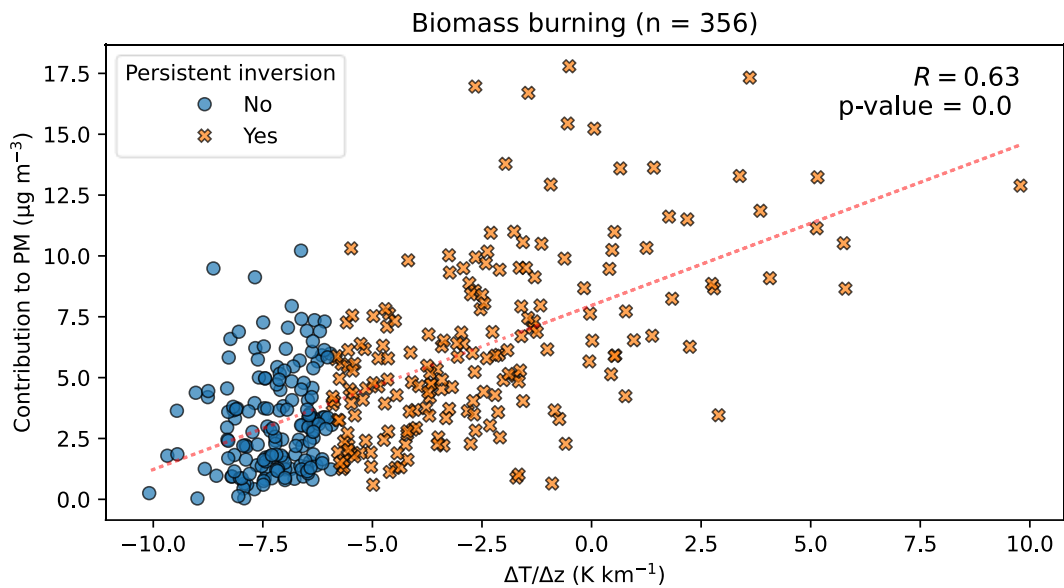
	Absolute trend ($\mu\text{g m}^{-3} \text{ yr}^{-1}$)	Relative trend ($\% \text{ yr}^{-1}$)	P-values	R^2
Aged sea salt	-0.01	-2.50	<<0.01	0.22
Biomass burning	-0.13	-5.48	<<0.01	0.98
Chloride rich	0.00	1.18	0.01	0.07
Industrial	-0.02	-5.36	<<0.01	0.40
MSA rich	0.08	6.63	<<0.01	0.64
Mineral dust	0.04	1.03	0.02	0.05
Nitrate rich	-0.11	-8.08	<<0.01	0.94
Primary biogenic	-0.01	-0.49	0.03	0.04
Primary traffic	-0.37	-12.85	<<0.01	0.94
Sulfate rich	-0.25	-6.89	<<0.01	0.70
PM_{10}	-0.73	-3.89	<<0.01	0.68
Rest	-0.11	-2.13	<<0.01	0.39

445
 446 **3.3.2. Potential influence of meteorology**

447 The STL deconvolution is inherently constructed to separate the yearly and seasonal variations from the long-
 448 term trends. While we discuss the long-term trends of the sources in other sections (3.3.1, 3.3.3, and 3.3.4), it is
 449 also interesting to evaluate the impact of the meteorology on the seasonal variations of the concentrations. It is
 450 well known that inversion layers in the lower atmosphere are extremely important for the modulation of the
 451 concentrations at the ground, particularly in the context of Alpine valleys during winter (Carbone et al., 2010;
 452 Glojek et al., 2022). In this section, we tried to better evaluate these impacts on the concentrations from the sources
 453 of PM in the case of our time series.

454 This was considered with the measurements of temperature along the slopes of the mountains very close to the
 455 city center (as described in section 2.2.3), for the winter periods of 2017-2023. It has been previously shown by
 456 Allard et al. (2019) that such measurements are representative of the temperature in the valley, despite the potential

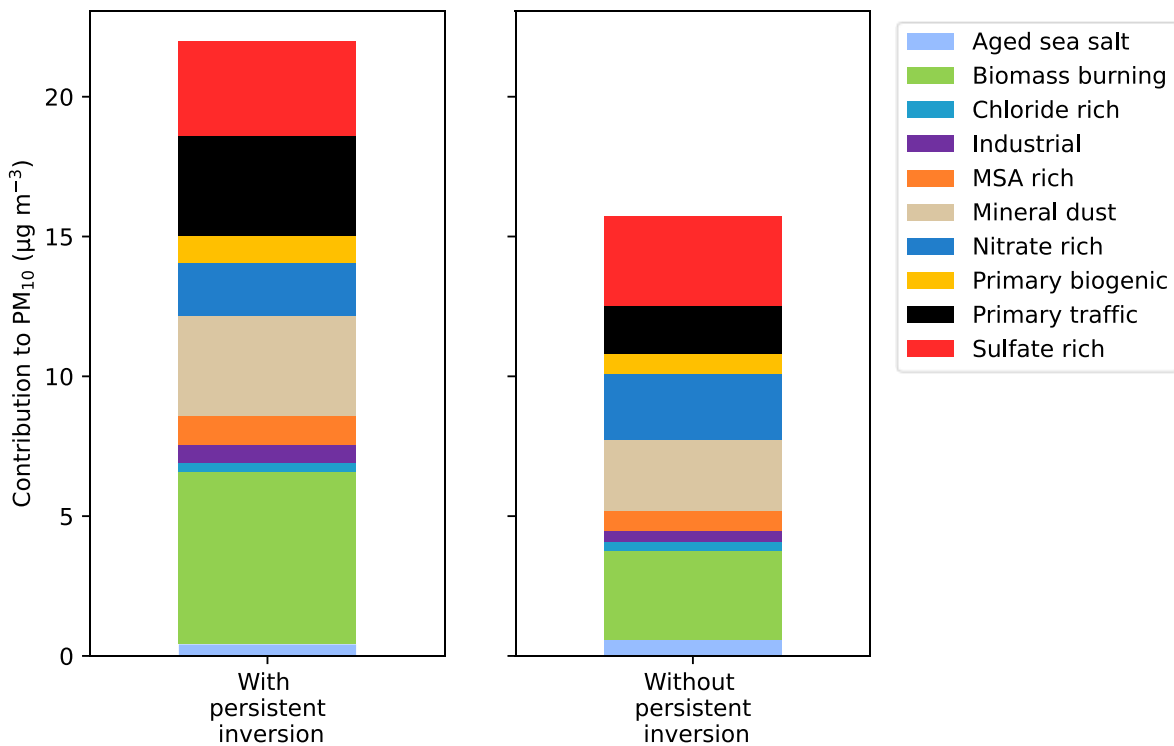
457 influence of wind slopes. We particularly considered the temperature gradient over the first 700 m above ground
 458 and the number of days with persistent inversion, as defined in section 2.2.3.
 459 The analysis of the relationship between the PM₁₀ and bulk temperature vertical gradients ($\Delta T/\Delta z$) in winter (Nov,
 460 Dec, Jan, Feb, Mar), summer (May, June, Jul, Aug), and transition season (remaining months) reveals that thermal
 461 inversion events and high PM₁₀ concentration are mainly occurring in winter time (Supplement S7, Figure S8)
 462 during the 5 years of the study. Periods of persistent temperature inversion were assessed based on the condition
 463 in Eq. 1, which detected 79 persistent inversion days in series from 4 to 22 consecutive days, for the winter periods
 464 2017-2023. A meaningful correlation is obtained between PM₁₀ concentrations and bulk temperature vertical
 465 gradient (r reaching 0.60, p<<0.001) for these winter months and even better when considering only the persistent
 466 inversion periods (r reaching 0.67, p<<0.001) for individual years (Table S7).



