1 Unveiling the optimal regression model for source

2 apportionment of the oxidative potential of PM₁₀

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14 Abstract

- 15 The capacity of particulate matter (PM) to generate reactive oxygen species (ROS) in vivo leading to oxidative
- stress, is thought to be a main pathway for the health effect of PM inhalation. Exogenous ROS from PM can be
- 17 assessed by acellular oxidative potential (OP) measurements as a proxy of the induction of oxidative stress in the
- lungs. Here, we investigate the importance of OP apportionment methods on OP repartition by PM₁₀ sources in
- different types of environments. PM₁₀ sources derived from receptor models (e.g. EPA PMF) are coupled with
- 20 regression models expressing the associations between PM₁₀ sources and PM₁₀OP measured by ascorbic acid
- 21 (OP_{AA}) and dithiothreitol assay (OP_{DTT}). These relationships are compared for eight regression techniques:
- 22 Ordinary Least Squares, Weighted Least Squares, Positive Least Squares, Ridge, Lasso, Generalized Linear
- 23 Model, Random Forest, and Multilayer Perceptron. The models are evaluated on one year of PM₁₀ samples and
- 24 chemical analyses at each of six sites of different typologies in France to assess the possible impact of PM source
- 25 variability on PM₁₀ OP apportionment. PM₁₀ SSource-specific OP_{DTT} and OP_{AA} and out-of-sample apportionment
- accuracy vary substantially by model, highlighting the importance of model selection depending on the datasets.
- 27 Recommendations for the selection of the most accurate model are provided, encompassing considerations such
- as multicollinearity and homoscedasticity.
- 29 Key words: Oxidative potential, source apportionment, OP apportionment.

1. Introduction

- 31 Ambient particulate matter (PM) is one of the key contributors to atmospheric pollution and is responsible for
- 32 approximately 7 million premature deaths worldwide yearly (WHO, 2021). Many epidemiological studies have
- 33 linked PM exposure to adverse health effects including (i) acute effects studies using time series and related studies
- 34 to evaluate the immediate impact of PM exposure (Bell et al., 2004; Dominici, 2004; Peng et al., 2009; Pope &
- 35 Dockery, 2006) and (ii) cohort studies aiming to evaluate the long-term effects of chronic PM exposure (Ayres et
- 36 al., 2008; Beelen et al., 2014; Crouse et al., 2012, 2015; Pelucchi et al., 2009; Yu et al., 2021). These studies
- 37 mainly focused on the association with PM mass concentrations. However, various research shows that the impacts
- 38 of PM also depend on other factors such as chemical composition, size distribution, particle morphology, and
- 39 biological mechanisms (Brook et al., 2010) (Crouse et al., 2012). PM's capacity to generate reactive oxygen species
- 40 (ROS) in vivo has recently been introduced as a pivotal indicator of PM biological mechanism, with direct

41 implications for oxidative stress and cellular damage (Akhtar et al., 2010; Ayres et al., 2008; Leni et al., 2020; Li

42 et al., 2008; Lodovici & Bigagli, 2011; Mudway et al., 2020; Nelin et al., 2012; Rao et al., 2018). The quantification

43 of the PM capacity to oxidize a biological media is called oxidative potential (OP) (Bates et al., 2019; Daellenbach

et al., 2020; Dominutti et al., 2023). Various acellular assays of OP have been introduced, differentiating ROS

45 generation mechanisms of PM (Calas et al., 2018; Dominutti et al., 2023). Dithiothreitol (DTT) and ascorbic acid

46 (AA) assays are two of the commonly used ones in the literature (Liu & Ng, 2023).

The relationship between PM chemical components and OP activities may identify which components are <u>the</u> most prone to generate ROS (Calas et al., 2018, 2019; Crobeddu et al., 2017; Godri et al., 2011; Janssen et al., 2014;

Szigeti et al., 2015, 2016; Yang et al., 2014). However, this research pathway struggles with the co-variation

between measured and unmeasured PM components (Calas et al., 2018; Weber et al., 2018). An alternative

approach is to examine the association between OP and sources of PM obtained using receptor models such as

chemical mass balance, positive matrix factorization (PMF), or principal components analysis. PMF is the most

popular method for its ability to quantify PM source contributions without extensive prior information on specific

sources at the site studied (Belis et al., 2013; Brown et al., 2015; Paatero & Hopke, 2009; Paatero & Tappert, 1994;

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Regression analysis is the most common and effective way to estimate the redox activity of receptor model-derived PM sources (Bates et al., 2015; Deng et al., 2022; Li et al., 2023; Liu et al., 2018; Shangguan et al., 2022; Verma et al., 2014; J. Wang et al., 2020; Yu et al., 2019). Generally, this is achieved by regression analyses to characterize the relationship between OP activities (nmol min⁻¹ m⁻³) and PM sources contribution (µg m⁻³). This approach provides the OP activities attributed to each microgram of each source (nmol min⁻¹ µg⁻¹), denoted as intrinsic OP, which can be used to calculate the contribution of each source for each observation day. Numerous regression models can be used for such OP source apportionment (SA), with multiple linear regression fitted by ordinary least squares (OLS) being the most common regression technique (Bates et al., 2015; Deng et al., 2022; Li et al., 2023; Liu et al., 2018; Shangguan et al., 2022; Verma et al., 2014; Y. Wang et al., 2020; Yu et al., 2019). Further, some studies exclude sources with negative intrinsic OP, assuming that negative OP activities are geochemically nonsensical (Bates et al., 2018; Weber et al., 2018). Additionally, weighted least square can be used to introduce a weighting term, usually using the OP analysis uncertainties to take into account the measurement uncertainties of the OP assays (Borlaza et al., 2021; Daellenbach et al., 2020; Dominutti et al., 2023; Fadel et al., 2023; in 't Veld et al., 2023b; Weber et al., 2021). Finally, non-linear models, such as multilayer perceptron, have been used to try to capture possible non-linearities between OP activities and PM sources (Borlaza et al., 2021; Elangasinghe et al., 2014; D. Wang et al., 2023). However, no study to date has compared the performance and applicability of these various regression models. Each model implies different assumptions which should be carefully considered when selecting a given model.

This study aims to evaluate the variability in PM₁₀ OP SA techniques by comparing eight regression techniques: multiple linear regression fitted by ordinary least squares (OLS), weighted least squares (WLS), positive least squares (PLS), Ridge regression (Ridge), Least Absolute Shrinkage and Selection Operator (Lasso), generalized linear model (GLM), random forest (RF), and multilayer perceptron (MLP). These techniques are applied to apportion PM₁₀ OP_{AA} and PM₁₀ OP_{DTT} to PM₁₀ sources at six sites in France. The PM₁₀ SA outputs have been published previously in Weber et al., (2021), using a harmonized PMF methodology based on one year of sampling with similar chemical analyses for a large set of chemical tracers. The results of the PM₁₀ OP SA models are compared with regard to the estimated intrinsic PM₁₀ OP of each source, the out-of-sample accuracy of the apportionment, and the assumptions inherent in each model. The most appropriate model at each site is compared with OLS to quantify the difference between choosing a model based on data characteristics vs. using the most common approach. Finally, this study provides guidelines for selecting the most suitable model in the strategy for OP contribution regarding sources of PM₁₀. This holds particular significance in the context of the implementation

of OP monitoring as a novel air quality metric as foreseen in research programs (such RI-Urbans) and in the process of the revision of the European Directive 2008/50/CE.

2. Methodology

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2.1. General organisation of this work

Figure 1 illustrates the general workflow of this work. Sections 2.2, 2.3, and 2.4 describe the methods used to analyse the temporal evolution of PM_{10} sources and PM_{10} OP-activities, identify collinearity among PM_{10} sources, and examine homoscedasticity in the relationship between PM₁₀ OP activities and PM₁₀ sources. Section 2.5 describes the eight regression techniques (OLS, WLS, PLS, Ridge, Lasso, GLM, RF, and MLP), used for PM₁₀ OP SA. Each technique is applied to each site separately using PM₁₀ OP_v (nmol min⁻¹ m⁻³) as the dependent variable and PM_{10} sources (µg m⁻³) as independent variables. The coefficient of the regression called the intrinsic PM_{10} OP of the source (nmol min⁻¹ μ g⁻¹), represents the capacity of each μ g of PM₁₀ from the given source to generate oxidative stress; the higher the intrinsic PM₁₀ OP of a source, the more redox-active. Each model is trained on a randomly selected (without replacement) 80% subsample of the dataset and validated on the remaining 20%. This process is repeated 500 times to estimate uncertainty, a method particularly needed for sources with strong seasonality. For WLS, PLS, Ridge, and Lasso models, PM₁₀ OP analytical errors were used as a weighting, implying that the PM₁₀ OP with the high analysis uncertainties has less influence on the model. These 8 regression techniques were applied to find the relationship between PM₁₀ OP and PM₁₀ sources, however, PLS, Ridge, and Lasso were performed 2 times, with and without weighting, consequently, there are 11 results of regression techniques that will be presented. Section 2.6 describes the statistical validation of the models using root mean square error (RMSE), mean absolute error (MAE), R-square (R²). The geochemical validation is based on the regression coefficient (the intrinsic PM₁₀ OP) of each source. These are calculated separately for the training and testing data and averaged across the 500 sampling iterations.