467
 468 **Figure 5.** Daily concentrations of biomass burning to PM₁₀ and daily temperature gradients ($\Delta T/\Delta z$) during the
 469 winter periods (from November to March) of 2017-2023. The dotted red line is the linear regression fit. The blue
 470 circle symbols denote days when persistent inversion does not occur, and the orange multiple symbol denotes days
 471 when persistent inversion occurs.

472
 473 The distribution between the daily PM₁₀ concentration and daily average $\Delta T/\Delta z$ in winter months revealed that
 474 the majority of PM₁₀ concentration peaks (in excess of $40 \mu\text{g m}^{-3}$) occur during the persistent inversion days
 475 (Figure S9). However, it also shows that a few high PM₁₀ concentrations could be found on the days without
 476 persistent inversion; meanwhile, the days with persistent inversion do not always have high PM₁₀ concentrations.
 477 This result is not surprising since the concentration of PM₁₀ is not only associated with thermal inversion events
 478 but also depends on other meteorological conditions (precipitation, heat deficit) and the variation of pollutant
 479 emissions (Carbone et al., 2010; Largeron and Staquet, 2016).
 480 Interestingly, the impact of persistent inversion days on PM₁₀ concentrations from the residential biomass burning
 481 source is larger than that for other sources or total PM₁₀ (Figure 5), with a higher correlation (0.63). In addition,
 482 the contribution of this source is systematically lower during non-inversion days, and large concentrations are
 483 essentially made during persistent days. The large impact of the inversions on the local sources is confirmed when
 484 comparing the source contribution of the inversion days vs non-inversion days (Figure 6). This figure
 485 shows both the large increase in average PM₁₀ concentrations and also the contributions of the local sources

486 (emissions from residential biomass burning, traffic, industries, mineral dust probably from resuspension) in the
 487 cases of inversion days during winter. Conversely, long-range transport sources (sulfate-rich, nitrate-rich) tend to
 488 be less important during these inversion days. A similar pattern is observed for the relative contribution of sources
 489 to PM (Figure S.10), in which the significant contribution of biomass burning, dust, industrial, and primary traffic
 490 is detected during inversion events. The trends of the two most important local anthropogenic sources (domestic
 491 biomass burning and traffic) are further discussed in the next sections.



492
 493 **Figure 6. Contribution of the different sources to the PM₁₀ composition for days with persistent inversion vs non-**
 494 **inversion days of the winters 2017-2023.**

495 3.3.3. Trend in biomass burning contributions

496 The trend of the domestic biomass burning PM₁₀ concentrations is investigated via an STL decomposition analysis
 497 on this PMF-derived source (Figure 7), indicating a statistically significant decreasing trend from 2013
 498 to 2023 (p-values <<0.01). The seasonal estimate shows the highest values in the winter season (Nov, Dec, Jan),
 499 with a visual trend to a smoothing of the peak concentrations; conversely, from Mar to Sept, the seasonal variations
 500 showed constantly lowest values. Extreme residual values were detected in the winter months of 2016, 2017, and
 501 2021, explained by high-concentration episodes of PM₁₀, where the concentration exceeded the European standard
 502 for PM₁₀ concentration in 24 hours (PM₁₀ concentration varied from 50 to 78 µg m⁻³). The linear fit line of the
 503 trend is highly significant with R² = 0.97, with a reduction of 134 ng m⁻³ yr⁻¹ (-5.5% yr⁻¹).

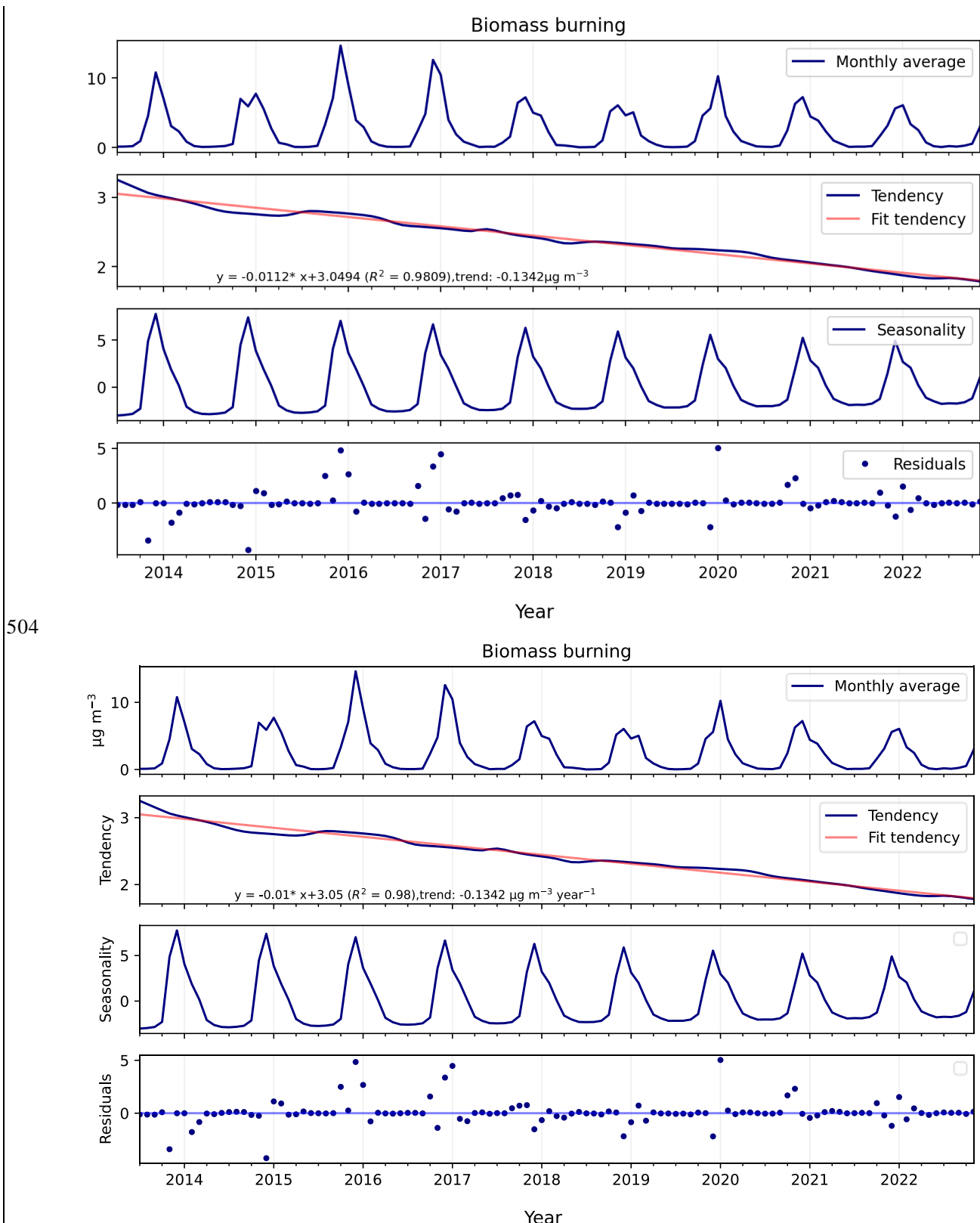
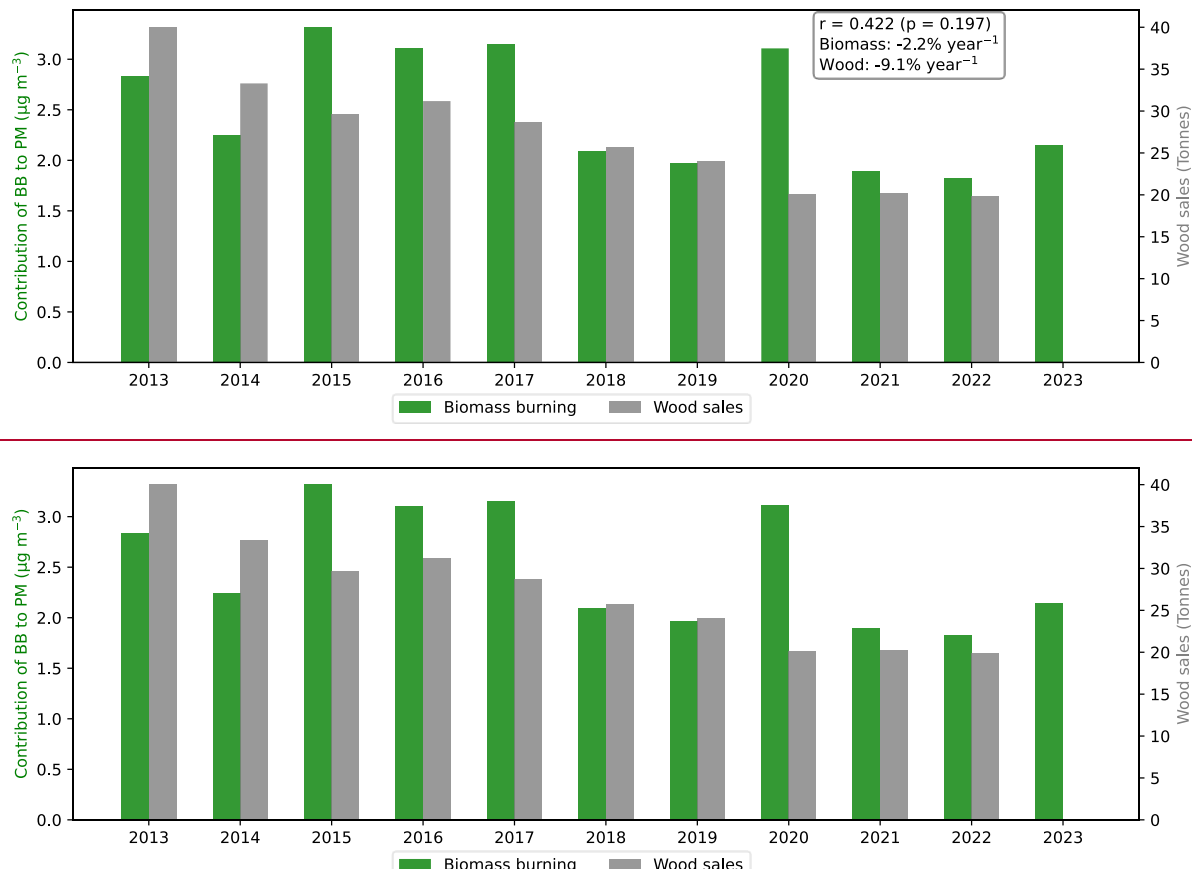


Figure 7. The season-trend (STL) decomposition of biomass burning

This reduction of biomass burning concentrations in Grenoble is 4 times higher than the results from a long-term study (2012 to 2020) in a French rural site - (Observatoire Pérenne de l'Environnement, OPE) (Borlaza et al., 2022a) - estimated at $33 \text{ ng m}^{-3} \text{ yr}^{-1}$ over the same period. Besides the study of Borlaza et al. (2022a), there are

510 no previous PMF studies describing any trend of biomass burning factors. Nevertheless, similar trends were found
 511 for concentrations of biomass burning tracers. In particular, Font et al. (2022) presented a downward trend of
 512 PM_{10} concentration from wood burning (a reduction from 1.5 to 3.8 % yr^{-1}) in urban sites in the United Kingdom
 513 from 2010 to 2021, by calculating the emission of wood burning from aethalometer measurement. Similarly, from
 514 2002 to 2018 in Norway, a downward trend of 2.8% yr^{-1} was also detected for levoglucosan (Yttri et al.,
 515 2021) (Esben Yttri et al., 2021). Additionally, Colette et al. (2021) modeled the trend of the emissions from
 516 different activities in Europe, showing that the trend of PM_{10} heating emissions was decreasing in the period 2000-
 517 2017, with mean rate values varying from 0.8 to 3.3% yr^{-1} for 30 European countries (EMEP monitoring sites).
 518 Even though the chemicals and the period of these studies differ, a decreasing trend is generally observed among
 519 European cities, including the one investigated here. Interestingly, the biomass burning source in Grenoble shows
 520 the strongest decreasing trend, with a reduction of 5.5% yr^{-1} .
 521 Since the biomass burning sources in Grenoble are related to residential heating, the observed reduction of the
 522 concentrations from this source could be linked to household behaviors (including appliance renovation) on top
 523 of the changes in meteorological conditions, lowering the overall heating demand. The average annual biomass
 524 burning sources PMF-derived is compared to the local PM_{10} emission inventory for residential heating (tonnes)
 525 in the Grenoble metropolis, estimated by the regional air quality monitoring agency (Atmo AuRA), to confirm
 526 the trend of biomass burning (Figure 8). This emission inventory has been available until 2022.