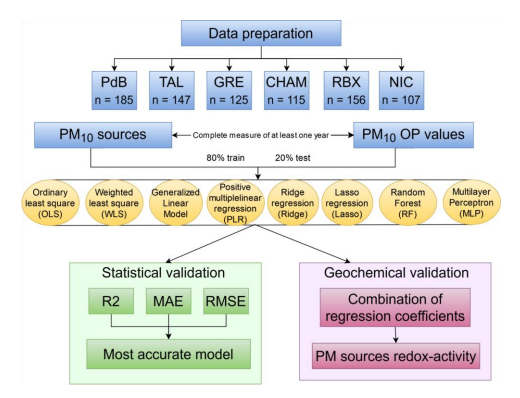


Figure 1. Workflow of the comparison of PM₁₀ OP sources apportionment methodology

2.2. Study sites and PM₁₀ sources

Six French sites are selected in this work for their different typologies: Roubaix and Nice (traffic sites within urban areas), Port-de-Bouc (industrial hotspot), Talence (urban background site), Grenoble and Chamonix (urban background sites in Alpine Valley). At each site, sampling was conducted over at least one year to capture the complete annual evolution of PM₁₀ and its components. These sites and sampling series were previously used and described by Weber et al. (2019).

In brief, daily filter samples were collected on pre-heated Pallflex quartz fibre filters every third day through high-volume sampling (DA80, Digitel). These filters were analyzed to determine PM's chemical species and OP activities. Further details regarding the chemical species and PM₁₀OP analyses methodology can be found in Weber et al. (Weber et al., 2019, 2021). Briefly, the elemental carbon (EC) and organic carbon (OC) were analyzed using the EUSAAR2 thermo-optical protocol with a Sunset Lab analyser. Major ionic components (Cl⁻, NO₃⁻, SO₄²⁻, NH₄⁺, Na⁺, K⁺, Mg²⁺, Ca²⁺) and methanesulfonic acid (MSA) were measured by ion chromatography (IC). Anhydro-sugars and saccharides (including levoglucosan, mannosan, arabitol, sorbitol, and mannitol) were analysed by high-performance liquid chromatography with pulsed amperometry detection (HPLC-PAD). Major and trace elements (Al, Ca, Fe, K, As, Ba, Cd, Co, Cu, La, Mn, Mo, Ni, Pb, Rb, Sb, Sr, V, and Zn) were determined by inductively coupled plasma atomic emission spectroscopy or mass spectrometry (ICP-AES or ICP-MS). Furthermore, colocated PM₁₀ measurements were conducted automatically at each site using the Tapered Element Oscillating Microbalance equipped with a Filter Dynamics Measurement System (TEOM-FDMS).

We used the PM₁₀ sources identified by Weber et al.₇ (2019), who performed a separate PMF for each site using a harmonized approach for all sites (same chemical species and measurement methods, same procedure to estimate uncertainties, same constraints on the preliminary solutions). Table 1 provides a data description, including the sampling duration, the number of samples collected, and the identified PM₁₀ sources at each site, while Figure 2 presents the localisation of the sites in France, together with the respective proportion of each PM₁₀ source at each site.

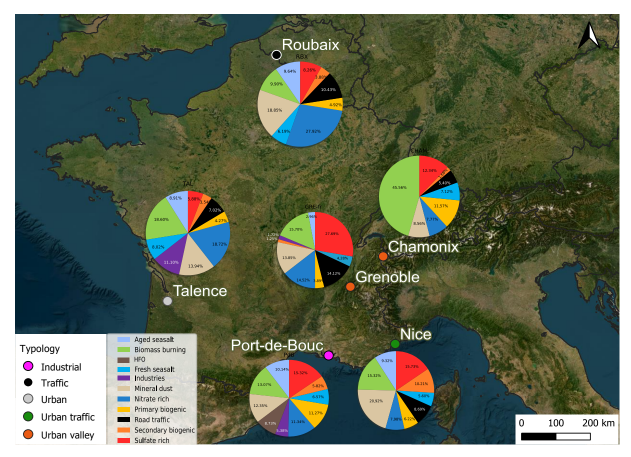


Figure 2. The location of the selected sites for this study. The small colored dots represent the typology of sites. The pie charts are the PM_{10} source apportionment for each site with the colors identifying the $PM_{\underline{10}}$ sources. Background photography from ESRI satellite.

Table 1. Data description

	PdB	TAL	GRE-fr	CHAM	RBX	NIC
Name	Port de Bouc	Talence	Grenoble	Chamonix	Roubaix	Nice
N of samples	185	147	125	115	156	107
Sampling dates	2014-06 to 2016-06	2012-02 to 2013-04	2017-02 to 2018-03	2013-11 to 2014-10	2013-01 to 2014-05	2014-07 to 2015-05
N of sources	10	10	10	8	9	9

2.3. OP analysis

 $\underline{PM_{10}}$ OP assays were performed on $PM_{\underline{10}}$ extracted from the filters using simulated lung fluid, as detailed in Calas et al. (2017, 2018). The AA assay involved ascorbic acid, a natural antioxidant in the lungs inhibiting lipid and protein oxidation in the lining fluid, using the method presented by Kelly & Mudway (2003) and further described by Calas et al., (2018). Conversely, the DTT assay used dithiothreitol (DTT) as a chemical surrogate for cellular reducing agents, specifically nicotinamide adenine dinucleotide and nicotinamide adenine dinucleotide phosphate

147 oxidase, thereby replicating in vivo interactions between PM₁₀ and biological oxidants (Calas et al., 2018; Cho et 148 al., 2005). Both assays measured the consumption of AA or DTT during the assay, i.e., the rate of the transfer of 149 electrons from AA or DTT to oxygen. The assays were conducted with 96-well plates of UV-transparent quality 150 (CELLSTAR, Greiner-Bio), and absorption measurements were acquired using a TECAN spectrophotometer, 151 Infinite M200 Pro, at the wavelengths of 265nm for the AA assay and 412nm for the DTT assay (Calas et al., 2017, 2018, 2019). Each sample extraction was subjected to four analyses; the PM10 OP activities in this study 152 153 represents the mean and the analysis uncertainty is the standard deviation of these four PM_{10} OP analyses. After 154 analysis, the PM₁₀ OP activities of each sample were blank-subtracted using lab and field blanks, and normalized 155 using the air sampling volumes and the mass concentration. The resulting OP_V represents the PM_{10} OP activities 156 due to PM₁₀ per cubic meter of air (nmol min⁻¹ m⁻³). To simplify the denotation of PM₁₀ OP, OP is used to represent 157 PM₁₀ OP throughout this article.

2.4. Collinearity and heteroscedasticity tests

The result of a regression model strongly depends on the characteristics of the dataset because each model makes assumptions about the data. Two critical assumptions in OLS regression analysis are that (1) there is little collinearity between independent variables (the PM₁₀ sources in this study), and (2) the variance of the regression residuals is constant (called homoscedasticity). These assumptions should be tested in different ways.

2.4.1. Collinearity

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Collinearity occurs when one or more of the independent variables is close to a linear combination of the other independent variables. When collinearity is present, small changes in the data can cause large changes in estimated coefficients, and the estimated standard errors of the coefficients are large. Variance Inflation Factor (VIF) is an indicator of the collinearity between the independent variables (Craney & Surles, 2002; O'Brien, 2007; Rosenblad, 2011). VIF of a specific source is calculated as:

$$VIF_{i} = \frac{1}{1 - R_{i}^{2}}, i = 1, ..., p - 1 (Eq1)$$

In this equation, p is the number of PM_{10} sources, R^2 is the coefficient of determination of a multiple linear regression model between the ith source and the other sources. VIF values of a PM₁₀ source present a range between 1, and ∞ . The higher the VIF values, the greater the collinearity between this PM₁₀ source and the other ones. A 173 VIF value between 5 and 10 is commonly interpreted as moderate collinearity, while values greater than 10 indicate high collinearity (Craney & Surles, 2002).