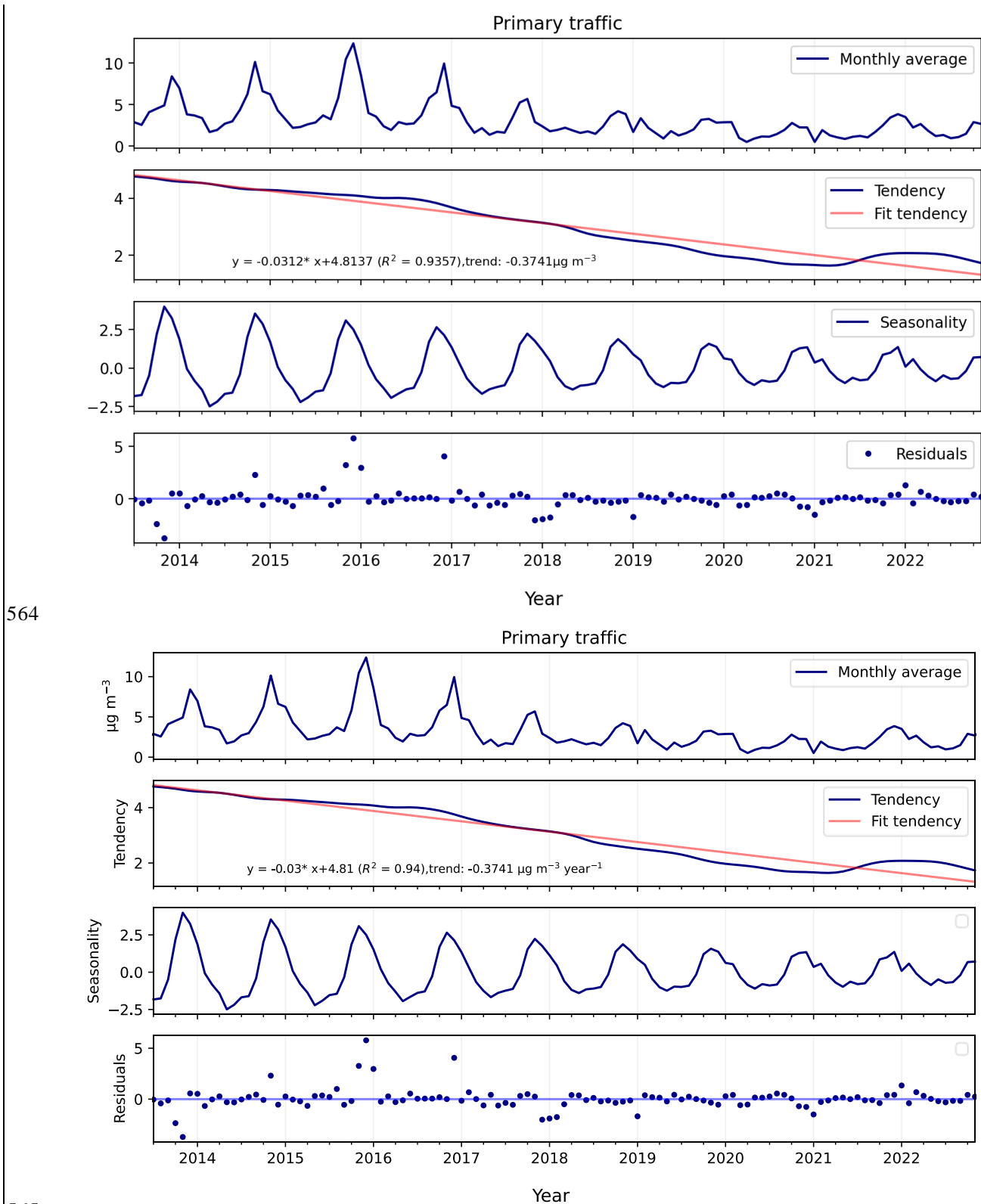
527
 528 **Figure 8. Comparison between annual average PM_{10} emission inventory based on the quantity of wood sales (in grey)**
 529 **in the Grenoble metropolis and the yearly average PM_{10} concentrations from the PMF biomass burning factor (in**
 530 **green).**

548 Except for the year 2020, the annual average of biomass burning agreed with the emission inventory,
549 demonstrating the consistency between the sources observed by the PMF model and the local inventory emission
550 data. Since 2015, the Grenoble metropolis has set up an air-wood bonus to encourage households to renew their
551 individual wood-burning appliance (fireplace or stove). It aims to replace all open fireplaces with closed
552 appliances in October 2024. The downward trend of biomass burning concentration could then be considered as
553 partly due to the implementation of dedicated action plans at the regional scale.

554 **3.3.4. Trends in traffic exhaust emissions**

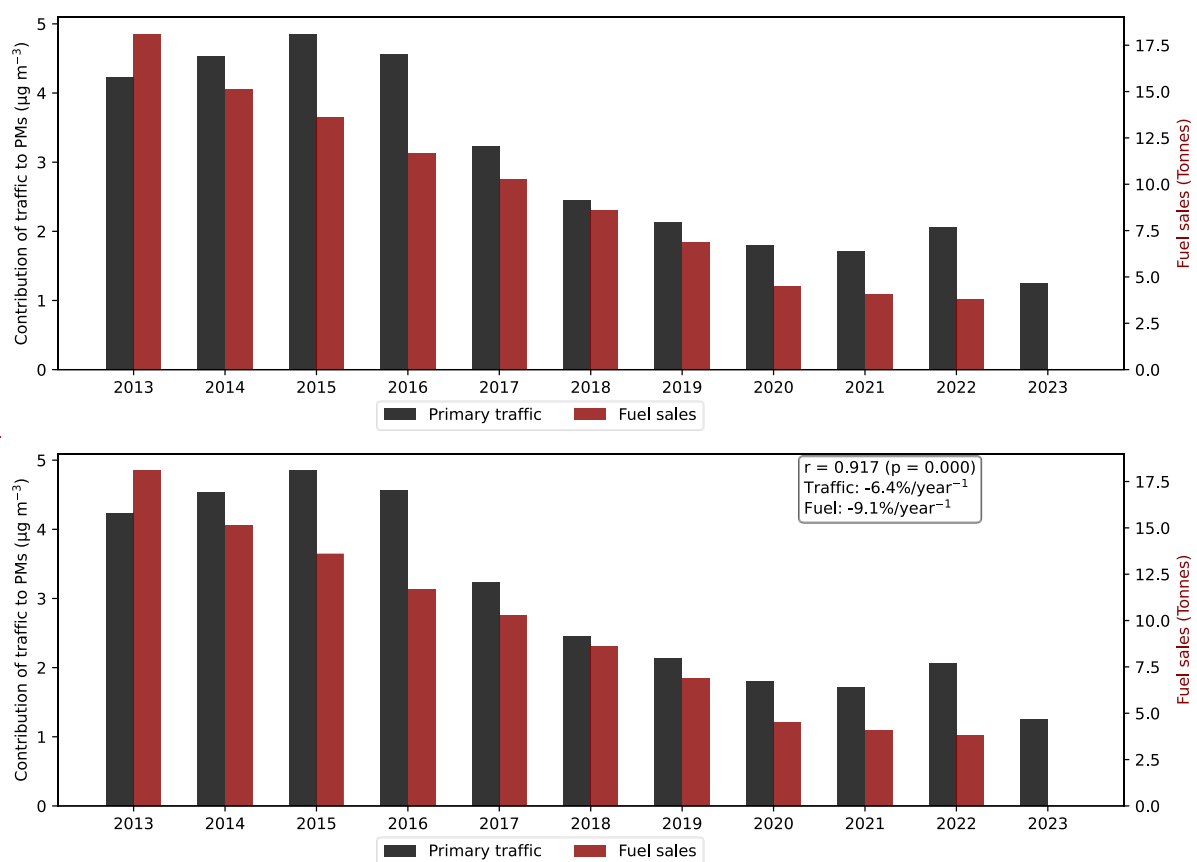
555 Similar to the time series of biomass burning concentrations, the traffic contribution was subjected to specific
556 STL analysis ([Figure 9](#)[Figure 9](#)). A significant downward trend of the concentrations of PM from traffic emission
557 is detected with a reduction of $374 \text{ ng m}^{-3} \text{ yr}^{-1}$ ($12.9\% \text{ yr}^{-1}$) (p-value $<< 0.01$). This reduction is almost 3 times
558 larger than that of the biomass burning concentrations. Traffic concentration before 2017 also showed a clear
559 seasonality with maxima in winter, which nearly disappeared from 2018 onward. It is striking that the same
560 behaviors (strong downward trend and smoothing) are also observed for NO_x concentration, another indicator of
561 traffic exhaust emission, which is also observed for NO_x seasonal patterns (see Supplement S6 and Figure S7).
562 Residuals show extreme values in the same month as biomass burning in 2016 and 2017, matching the PM_{10}
563 episode. The traffic trend closely follows a linear regression fit line, with $R^2 = 0.94$.

Mis en
(France)



The downward traffic trend observed in this study is consistent with another long-term study (2012-2020) of a rural site in France, which showed a traffic trend ~~of $-6.5\% \text{ yr}^{-1}$ (58% total reduction) of $0.1 \mu\text{g m}^{-3} \text{ yr}^{-1}$~~ (Borlaza et al., 2022a). This is aligned with other results of fossil fuel black carbon in several rural sites in France (Font et al., 2025), or EC over many rural sites in Europe (Aas et al., 2024). Additionally, our result also agrees with other

571 studies, like that by Pandolfi et al. (2016), which indicated a downward trend of traffic sources in an urban site in
 572 Spain, with a reduction of $0.11 \mu\text{g m}^{-3} \text{ yr}^{-1}$ (5.6% total reduction), which is lower than that of our study.
 573 Finally, the trend of traffic emission to PM_{10} in 30 European countries was modeled as reported by Colette et al.
 574 (2021), showing a downward trend with a reduction from 2.3 to 3.5% yr^{-1} from 2000 to 2017. As for biomass
 575 burning, the Grenoble supersite seems then experiencing faster reductions in primary traffic PM loadings than
 576 most of others European cities.
 577 Furthermore, the PMF-derived traffic factor was compared to the local PM_{10} traffic emission inventory by fuel
 578 type (provided by Atmo AuRA), revealing very similar trends (Figure 10). In addition, this source is
 579 also compared to the PM_{10} emission by the transport sector (kilotonnes) over France, which was assessed from
 580 the emission inventory data of CITEPA (Figure S11), also confirming the concomitant reductions of traffic
 581 emissions and contributions to PM_{10} in ambient air.



582
 583
 584 **Figure 10. Comparison between annual average PM_{10} emission inventory based on the quantity of fuel sale (red bar)**
 585 **in the Grenoble metropolis and the yearly average PM_{10} concentrations from the PMF-derived traffic source**
 586 **contributions (black bar).**

587 This traffic trend may be separated into three parts. Between 2014 and 2016 with a slow decrease trend of -3% yr^{-1} ;
 588 from 2016 to 2021, with an average reduction of 10% yr^{-1} , and a mild increasing trend of approximately 3% yr^{-1} in the last three years of the study. The beginning of this increase coincides with the post-lockdown period, when
 589 transportation activities were back to normal, resulting in a fairly similar contribution of traffic sources compared
 590 to that in the pre-lockdown period.
 591 Besides the implementation of the two versions of the Euro 6 emission standards (introduced in 2015 and 2018,
 592 respectively), local emission abatement strategies decided by Grenoble municipality from 2016 onwards might be

594 the main drivers for the observed decreasing trends (City's low emission zone
595 <https://zfe.grenoblealpesmetropole.fr/> last assessed: 21/05/2025).

596 **3.4. Trends in PM₁₀ OP sources**

597 In this section, the sources of OP are assessed using regression techniques, which are presented in section 2.6.
598 The most appropriate model is selected based on characteristics of PMF-derived sources and OP_v, as shown in
599 section 3.4.1. Intrinsic OP derived from the best regression model, indicating the highest redox-active PM sources,
600 is presented in section 3.4.2. Finally, section 3.4.3 provides the trend of OP sources, highlighting which sources
601 are the drivers of OP trends.