2.4.2. Heteroscedasticity

Heteroscedasticity occurs when the variance of regression residuals is not constant but varies for different values of the dependent variable. In this case, the estimated standard errors of the regression coefficients are not reliable. The Goldfeld-Quandt test was developed by Goldfeld & Quandt (1965) to evaluate residual variance in a regression model. To implement the Goldfeld-Quandt test, an OLS regression was performed between OP and PM₁₀ sources to identify the residual of OP prediction. Next, the PM₁₀ sources and residual corresponding are divided into three segments: the upper segment is the group with higher PM₁₀ sources concentration, the lower segment is the group with lower PM₁₀ sources concentration, and the middle segment, constituting 10% of the moderate PM₁₀ concentration, is excluded. A subsequent regression analysis is then conducted on the two remaining subgroups to determine the ratio of residual sums of squares. Finally, an F-test is conducted on this ratio to assess whether the variances are the same, with a p-value below 0.05 interpreted as evidence of heteroscedasticity.

The Variance Inflation Factor (VIF) and the Goldfeld-Quandt test were performed in Python 3.9, using the statsmodels 0.14.0 package (Seabold & Perktold, 2010).

2.5. Regression models

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The fundamental principle of regression models in this study is to use the PM₁₀ sources to predict OP activities by identifying the parameters (coefficients and residuals) that minimize an error term (Hastie, 2009). A simple regression model can be represented by Eq. 2, which defines the estimated function of the regression model, and

193 Eq. 3, which estimates the residuals.

$$\hat{y} = f(X) + e(Eq2)$$

$$e = y - \hat{y}(Eq3)$$

Here, \hat{y} is the estimated OP (nmol min⁻¹ m⁻³), X are the PM₁₀ source contributions (μ g m⁻³), y is the observed OP (nmol min⁻¹ m⁻³), and e denotes the residuals (nmol min⁻¹ m⁻³). Each model has certain assumptions and a minimization term, as presented below.

Ordinary least squares (OLS):

OLS is a linear regression technique that minimizes the residual sum of squares. This model is based on several assumptions: (1) **Linearity:** The relationship between OP and PM₁₀ sources is linear. (2) **Independence:** The PM₁₀ sources must be independent, with no collinearity. (3) **Homoscedasticity:** The variance of residuals is constant across all values of PM₁₀ sources. (4) **Normality:** The residuals are normally distributed. In the OLS model, the estimated equation and objective to minimize are defined as follows:

$$\hat{y} = \beta_0 + \sum_{i=1}^{p} \beta_i * x_i (Eq4)$$

206 Minimize:
$$\sum_{i=1}^{m} (y_i - \hat{y_i})^2$$
 (Eq5)

Here, the β_0 denotes the intercept (nmol min⁻¹ m⁻³), β_i represents the regression coefficient (intrinsic OP, nmol min⁻¹ μ g⁻¹) of source i, x_i is the concentration of source i (μ g m⁻³), p is the number of PM₁₀ sources, and m is the number of observations.

210 Weighted least square (WLS):

The assumptions and the minimization term in WLS closely align with those in OLS. The only difference is that WLS accounts for heteroscedasticity by introducing a weighting term for individual OP observations, whose variance is assumed to be related to the variance of the residuals. The estimation equation in WLS is the same as that of OLS, but the objective to minimize is expressed as:

215 Minimize:
$$\sum_{i=1}^{m} (y_i - \hat{y}_i)^2 * w_i (Eq6)$$

$$w_i = \frac{1}{SD_i^2}$$

With w_i being the weight assigned to each observation, and SD_i is the OP analysis variance of each observation.

Positive least square (PLS):

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The assumptions for PLS primarily include linearity, independence, and normality. PLS can be applied with weighting, if there is heteroscedasticity in the data. PLS extends OLS with the constraint that the regression coefficients must be non-negative. The estimation equation and the error term, PLS, are similar to OLS (without weighting) and WLS (applying weighting). To ensure the positivity of coefficients, a specific condition must be met:

$\beta_i \geq 0$, $\forall i$ in PM sources (Eq7)

225 Ridge:

Shrinkage methods such as Ridge regression try to produce a more interpretable model or reduce error in the presence of collinearity by selecting a subset of the independent variables. Ridge regression is introduced by Hoerl & Kennard (1970), which incorporates a penalty term that shrinks the coefficients towards zero. The Ridge regression minimizes the residual sum of squares plus a penalty term proportional to the sum of squares of the coefficients (L2 regularization) as shown in Eq 8 and Eq 9. Consequently, Ridge regression reduces the influence of a PM₁₀ source that exhibits minimal impact on OP prediction without excluding it from the model.

232 Minimize:
$$\sum_{i=1}^{m} (y_i - \hat{y}_i)^2 + \lambda * \sum_{j=1}^{p} \beta_j^2$$
 (Eq8)

233 Minimize:
$$\frac{1}{2m} \sum_{i=1}^{m} w_i (y_i - \widehat{y}_i)^2 + \lambda * \sum_{j=1}^{p} \beta_j^2 (Eq9)$$

where λ is the parameter representing the amount of shrinkage, the larger λ , the greater the shrinkage. The hyperparameter tuning was implemented with different values of λ (5, 1, 0.5, 0.1, 0.01, 0.005, 0.001, 0.0005, 0.001). The best λ for every site varied from 0.005 to 0.01 and in this study, 0.01 was selected. Ridge can be

applied with weighting to account for heteroscedasticity. Least Absolute Shrinkage and Selection Operator (Lasso):

Lasso (Tibshirani, 1996) is a shrinkage method that uses a penalty term proportional to the sum of the absolute regression coefficients (L1 regularization). This penalty term shrinks the coefficients of a source with a low impact on OP prediction to zero, effectively removing it from the model. This results in a sparse model that may be easier to interpret and may reduce error on out-of-sample data. However, Lasso is more sensitive to outliers than ridge regression and is less stable when data are collinear. Lasso can be applied with weighting to account for heteroscedasticity.

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$$Minimize: \sum_{i=1}^{m} (y_i - \hat{y_i})^2 + \lambda * \sum_{j=1}^{p} |\beta_j| (Eq.910)$$

$$m$$

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$$Minimize: \frac{1}{2m} \sum_{i=1}^{m} w_i (y_i - \hat{y}_i)^2 + \lambda * \sum_{j=1}^{p} |\beta_j| (Eq11)$$

Similar to Ridge, λ is the parameter representing the amount of shrinkage. λ is selected as 0.01 in this study by running the hyperparameter tuning using the same values as for Ridge.

Generalized linear model (GLM):

Generalized linear models, as introduced by McCullagh (1989), provide a framework for regression analysis that can contain non-normal error distributions and capture non-linear relationships between OP activities and PM_{10} sources. GLM allows for error variance that is a function of the predicted value, hence accounting for heteroskedasticity. Key assumptions underlying GLM include (1) independence, (2) the non-normal distribution of OP, and (3) the relationship between the PM_{10} sources and the transformed OP (logarithm in this study) is linear. The mathematical expression for GLM can be represented as follows:

$$log(\hat{y}) = \beta_0 + \sum_{i=0}^{p} \beta_i * x_i (Eq102)$$

where β_0 denotes the intercept, β_i represents the regression coefficient of source i, and x_i is the concentration of source i.

Random forest (RF):

RF, an ensemble learning method introduced by Breiman (2001), combines multiple decision trees to make predictions. In the reference implementation, each tree is grown on a bootstrap sample of the data and a random subset of the available features is evaluated at each node to choose the best split. The predictions of all trees are averaged to give the forest's final prediction. RF is customizable via hyperparameters such as the number of trees, the size of the bootstrap sample, and the number of features to evaluate at each node. The hyperparameters tuning used 5-fold cross-validation on the training data for hyperparameter tuning. The training dataset was separated into 5 parts: 4 parts were used for training, and the remaining was used for validation. This process was repeated 5 times, and the hyperparameter value producing the lowest mean RMSE across the 5 parts was selected. The hyperparameters tuning is shown in section \$1.1 Supplement. The hyperparameters of RF in this study were chosen by tuning, as shown in section \$1.1 Supplement.

RF does not assume a specific equation to express the relationship between OP activities and $PM_{\underline{10}}$ sources, with the result that intrinsic OP could not be computed in this regression model. Nevertheless, RF can estimate the relative importance of each $PM_{\underline{10}}$ source in OP prediction. This study estimated the permutation importance of each $PM_{\underline{10}}$ source as the mean increase in the mean squared error of predicted OP when the values of the $PM_{\underline{10}}$ source were permuted.