602 **3.4.1. Selection of the most appropriate model**

603 Following the methodology exposed in Ngoc Thuy et al. (2024), the characteristics of the dataset, including
604 collinearity and heteroscedasticity, are tested in order to select a satisfactory inversion model for OP_{DTT} source
605 apportionment (SA) and OP_{AA} SA (Table S8). The OP SA can be applied for the 11-year PMF solution since the
606 source profiles have been demonstrated to be homogenous over the years. Consequently, the OP^m should be
607 substantially homogenous over the years (Ngoc Thuy et al., 2024), and it is unnecessary to perform the OP SA
608 for each year separately. The characteristic tests indicate that the weighted positive least squares (wPLS) and
609 weighted least squares (WLS) could be suitable models for both OP_{AA} and OP_{DTT} SA. The average accuracy
610 metrics of the testing dataset in 500 iteration runs (including R², RMSE, MAE) of wPLS and WLS were compared
611 to select the most appropriate model (Table S9). Finally, WLS was chosen due to the highest R² and lowest error
612 for both OP_{AA} and OP_{DTT} prediction. The comparison between observed and predicted OP_{AA} and OP_{DTT} showed
613 a good correlation between measured OP and WLS predicted OP, with R² = 0.80 and 0.70 for OP_{AA} and OP_{DTT},
614 respectively (Figure S12 and S13), with n = 1570 for OP_{AA} and OP_{DTT}.

615 In addition, the study revealed good performance of Multiple Layer Perceptron (MLP) and Random Forest (RF)
616 for the training and testing datasets (Table S10). These neural network models were overfitting the results of OP
617 SA for the 6 French sites tested in Ngoc Thuy et al. (2024) since the number of samples was lower than 200 for
618 individual sites. The present study confirmed the conclusion of Ngoc Thuy et al. (2024), demonstrating that a
619 higher number of samples improved the performance of the neural network model. However, such non-linear
620 models do not provide values for intrinsic OP, and cannot be selected for the final results at this stage. However,
621 such non-linear models do not provide values for the intrinsic OP, which is basically the regression slope of the
622 regression. Since the objectives of MLP and RF are not to define a "slope" but to better predict OP, therefore, the
623 "slopes" of such models actually constantly vary with the input data to ensure the best performance of the model.
624 Since the OP intrinsic is not defined, these models cannot be selected for the final results at this stage.

625 **3.4.2. Intrinsic OP of PMF-derived sources**

626 The intrinsic OP of 1 μ g PM₁₀ source (OP^m nmol min⁻¹ μ g⁻¹) is investigated thanks to the WLS technique, resulting
627 in 500 values of OP^m for each source (Table 2 and Table S11). The anthropogenic sources, including
628 biomass burning, industrial, and traffic, have the dominant intrinsic OP_{DTT} and OP_{AA}, which is consistent with the
629 study in 2017-2018 in Grenoble (Borlaza, 2021) and results obtained at other French sites (Ngoc Thuy et al., 2024;
630 Weber et al., 2021) and EU sites (Fadel et al., 2023; Veld et al., 2023). The different ranking of the intrinsic OP

631 of the sources according to the two assays is also aligned with previous results (Weber et al., 2021). While intrinsic
632 OP_{AA} of biomass burning is highest (0.76 nmol min⁻¹ µg⁻¹), followed by industrial (0.48 nmol min⁻¹ µg⁻¹) and
633 traffic (0.38 nmol min⁻¹ µg⁻¹), the order of intrinsic OP_{DTT} is industrial (0.52 nmol min⁻¹ µg⁻¹), traffic (0.38 nmol
634 min⁻¹ µg⁻¹) and biomass burning (0.14 nmol min⁻¹ µg⁻¹). The intrinsic OP_{DTT} of biomass burning is also lower than
635 that of OP_{AA}, as reported by Borlaza et al. (Borlaza et al., 2021), suggesting the synergistic and antagonistic effects
636 between some elements, quinones, or bioaerosols, decreasing the overall intrinsic OP_{DTT} of this source
637 (Pietrogrande et al., 2022; Samake et al., 2017; Xiong et al., 2017).

638 The other anthropogenic sources, including nitrate-rich and sulfate-rich, have lower intrinsic OP than
639 anthropogenic sources associated with combustion (traffic and biomass burning), as reported by Daellenbach et
640 al. (2020). ~~The natural sources have a negligible intrinsic OP (lower than 0.03 nmol min⁻¹ µg⁻¹). The natural~~
641 ~~sources have a negligible intrinsic OP (lower than 0.03 nmol min⁻¹ µg⁻¹ for OP_{DTT} and 0.2 nmol min⁻¹ µg⁻¹ for~~
642 ~~OP_{AA}~~). These findings highlight the high impact of the anthropogenic sources, verified for the overall period 2013-
643 2023.

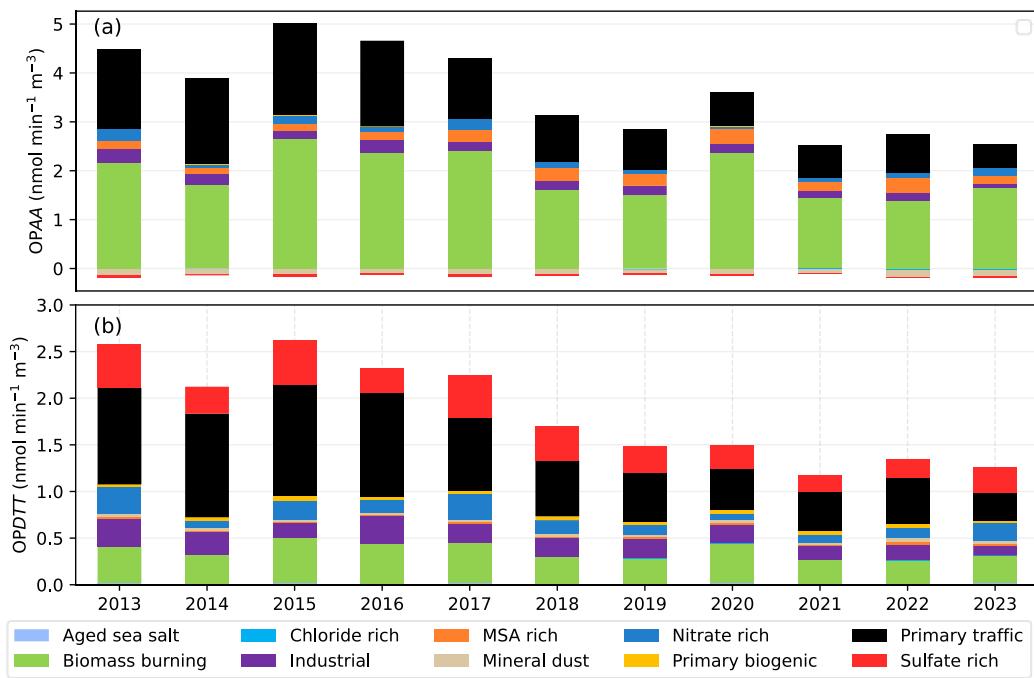
644 **Table 2. Intrinsic OP_{AA} and OP_{DTT} (nmol min⁻¹ µg⁻¹) of PM₁₀ sources (mean ± std of 500 iterations)**