Multilayer perception (MLP):

MLP is an artificial neural network that consists of multiple layers of interconnected nodes or neurons organized in a feedforward structure (Akhtar et al., 2018; Bourlard & Wellekens, 1989; Chianese et al., 2018). These layers include an input layer (PM_{10} sources), one or several hidden layers, and an output layer (OP_{AA} or OP_{DTT} activities). In MLP, the neurons in the hidden layers are linked with the previous neurons by the connection weight, where every neuron is independent and has a different weight. The output of each neuron depends on its inputs and an activation function, which, if non-linear, allows the model to capture non-linear relationships. The implementation of MLP includes three steps: (1) forward pass to training model: the input is passed to the model, multiplied with an initial weight, add bias at every layer, then calculate output of the model. (2) error calculation: after applying step 1, the output of the model and the observed data are used to calculate the error. (3) backward pass: the error is propagated back through the network, and then the weights are adjusted to minimize overall error. These 3 steps are repeated until the error is minimized.

The choice of hyperparameters to ensure the MLP model's robustness is processed by hyperparameter tuning <u>using</u> 5-fold cross-validation as shown in section S1.2 of the supplement. Thanks to hyperparameter tuning, the two hidden layers and a logistic sigmoid activation function were selected in this study to capture the non-linear relationships between OP activities and PM₁₀ sources, and shown in section S1.2 of the supplement. Thanks to hyperparameter tuning, the two hidden layers and a logistic sigmoid activation function were selected in this study to capture the non-linear relationships between OP activities and PM sources.

All regression models were performed using the Python package statsmodels 0.14.0 (Seabold & Perktold, 2010) and scikit-learn 1.3.1 (Pedregosa et al., 2011).

2.6. Performance of the models

The performance metrics R-square (R²), mean absolute error (MAE), and root mean square error (RMSE) were used to assess the goodness of fit of models as described by Kuhn & Johnson (2013). R² quantifies the model's ability to explain the variance in the data. R² equal to 1 indicates a perfect fit. RMSE represents the aggregation of the individual differences between predicted OP and measured OP, while MAE assesses the average magnitude of errors between them. Lower RMSE and MAE values indicate a better fit, with a perfectly fitting model yielding an RMSE or MAE of 0. Eq132, Eq143, and Eq154, respectively, define R², MAE, RMSE. These indicators are computed for the training and testing data of each sampling iteration and averaged across the 500 sampling iterations.

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$$R^{2} = 1 - \frac{\text{Sum of Squared Residuals}}{\text{Total Sum of Squares}} = 1 - \frac{\sum_{i=0}^{m} (y_{i} - \widehat{y_{i}})^{2}}{\sum_{i=0}^{m} (y_{i} - \widehat{y_{i}})^{2}} (Eq123)$$

$$MAE = \frac{\sum_{i=0}^{m} |y_{i} - \widehat{y_{i}}|}{m} (Eq14)$$

$$RMSE = \sqrt{\frac{\sum_{i=0}^{m} (y_{i} - \widehat{y_{i}})^{2}}{m}} (Eq15)$$
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3. Result and discussion

Assessments of collinearity and homoscedasticity are addressed in Section 3.1. Model performance, including key performance metrics and identification of the optimal model, is detailed in Section 3.2. Section 3.3 compares the intrinsic OP estimated by the different models. Section 3.4 compares intrinsic OP between the combined best-fit and reference models. Lastly, Section 3.5 proposes recommendations for selecting an appropriate model.

3.1. Dataset characteristics

The contributions of identified sources (µg m⁻³) and the OP_v activities (nmol min⁻¹ m⁻³) in each site are presented in Figure 3, illustrating variations in annual average OP activities and PM₁₀ source contributions by sites. Most sites, including traffic and industrial ones, show higher OP_{DTT} activities than OP_{AA}. Conversely, for the alpine valley sites, CHAM presents higher OP_{AA} than OP_{DTT}, while GRE-fr experiences similar levels of OP_{AA} and OP_{DTT} of the 2 OPs. Additionally, the average OP activities in every site are not proportional to the average PM concentration. For instance, CHAM and NIC had lower PM₁₀ concentrations but higher OP activities than other sites, while TAL showed high PM₁₀ concentrations but relatively lower OP activities.

The variations observed in the levels of PM₁₀ and OP across six sites can be attributed to distinctions in identified sources and their respective contributions. These disparities are contingent upon the unique typologies of each site, which are discussed in Weber et al., 2021. Further, we can observe a significant seasonality in the OP activities (Table S.1). Strong seasonality of OP in Alpine valley sites has been addressed in previous studies (Borlaza et al., 2021; Dominutti et al., 2023; Weber et al., 2018, 2021), with thermal inversions during winter increasing pollutants concentrations and OP activities compared to summer. Conversely, OP activities in cold and warm periods in other sites are not significantly different.

The PM₁₀ sources and their repartition vary among sites (Figure 3) because of the difference in typology and local activities. For instance, in the industrial site (PdB), two specific sources are identified: shipping emissions (HFO) with an annual mean contribution of 1.39 μ g m⁻³ and industrial sources at 0.86 μ g m⁻³. The urban background site TAL also appears to be influenced by industrial sources (2.34 μ g m⁻³), which might, however, be partly due to biases induced by the application of the harmonized receptor model protocol (Weber et al., 2019). Note that the application of a site-specific PMF procedure for this site leads to a much lower contribution of this source category but relatively similar contributions of other sources (Favez, 2017). GRE-fr, an urban background site in an alpine

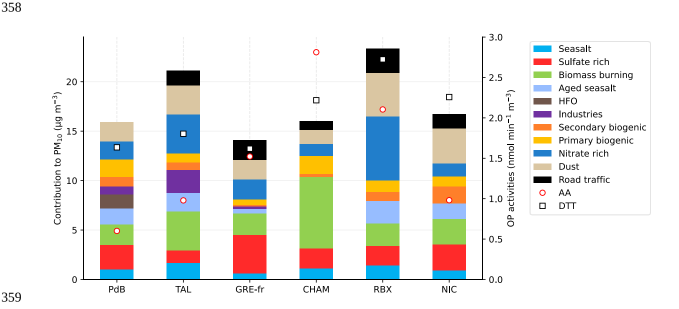
valley, presents significant long-range transport sources, with secondary sulfate contributing 3.90 µg m⁻³ followed by biomass burning at 2.21 µg m⁻³. As expected, biomass burning is an abundant source in CHAM, accounting for 7.28 µg m⁻³ of the PM contribution, while the traffic sites RBX and NIC displayed high contributions of traffic sources (at 2.43 μg m⁻³ and 1.45 μg m⁻³ respectively).

The presence of multicollinearity and homoscedasticity were tested to assess the data characteristic of every site. The only site with evidence of collinearity was NIC, where the VIF of the traffic source was equal to 5.0. For all other sites, VIF values are below 5, indicating limited collinearity among sources. This is expected, as the PMF analysis is constrained to avoid collinearity between sources. VIF values for each site can be found in Table S.2.

The presence of heteroscedasticity is commonly found when the dependent variable (or OP in this study) exhibits a large difference between the minimum and maximum values or when the error variance varies proportionally with an independent variable (PM₁₀ sources). The heteroscedasticity was assessed by applying the Goldfeld-Quandt test. Table 2 presents the p-values of the Goldfeld-Quandt test, indicating homoscedasticity of OP prediction when p >0.05. This test reveals that heteroscedasticity was detected in CHAM, GRE-fr, NIC for OP_{AA} and in CHAM and TAL for OPDTT (Table 2). We observed a large difference between the cold and warm periods for both OPAA and OPDTT in CHAM, similar to what was seen for OPAA in GRE-fr (Table S1), which can be the reason for the presence of heteroscedasticity. For NIC and TAL, there is an insignificant difference between the cold and warm periods, which indicates the presence of heteroscedasticity may be because of the relationship between the PM₁₀ sources and error variance. When heteroscedasticity is detected, unweighted regression for OP prediction according to sources may not accurately reflect the uncertainty of each source's intrinsic OP. The scatterplots representing the relationship between the regression analysis residuals and the fitted values (for observed OP) are available in Figures S.1 and S.2, Supplement.

Table 2. The p-value of the Goldfeld-Quandt heteroscedasticity test

	PdB	TAL	GRE-fr	СНАМ	RBX	NIC
AA	0.15	0.78	<< 0.001	<< 0.001	0.44	0.002
DTT	0.59	<< 0.001	0.189	<< 0.001	0.56	0.91



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Figure 3. The contribution of sources to PM_{10} and the OP activities in 6 sites. The left y-axis and bar show the contribution of PM sources in μg m⁻³. The right y-axis, circles and squares showed the mean OP_v activities in nmol min⁻¹ m⁻³, with red circle for OP_{AA} and black square for OP_{DTT} .

3.2. The performances of regression models

The 11 regression models, with or without weighting for some of them, were tested by comparing their performance metrics between the measured and reconstructed OPs. For each run (n = 500 iterations), the R^2 , RMSE, and MAE were computed for the testing and training dataset, resulting in 500 values for each performance metric. Figure 4 presents the mean R^2 values of the training data sets, the mean and the standard deviation of the testing datasets of the OP_{AA} models across the 500 sampling iterations, and Figure 5 presents the mean RMSE and MAE. The same result pattern was found for OP_{DTT}, as presented in the tables S.3, S.4, S.5, Supplement. The WLS, wPLS, wRidge, and wLasso models incorporated weighting, while the OLS, PLS, Ridge, Lasso, GLM, RF, and MLP models were unweighted.

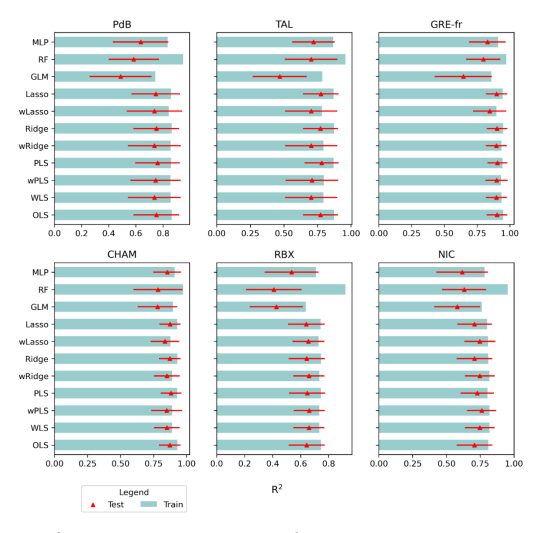


Figure 4. The R^2 of 11 OP_{AA} models in 6 sites. The mean R^2 of training data is shown in a blue bar, the mean R^2 of testing data is shown by a red triangle, and the red bar is the standard deviation of the R^2 of the testing data. The y-axis represents the models, and the x-axis denotes the R^2 values.

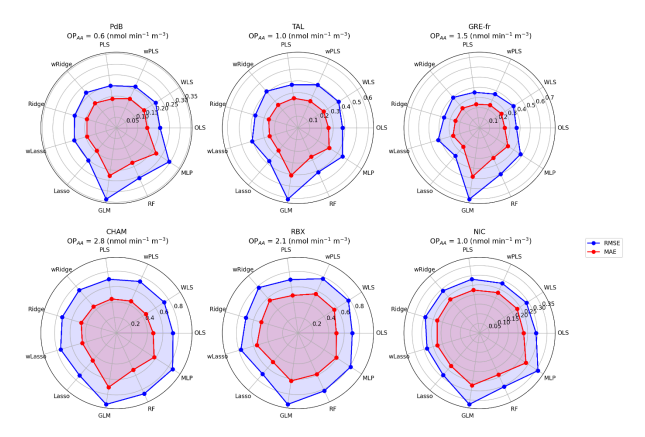


Figure 5. The MAE and RMSE of 11 OP_{AA} models in every site for the testing data. Blue and red lines present the RMSE and the MAE, respectively. The values in the figure are the mean of RMSE and MAE of 500 iterations.

OP predictions across all sites are statistically validated, with testing R^2 values observed in RBX, NIC, PdB, TAL, CHAM, and GRE-fr being 0.66, 0.76, 0.76, 0.78, 0.87, 0.90, respectively. The lowest mean test set RMSE values are 0.70, 0.28, 0.21, 0.37, 0.70, 0.31 nmol min⁻¹ m⁻³, respectively, for the same sites. The lowest mean test set MAE values are 0.49, 0.23, 0.14, 0.25, 0.45, and 0.21 nmol min⁻¹ m⁻³, respectively. Notably, the GLM model exhibits for all sites the lowest R^2 values and the highest RMSE (Table S.3, S.4, S.5, Supplement). These results strongly suggest that the relationship between OP_{AA} and PM₁₀ sources is not log-linear.

Differences in MAE, RMSE, and R² between the training and testing database for RF and MLP are significant across the sites. Notably, RF displays a large difference in R², with a gap of up to 0.6 in RBX (R² training: 0.92, R² testing: 0.27). Similar gaps were found in RMSE and MAE. RF consistently performed best on the training set, characterized by the highest R² and the lowest MAE and RMSE values, but had lower set test R² values than the other models (except GLM). Conversely, MLP exhibited training R² values comparable to other models but lower test R². These findings suggest overfitting: the flexible algorithms identify relationships in the training data that do not generalize to the testing data. This observation may be attributed to the limitations of data coverage, possibly failing to fully represent the underlying relationships, leading to poor performance in testing datasets (Benkendorf & Hawkins, 2020; Hawkins, 2004; Hernandez et al., 2006; Matsuki et al., 2016; Raudys & Jain, 1991; Stockwell & Peterson, 2002; Wisz et al., 2008). Pearce and Ferrier (2000) recommended that the minimum number of samples for robust performance should be over 250 for GLM model, while (Raudys & Jain, 1991) showed that the minimum number of sample are based on the complexity of the model and the number of predictors. Additionally, Harrell (2016) suggested that the number of predictors (PM sources) should be below the number of samples divided by 15, a threshold not reached in this analysis. For example, in NIC, the minimum number of samples should be 135 for the training set (9 PM sources x 15), while in total, we have only 107 samples. Therefore, we

can also recommend that, for optimal performance of RF, and MLP, the number of samples and PM sources should satisfy these thresholds.

The WLS, OLS, wPLs, wRidge, and wLasso models show more robust performances with fewer differences between the training and testing data. At most sites, there is very little difference between the R², RMSE, and MAE of OLS and Ridge, with or without weighting, and often PLS and Lasso as well. This consistency is observed even in the collinearity case of NIC, where VIF = 5. The difference between these models is a maximum of 0.06 in R², 0.01 in MAE and 0.1 in RMSE, indicating that these models work well for OP prediction. Nevertheless, it is worth noting that every model exhibits different assumptions that have to be respected. The assumption violations may lead to unreliable regression coefficients (intrinsic OP) even though the prediction is good (Cohen et al., 2002; Williams et al., 2013).

The best model for each site was selected based on both data characteristics (collinearity and heteroscedasticity) and testing data performance. For sites with collinearity, the Ridge, Lasso were considered most appropriate. For sites with heteroscedasticity, models with weights were considered the most appropriate. For sites with neither collinearity nor heteroskedasticity, OLS and PLS were considered most appropriate. Tables 3 and 4 present the best OP_{AA} and OP_{DTT} prediction models for each site. It follows that the best model is not necessarily the same one for both series of OP for a given site. As a rule, the model that exhibits the best performance metrics (the best model by error in Table 3 for OP_{AA} and Table 4 for OP_{DTT}) is suited to the best model chosen by data characteristics; therefore, choosing a model according to data characteristics help to more reliable in OP predictions.

Table 3. Criteria to select the best model for OPAA

	PdB	TAL	GRE-fr	СНАМ	RBX	NIC
Collinearity	No	No	No	No	No	Yes
Heteroscedasticity	No	No	Yes	Yes	No	Yes
Best model by	OLS/	OLS/	WLS/	WLS/	OLS/	wRidge/
characteristic	PLS	PLS	wPLS	wPLS	PLS	wLasso
Best by error	PLS	PLS	wPLS	wPLS	OLS	wRidge

Table 4. Criteria to select the best model for OP_{DTT}

	PdB	TAL	GRE-fr	СНАМ	RBX	NIC
Collinearity	No	No	No	No	No	Yes
Heteroscedasticity	No	Yes	No	Yes	No	No
Best model by	OLS/	WLS/	OLS/	WLS/	OLS/	Ridge/
characteristic	PLS	wPLS	PLS	wPLS	PLS	Lasso
Best by error	OLS	wPLS	PLS	wPLS	PLS	Ridge

3.3. Effect of the choice of a model on intrinsic OP

It is particularly important to try to define the best way of calculating the more accurate PM sources intrinsic OP and the contribution of sources to OP, since these values are fundamental inputs in all the works of large-scale modelling of OP with chemical transport models (CTM) (Daellenbach et al., 2020; Vida et al., 2024). Figures 6 and 7 show the variations of intrinsic OP for all the models, focusing on the results of NIC as an example. The evaluation of the 5 other sites is presented in Fig S.3 to Fig S.7 for OP_{AA} and Fig S.8 to S.12 for OP_{DTT}. The differences in equations, error term minimizations, and assumptions can explain the differences in intrinsic OP per µg of source among the eight regression models. While the R², RMSE, and MAE values are similar among models (except for GLM, RF, and MLP), the intrinsic OP values significantly differ between the models with and without weighting and between the linear and non-linear regression models. The average intrinsic OP of 500 iterations is discussed in this section since these values are usually used to calculate the contribution of the PM₁₀ source to OP in prior studies (Borlaza et al., 2021; Dominutti et al., 2023; Weber et al., 2018). The mean and standard deviation of intrinsic OP_{AA} and OP_{DTT} for the 6 sites are shown in Table S.6 and S.7, respectively.