Source	OP _{AA}	OP _{DTT}
Aged sea salt	-0.02 ± 0.07	0.03 ± 0.02
Biomass burning	0.76 ± 0.13	0.14 ± 0.09
Chloride rich	-0.07 ± 0.09	0.01 ± 0.02
Industrial	0.48 ± 0.14	0.52 ± 0.08
MSA rich	0.20 ± 0.04	0.01 ± 0.02
Mineral dust	-0.03 ± 0.06	0.01 ± 0.02
Nitrate rich	0.09 ± 0.16	0.11 ± 0.12
Primary biogenic	0.00 ± 0.04	0.02 ± 0.03
Primary traffic	0.38 ± 0.10	0.24 ± 0.07
Sulfate rich	-0.01 ± 0.08	0.09 ± 0.04

645 **3.4.3. Trends in OP**

646 The trend of OP is first presented by the yearly average contribution of sources to OP_{AA} and OP_{DTT} ([Figure 4](#)
647 [Figure 4](#)), indicating a reduction of OP values over the years. Overall, the yearly average of the OP_{AA}^v and OP_{DTT}^v is
648 decreasing and reached its lowest values in 2021 (2.41 and 1.17 nmol min⁻¹ m⁻³ for OP_{AA} and OP_{DTT}, respectively).
649 From 2018 onward, both assays consistently exhibited lower OP^v values than in preceding years. [Although OP^v](#)
650 [is normalized to PM₁₀ mass concentration](#) [Although OP_v is calculated using PM₁₀ concentration](#), implying that a
651 decrease in PM₁₀ concentration generally reduces OP^v, the contribution of sources to OP is different from that of
652 PM₁₀. While dust and sulfate-rich are dominantly contribute to PM₁₀, biomass burning is the most important
653 contributor to OP_{AA} (1.87 ± 2.7 nmol min⁻¹ m⁻³), and primary traffic is commonly assessed as the largest
654 contributor to OP_{DTT} (0.71 ± 0.70 nmol min⁻¹ m⁻³). The industrial mass contribution is 10 times lower than that of
655 the sulfate-rich. However, industrial emissions appear to contribute much more to OP_{AA} and equally to OP_{DTT} than
656 the sulfate-rich factor. This finding was also observed in 2017-2018 at the same site in Grenoble (Borlaza, 2021).
657 This significant contribution of traffic and biomass burning over the years is more evident when considering
658 relative contribution (Figure S15). These results again emphasize the importance of considering not only the mass
659 concentration but also its redox activity in evaluating the potential adverse health effects of a source of PM.

660 In addition, the temporal evolution of OP_{AA} and OP_{DTT} did not exactly follow PM_{10} trends, especially for the
 661 period of 2016-2017 and 2019-2020. Regarding the period between 2016 and 2017, a dramatic increase in PM_{10}
 662 concentration is observed, principally due to the higher contribution of nitrate and sulfate-rich. On the other hand,
 663 OP_{AA} and OP_{DTT} values remained fairly unchanged between 2016 and 2017. Focus on 2019 and 2020, the PM
 664 concentration and OP_v values are identical (less than $0.001 \mu\text{g m}^{-3}$ and $\text{nmol min}^{-1} \text{m}^{-3}$ of difference, respectively),
 665 while OP_{AA}^v presents a remarkable difference ($\Delta = 0.8 \text{ nmol min}^{-1} \text{m}^{-3}$). Indeed, the discrepancy between 2019
 666 and 2020 in OP_{AA}^v is principally attributable to a higher contribution to biomass burning, which is the dominant
 667 driver of OP_{AA}^v . Overall, the downward trend of OP_{AA} and OP_{DTT} is different from PM_{10} , since the driven sources
 668 of OP and PM are different.



669
 670 **Figure 11. Yearly average contribution of sources to (a) OP_{AA}^v and (b) OP_{DTT}^v**

671 The yearly average may not be properly representative of the trends of OP; therefore, a STL deconvolution was
 672 performed for OP_{AA}^m and OP_{DTT}^m (Figures S16, S17, respectively) to investigate the trend of OP^m over the 11
 673 years of the study. Indeed, considering the trend of the intrinsic OP^m confirms that the downward trend of some
 674 sources leads to a change in the trend of OP_{AA}^m and OP_{DTT}^m .

675 An insignificant linear trend is observed for OP_{AA}^m (fit line: $R^2 = 0.4$, p-values $<< 0.01$), yet its average intrinsic
 676 activity still exhibits a decreasing value, with the annual mean falling by approximately $0.002 \text{ nmol min}^{-1} \text{m}^{-3}$
 677 (2.5 %) across the study period. Interestingly, the seasonality of OP_{AA}^m exactly matches the seasonality of
 678 biomass-burning concentrations, pointing out that the high values of OP_{AA}^m in winter align with biomass-burning
 679 activities. The trend line of OP_{AA}^m did not match the trend of biomass burning nor that of the traffic or industrial
 680 emissions, suggesting the synergistic effect between sources, as well as the influence of the other sources outside
 681 of the winter season, such as MSA-rich and primary biogenic, which get a high ranking of OP_{AA}^m (Table 2).
 682 Conversely, the OP_{DTT}^m showed a significant downward trend ($R^2 = 0.6$, p-value $<< 0.01$), with a reduction of 0.005
 683 $\text{nmol min}^{-1} \mu\text{g}^{-1}$ (6.5%) across 11 years. The seasonality of OP_{DTT} is different from that of biomass burning and
 684 OP_{AA}^m , since biomass burning is not the main driver of OP_{DTT} (only ranked third), indicating a lower influence of
 685 this source on OP_{DTT}^m compared to OP_{AA}^m . Interestingly, a slight increase in OP_{DTT}^m from 2021 onward is also

686 observed, which is associated with PM_{10} and traffic, suggesting that traffic emission could be the main driver for
687 increasing PM_{10} concentration and OP_{DTT^m} from 2021. Overall, the relative decrease of OP_{DTT^m} exceeds that of
688 OP_{AA^m} could be explained by the 4th most important contributor to these OPs. All four leading contributors to
689 OP_{DTT^m} show significant reductions, whereas MSA-rich factor, one of the top four contributors to OP_{AA^m} , has an
690 increasing trend. These findings again underscore that trends in OP^m are governed by the evolution of the sources
691 most active in each assay. Thus, the decrease in the magnitude of the OP_m depends on how its dominant redox-
692 active sources evolve over time.