Intrinsic OP_{AA} of PM₁₀ sources at NIC is the same between WLS and wRidge and between the OLS and Ridge, revealing that the moderate collinearity of the road traffic source did not affect the estimated intrinsic OP_{AA}. PLS sets the intrinsic OP_{AA} of some sources to zero, therefore producing slightly different results. Lasso regression sets the intrinsic OP_{AA} of some sources to zero and shrinks the estimates for all other sources toward zero. GLM produces intrinsic OP_{AA} values that represent a multiplicative change on the log scale, so they are not directly comparable to the other models. However, the direction and importance of the sources are similar to the other models. Whatever the model, road traffic appears as the source with the highest intrinsic OP_{AA}, followed by biomass burning, aged salt, salt and sulfate-rich sources, in NIC. Traffic and biomass burning sources have been similarly recognized as significant contributors to OP_{AA} in prior studies (Borlaza et al., 2021; Dominutti et al., 2023; Stevanović et al., 2023). The intrinsic OP of the dominant sources is stable, indicating that all these models could give the same information about the intrinsic OP of the main sources. Conversely, the differences are larger between models for the sources with small to very small intrinsic OP (MSA rich, primary biogenic, nitrate-rich, dust), whose intrinsic OP varies from positive to negative among models.

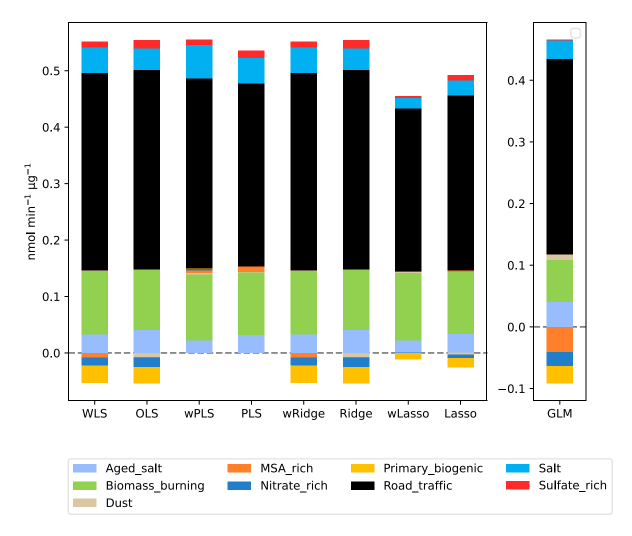


Figure 6. Intrinsic OPAA values of the different PM₁₀ sources at Nice were obtained with the different models.

The OP_{DTT} intrinsic values in NIC (Figure 7) display minimal variation among the WLS, wPLS. This consistency is linked to the absence of negative intrinsic values. On the other hand, even though there is the presence of moderate collinearity, wRidge still has the same result as WLS and wPLS. In line with the OP_{AA} results, the wLasso and GLM models exhibit distinct responses compared to the other models. The intrinsic OP_{DTT} of all sources varies depending on the presence or absence of weighting. While the WLS models tend to amplify the influence of some sources (aged sea salt, primary biogenic, sea salt, and sulfate-rich), the OLS reduces the intrinsic OP_{DTT} of these sources. Conversely, MSA-rich, nitrate, and road traffic sources undergo less influence in WLS but higher in OLS. Different from OP_{AA} , OP_{DTT} prediction shows more variation among models, highlighting the effect of choosing a model on evaluating the intrinsic OP_{DTT} of PM_{10} sources.

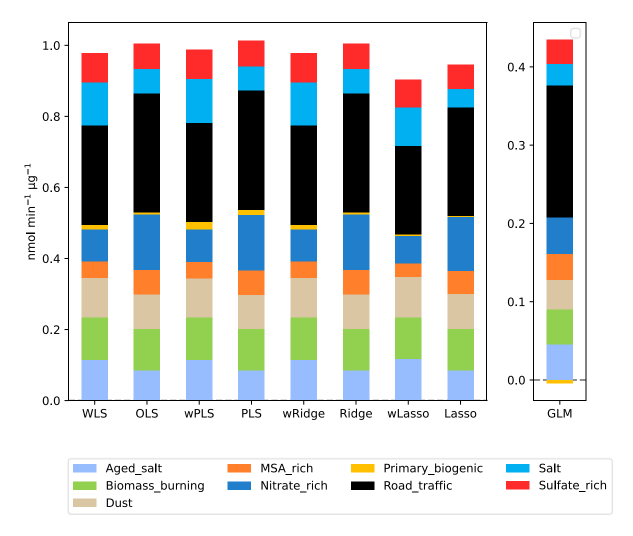


Figure 7. The variations of the intrinsic OP_{DTT} of the different $PM_{\underline{10}}$ sources at Nice were obtained with the different models.

The comparison of intrinsic OP among regression models in NIC demonstrated that OP_{DTT} and OP_{AA} intrinsic values exhibit variation across different models with and without weighting, illustrating that the choice of the model significantly influences the values obtained for intrinsic OP of $PM_{\underline{10}}$ sources (A similar pattern is observed for all other sites and shown in Fig S.3 to Fig S.7 for OP_{AA} and Fig S.8 to S.12 for OP_{DTT}). Because of the difference in OP-intrinsic OP across models, a comparison between the best-performing and most commonly used models (OLS) is presented in the following section to elucidate the advantage of choosing a model based on data characteristics (section 3.4).

3.4. Comparisons between the best site-specific model and OLS

In this section, the intrinsic OP of the best model is selected for each site as discussed in Section 3.2, and the intrinsic values of each source are compared to the ones returned by the OLS model. The OLS model is used as a representative of usual practices that do not consider the database characteristics (Williams et al., 2013). Each PM_{10} source's average intrinsic OP value is calculated from all the 500 bootstrapping iterations for all sites where that particular source is identified. Intrinsic OP values obtained in this way from the best model (the best model presented in Table 3 for OP_{AA} and Table 4 for OP_{DTT}) encompassing all six sites are called **intrinsic OP of the best model**, and the intrinsic OP values derived from the OLS from all six sites are called **intrinsic OP of the reference model**.

A meaningful comparison of the two series of intrinsic values requires two conditions. First, intrinsic OP values should be consistent across all sites. While recognizing that intrinsic OP values depend on diverse factors, we assumed the sites share fairly uniform PM₁₀ chemical source profiles in France. This is demonstrated by evaluating the Pearson distance and standardized identity distance similarity indicators of the source chemical profiles (Belis et al., 2015; Weber et al., 2019), and Figure S.13 indicates consistent profiles of sources for the 6 sites. Consequently, we could expect to observe minimal divergence in intrinsic OP values among these sites. Second, we postulate that negative intrinsic OP values are possible since previous studies have reported that total PM₁₀ intrinsic OP can be modulated due to the synergetic/antagonistic effects involving, for example, soluble copper, quinones, and bacteria (Borlaza et al., 2021; Pietrogrande et al., 2022; Samake et al., 2017; S. Wang et al., 2018; Xiong et al., 2017). Samake et al. (2017) demonstrated that the presence of bacterial cells in aerosol decreases the redox activity of Cu and 1,4-naphthoquinone, with a maximum decreasing of 60% compared to the oxidative reactivity considered individually. Pietrogrande et al. (2022) indicated that the mixture of Cu, Fe, 9,10phenanthrene quinone and 1,2-naphthoquinone reduces the rate consumption of AA and DTT, up to 50% depending on the quantity of each chemical. Wang et al. (2018) reported that the mixing of Cu and naphthalene secondary organic aerosol (SOA) and phenanthrene SOA only got half of DTT rate consumption compared to the consumption when considered separately. Xiong et al. (2017) showed the presence of antagonists in the interaction of Fe and quinones, nevertheless, much lower than those in the other studies (under 10%). These references reported that the antagonistic effects of a mixture can significantly reduce the consumption rate of OP_{DTT} and OP_{AA}, and this impact varies widely from 10% to 60% depending on the type of chemical species and the quantity of each species in the mixture. These last studies showed that the impact of synergistic and antagonistic effects cannot exceed 60% of the intrinsic OP value when assessed independently for each chemical. Consequently, we consider here that the intrinsic OP value of an individual site for a given source could be negative only within a range of at most 60% of the mean combined intrinsic OP value of this source across all sites. Negative intrinsic OP exceeding this criterion may result from the mathematical construction of the model. The comparison of intrinsic OP_{AA} of the best and reference model is presented in 3.4.1 and that of OP_{DTT} is shown in 3.4.2.

505 3.4.1. OP_{AA} activities

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The results of the comparison of OP_{AA} intrinsic values (Figure 8 and Table S.8) show that the anthropogenic sources get the highest intrinsic OP values in both the best and reference models. Among these sources, road traffic appears as the most prominent potent fraction, followed by biomass burning, HFO, and industrial. These results are aligned with prior research (Calas et al., 2019; Daellenbach et al., 2020; Dominutti et al., 2023; Fadel et al., 2023; Fang et al., 2016; in 't Veld et al., 2023; Weber et al., 2018; Zhang et al., 2020) which has highlighted the sensitivity of OPAA to concentrations of metals, black carbon, and organic carbon. The differences between the best and reference models were insignificant for these sources, demonstrating that the best and reference models consistently captured similar patterns for the most critical sources of OP activities.

However, the interquartile ranges (IQR) of the intrinsic OP values are consistently narrower for the best models across all sources, accounting for less divergence in intrinsic OP values across sites. Moreover, the median intrinsic OP values obtained from the best model closely approximated the mean values, indicating the absence of extreme intrinsic OP values. For instance, in the case of road traffic, the mean and median values were 0.24 and 0.23 nmol min⁻¹ µg⁻¹, respectively. Conversely, the reference model exhibited a large difference between the mean and median values, implying lower consistency across sites and sampling iterations. The same result was observed in biomass burning source, in which the median and mean intrinsic OP in the best model had fewer discrepancies. Further, the biomass burning intrinsic OP in GRE-fr of the best model is more consistent with those in other sites (best: 0.30 nmol min⁻¹µg⁻¹, reference: 0.35 nmol min⁻¹µg⁻¹).

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When considering sources with low intrinsic OP, the variability can be larger between the two methods. As an example, for the sulfate-rich sources, the median intrinsic OP values were positive (0.002 nmol min⁻¹ µg⁻¹), while the mean intrinsic OP values were negative (-0.008 nmol min⁻¹ μ g⁻¹). The mean intrinsic OP in the best model exhibited fewer negative values in individual sites than in the reference model (for aged salt, salt, primary biogenic, MSA rich, sulfate-rich and nitrate-rich). In addition, the best model showed the less disparate intrinsic OP among individual sites, for instance, the aged salt sources in GRE-fr and the primary biogenic and salt sources in CHAM. highlighting the advantage of considering the data in model selection. For example, Furthermore, the best model displayed an intrinsic OP meaningful in terms of geochemicals, which showed in the source of salt, primary biogenic, sulfate-rich. For instance, in the reference model, Tthe mean average intrinsic OP values of the primary biogenic source revealed a negative intrinsic OP in NIC (-0.03 nmol min⁻¹ μ g⁻¹), the intrinsic OP of salt in GRE-ft (-0.07 nmol min⁻¹ μ g⁻¹) as well as the sulfate-rich source in CHAM (-0.05 nmol min⁻¹ μ g⁻¹). This negative value represented represented a 100% reduction compared to the mean intrinsic OP of all sites. Moreover, In the OLS model, the negative intrinsic OP was observed in NIC (Primary biogenic), and some extreme values in GRE-fr (aged salt, salt), CHAM (salt, primary biogenic, MSA-rich), NIC (where heteroscedasticity was presented) in the OLS model, underscores that the model assumptions on data characteristics proving false could impact the accuracy of OP prediction. Consequently, these results highlight the advantage of considering the data in model selection.

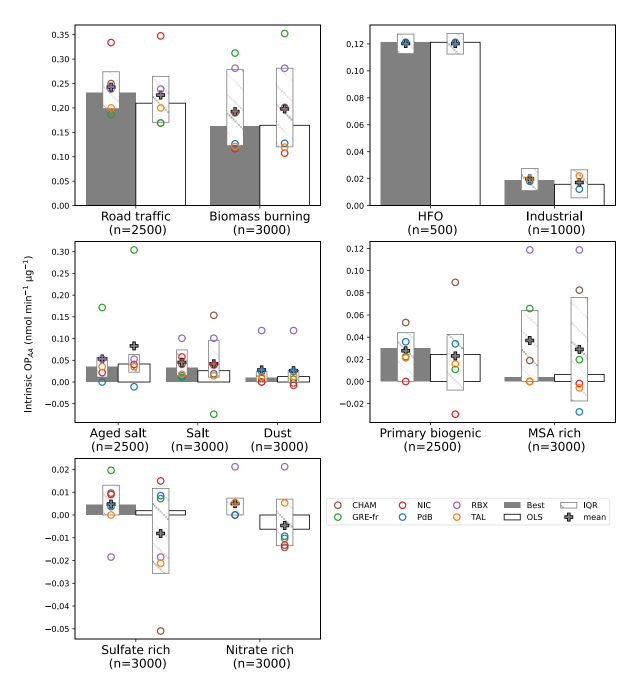


Figure 8. Intrinsic OP_{AA} estimated by the best and the reference methods in the 6 sites. The y-axis represents the intrinsic OP values in nmol min⁻¹ μ g⁻¹, the x-axis represents the sources. The grey bars are the median intrinsic OP values of the best models in the 6 sites (n = 500 bootstrapping * number of sites where the given source is detected) for each source. The white bars are the same median intrinsic OP values for the reference (OLS) model. The grey plus symbol represents the mean of OP intrinsic OP values. The hatched bars are the interquartile ranges of the intrinsic OP values. The dots represent the mean intrinsic OP of all sites, including grey – Chamonix, green – Grenoble, red – Nice, blue – Port-de-Bouc, purple – Roubaix, and orange-Talence.

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The detailed comparison of intrinsic OP_{AA} between the best and reference models is categorized into four groups and discussed in detail in section S9. These groups include (1) anthropogenic sources without nitrate and sulfate (road traffic, biomass burning, HFO, industrial), (2) natural inorganic sources (aged sea salt, sea salt, dust), (3) biogenic sources (primary biogenic, MSA rich), and (4) nitrate and sulfate-rich sources.

3.4.2. OP_{DTT} activities

Similar to OP_{AA}, for OP_{DTT} the IQR of the best model is narrower for most of the sources than the IQR of the reference model (OLS). Except for the road traffic, industrial, and MSA-rich, the IQR is slightly higher in the best model (Figure 9 and Table S.9). In the two models, the mean intrinsic OP is essentially unchanged, where the traffic is the most critical source (0.27±0.10), followed by HFO (0.18±0.01), biomass burning (0.12±0.03), dust (0.12±0.07), primary biogenic (best: 0.10±0.06, reference: 0.12±0.08) and MSA rich (best: 0.11±0.09, reference: 0.09±0.09). The remaining sources, such as sea salt, sulfate rich, industrial, and nitrate rich, show a negligible contribution to OP_{DTT} with an intrinsic OP_{DTT} from 0.02 to 0.08. The minimum difference between the two models in the dominant sources again confirms the conclusion in the OP_{AA} comparison, demonstrating the similar pattern of the best and the reference model in the most crucial sources of OP. For both best and reference, OP_{DTT} activities showed sensitivity to more sources than OP_{AA}, as discussed in previous studies (Borlaza et al., 2021; Calas et al., 2019; Dominutti et al., 2023; Fadel et al., 2023). The traffic, HFO and biomass burning sources highlighted in are the most contributing to OP_{DTT} to organic compounds, as mentioned in . The intrinsic OP of dust and MSA rich have been shown to vary in the literature, indicating the different effects of the compositions to generate DTT ROS.

While the best and reference models give the same mean intrinsic OP_{DTT} of all sites, the mean OP_{DTT} at each individual site can vary substantially between the two models. The best model exhibited the positive intrinsic OP for all sources, while the reference model displayed negative intrinsic OP in RBX (MSA-rich and sulfate-rich). Especially in the case of sulfate-rich in RBX, the negative intrinsic OP in the reference model passed the threshold of negative value, which presented a 110% reduction compared to the mean intrinsic OP of all sites. This is also found in the OP_{AA} comparison, which confirmed that the best model generates a geochemical meaningful OP intrinsic. In addition, the best model exhibited consistent intrinsic OP across sites, especially for the source of dust, salt, primary biogenic, sulfate-rich in TAL (heteroscedasticity is presented in this site), where intrinsic OP in TAL in the best model is more similar to the other sites. For instance, the reference model presented that the intrinsic OP in TAL is 0.20 nmol min⁻¹ µg⁻¹, far from the mean of all sites (0.07 nmol min⁻¹ µg⁻¹). We observed the same for OP intrinsic of nitrate-rich source in CHAM (where the heteroscedasticity is detected), which displayed the less dissimilar of CHAM with the other site in the best model. This again validates the conclusion in OP_{AA} comparison, demonstrating that respecting model assumption is essential to obtain a robust OP SA result.