693 Considering the volume-based metrics (OP_v), a downward trend is detected for OP_{AA} and OP_{DTT} . PM_{10} decreased
694 by 3.9 % over the decade, which is consistently comparable to OP_{AA^v} (4.9 %) and OP_{DTT^v} (5.3 %). This good
695 agreement partially reflects the influence of the PM mass concentration since ~~these OP^v values are normalized to~~
696 ~~PM₁₀ mass concentration~~ ~~these OP_v values are calculated using PM10 concentration~~. However, the slight
697 difference in the relative downward trend could be related to the most driven sources of OP and PM, as discussed
698 above.

699 Finally, the impact of persistent inversion days on the OP^v is also investigated to assess the association between
700 the redox activity of PM sources and thermal inversion. A comparison of the source's contribution to OP^v (for
701 both AA and DTT) between the period with and without persistent inversions is carried out and shown in Figure
702 S14. The comparison confirms the larger increases in average OP_{AA} (85.1%) and OP_{DTT} (63.8 %) compared to
703 that of PM_{10} (39.6%) for the persistent inversion periods. The higher values of OP_{AA} and OP_{DTT} are related to the
704 larger increases in the contribution of local anthropogenic sources, with BB impacting most the OP_{AA} values while
705 traffic significantly influences OP_{DTT} . This result again highlights the potential effect of persistent inversion on
706 the PM10 source's contribution, but all the more of their redox-active properties, which could be associated with
707 the health-relevant metrics (Tassel et al., 2025 in progress).

708 Over the decade, anthropogenic sources have driven OP, with biomass burning impacting OP_{AA} and traffic/
709 industrial sources dominating OP_{DTT} . Frequent thermal inversion in Alpine valley strongly amplifies OP, which is
710 more significant than the mass of PM_{10} itself. Finally, OP_v and intrinsic OP trends over the decade do not align
711 with that of PM_{10} mass, emphasizing the need to prioritize redox-active components over the bulk PM
712 concentration in air quality policy.

713 4. Conclusions

714 Thanks to long-term PM_{10} observations with a detailed set of chemical markers, a comprehensive source
715 apportionment was performed to identify the evolution of PM_{10} sources in Grenoble (France). This is one of the
716 very few studies in Europe that could assess over 11 years of PM_{10} sources and the only study so far investigating
717 trends in PM_{10} -related OP. The trend of PM_{10} sources, especially anthropogenic sources such as biomass burning
718 and primary traffic, was evaluated and linked to the meteorology and emission reduction policies. In addition, the
719 trend of OP^m , OP^v , and sources of OP revealed that the trend of OP depends on the source that drives OP. The
720 analysis of these trends confirms the improvement of the air quality at the Grenoble supersite from 2013 to 2023,
721 and objectivates the main sources that are involved in their concentration's decrease.

722 Nevertheless, the following methodological limitations in this long-term study shall be kept in mind:
723 - Daily concentrations of metal elements were only analyzed for some periods (2013, 2017-2018, 2020-2021),
724 while the remaining data were derived from weekly sampling. An imputation technique was implemented to

725 impute daily concentrations. The PMF result demonstrated the stability of most chemical profiles at Grenoble
726 from 2013 to 2023, compared to those previously published (Borlaza et al., 2021), despite these uncertainties in
727 the imputed metal concentrations.

728 - The process of implementing such a PMF analysis strategy is not straightforward. A combined PMF approach
729 could be used for datasets with different time resolution (Via et al., 2023). This approach would allow combining
730 the 7-day and daily filter samples into a PMF without performing imputation.

731 - The lack of a secondary biogenic organic aerosol tracer in long-term observations prevents the identification of
732 the BSOA source, which could make up about 10% of the total mass of PM₁₀ on a yearly average, as observed in
733 previous work at the site (Borlaza et al., 2021), which used 3-MBTCA and picnic acid for the yearly period of
734 observation.

735 Thanks to long-term PM₁₀ observations with a detailed set of chemical markers, a comprehensive source
736 apportionment was performed to identify the evolution of PM₁₀ sources in Grenoble (France). This is one of the
737 very few studies in Europe that could assess over 11 years of PM₁₀ sources and the only study so far investigating
738 trends in PM₁₀-related OP. The trend of PM₁₀ sources, especially anthropogenic sources such as biomass burning
739 and primary traffic, was evaluated and linked to the meteorology and emission reduction policies. In addition, the
740 trend of OP^m, OP^v, and sources of OP revealed that the trend of OP depends on the source that drives OP. The
741 analysis of these trends confirms the improvement of the air quality at the Grenoble supersite from 2013 to 2023,
742 and objectivates the main sources that are involved in their concentration' decrease.

743 Overall, a total of ten sources were identified, including aged sea salt, biomass burning, chloride-rich mineral
744 dust, MSA-rich, nitrate-rich, industrial, primary biogenic, and primary traffic. The source chemical profiles are
745 consistent with those presented in 2017-2018 (Borlaza et al., 2021), demonstrating that the sources of PM₁₀ in
746 Grenoble were relatively stable during our study period. The trend of PM₁₀ sources was investigated using STL
747 decomposition, which reveals a downward trend for all the PM₁₀ sources over 11 years, especially for the
748 anthropogenic sources. Extending PMF outputs to oxidative potential apportionment showed that biomass
749 burning, traffic, and industrial emissions dominate redox activity in both the ascorbic acid (AA) and dithiothreitol
750 (DTT) assays. Trend analysis of volume- and mass-normalized OP metrics indicates that biomass burning governs
751 the long-term behavior of OP_{AA}. In contrast, traffic is the principal driver of OP_{DTT} assay, underscoring source-
752 specific control of PM₁₀ OP in the Grenoble atmosphere.

753 Both of these anthropogenic sources, as well as their influences on PM₁₀ OP, showed significant decreasing trends
754 concomitantly to the implementation of emission reduction strategies (both at the national and regional levels)
755 that should be reinforced to reach the goals of the European zero pollution action plan and the recently revised
756 Directive on ambient air quality (22024/2881/EU). The continuation of these measurements will take place in the
757 coming years, with this site being selected as one of the supersites for the new EU Air Quality directive.

758 **Data availability**

759 The datasets could be made available upon request by contacting the corresponding authors.

760 **Author contributions**

761 VDNT performed the source apportionment and the trend of sources, and the result visualisation. GU, JLJ
762 mentoring, supervision, validation of the methodology and results. RE, SD, CV, and AN contributed to data
763 acquisition (analytical investigation on samples) and data curation. OF, JLJ, and GU acquired funding for the
764 original PM sampling and analysis. VDNT wrote the original draft. All authors reviewed and edited the
765 manuscript.

766 **Competing interests**

767 The authors declare that they have no conflict of interest.

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779

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