For the sulfate rich source, the reference model showed a negative intrinsic OP_{DTT} in RBX (0.09 nmol min $^{\downarrow}\mu g^{-}$), while the best model showed an intrinsic of 0. On the other hand, the reference model presents that the intrinsic OP in TAL is 0.20 nmol min $^{\downarrow}\mu g^{-}$, far from the mean of all sites (0.07 nmol min $^{\downarrow}\mu g^{-}$). The best model, conversely, shows a more consistent intrinsic OP in TAL compared to the other sites. A similar result was also found in primary biogenic sources, where the reference model overestimates the intrinsic OP of this source in TAL compared to the other sites. The reason is that heteroscedasticity was detected in TAL, which does not satisfy the assumption of OLS.

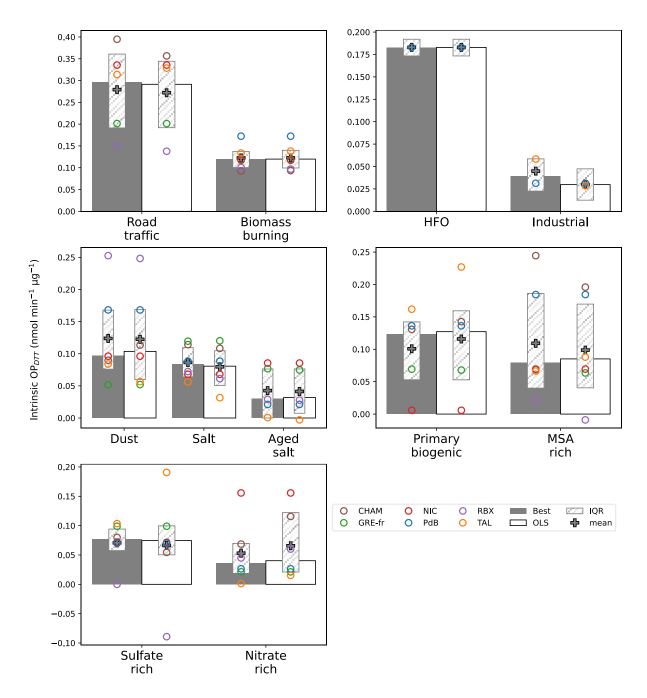


Figure 9. Intrinsic OP_{DTT} was estimated by the best and the reference methods in the 6 sites. The y-axis represents the intrinsic OP values in nmol min⁻¹ μ g⁻¹, the x-axis represents the sources. The grey bars are the median intrinsic OP values of the best models in the 6 sites (n = 500 bootstrapping * number of sites where the given source is detected) for each source. The white bars are the same median intrinsic OP values for the reference (OLS) model. The grey plus symbol represents the mean of intrinsic OP values. The hatched bars are the interquartile ranges of the Intrinsic OP values. The dots represent the mean intrinsic OP of all sites, including grey – Chamonix, green – Grenoble, red – Nice, blue – Port-de-Bouc, purple – Roubaix, and orange-Talence.

The comparison of intrinsic OP between the best models and the reference model highlights the importance of considering the database characteristics when selecting a model for OP SA. For all the datasets studied here, using the best model for each site delivered more robust results with reduced uncertainty, reduced differences in intrinsic

OP across sites, and provided a more geochemically meaningful intrinsic OP. The recommendation for selecting a model based on the characteristics of the database is presented in section 3.5.

3.5. Guidelines for the selection of regression model for OP SA.

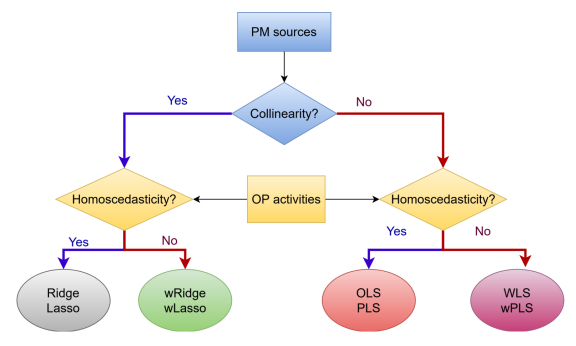


Figure 10. Workflow in model selection considering the characteristics of data

Our results have highlighted the benefits of choosing a model that matches the characteristics of the data to improve the robustness of OP SA method. For this reason, this section develops a workflow to help make model selection decisions. Before selecting a regression for OP SA, the first question is whether the $PM_{\underline{10}}$ sources are collinear and the second is whether the residual variance of the regression between OP and $PM_{\underline{10}}$ mass is constant. These two questions represent the characteristics of $PM_{\underline{10}}$ sources and OP activities, which vary according to the study site.

For data exhibiting collinearity between sources and generating a residual variance that varies according to the value of the PM_{10} sources, weighted regularisation regression can help to reduce collinearity and to match the model assumption about the residual. On the other hand, the unweighted Ridge and Lasso are introduced for data showing collinearity and homoscedasticity. Additionally, data with no collinearity are suitable for OLS and unweighted PLS in the case of homoscedasticity, while WLS, weighted PLS are used for data with heteroscedasticity.

If the number of predictors (PM_{10} sources) is below the number of samples divided by 15, RF and MLP can also be employed to capture possible non-linear relationships between the OP and PM_{10} sources. However, cross-validation must be used to ensure that there is no over-fitting. In addition, these models do not estimate intrinsic OP (nmol min⁻¹ μ g⁻¹) but only the importance of each PM_{10} source to the OP prediction. This is a large drawback since the intrinsic OP of sources is a must for the modelling effort of OP with CTM. However, RF and MLP could be useful for OP prediction in the case of larger datasets generated by online instruments.

For each data characteristic there is more than one model that suits. Out-of-sample performance metrics should be employed to identify the most accurate of these models.

Finally, these techniques of OP apportionment could not be well performed with uncertain PMF-derived sources. The PMF results sometimes do not adequately represent PM mass concentration for several reasons, such as the lack of a trace species to identify a source, an insufficient sample size, the source contribution being too small to

be identified (under 1%), or collinearity matters. The important information could be missed because of these problems in PMF implementation, which is apprehended by the model's low accuracy. Our study did not encounter this problem since the PMF is harmonized and performed according to European recommendations which could well perform the regression technique and allow to obtain a very satisfactory successive OP modelled in comparison to observations after regression techniques (R² from 0.7 to 0.9). However, this problem could potentially happen, and for these cases, we could recommend either subtracting the total source contribution from PM mass concentration to get a part that PMF cannot simulate. The information in this part may contain vital sources. Alternatively, it is possible to re-execute the PMF to validate the result and ensure the robustness of the chemical profile and the contribution of sources.

Limitations and perspectives of the study:

- This study compares eight regression models but is not exhaustive; further research could add more regression techniques to evaluate result variations across models. The potential techniques that could be applied for OP SA are gradient boosting techniques for resolving regression models, or supervised machine learning techniques which allows the investigation of linear and non-linear regression relationships. However, the consistently strong performance of ordinary linear regression across six locations in France suggests that there may be little to gain from applying more complex models in areas with similar PM₁₀ sources.
- PMF coupled with a regression model remains a popular approach for OP SA. Notably, the uncertainties
 in PMF are typically addressed in chemical profiles, but not in contributions. Incorporating uncertainty
 from variations in contribution into models could enhance their robustness compared to relying only on
 absolute PMF results.
- Observations ranged between 100 and 200 samples at each site, which may be insufficient to obtain fair performance of GLM, decision trees and neural network models. Such a number of samples is sufficient to address SA through PMF model for offline analyses. Therefore, such study outlines well the limitations of GLM, RF, MLP for such types of datasets. Future investigations should be performed in an extended dataset, such as long term or real time measurement data, to investigate the performance of such machine learning algorithms. Observations ranged between 100 and 200 samples at each site, which may be insufficient to obtain a fair performance of GLM, decision trees and neural network models even though this number of samples is sufficient to address SA through the PMF model for offline analyses. Therefore, this study outlines well the limitations of GLM, RF, and MLP for offline datasets. Future investigations should be performed in an extended dataset, such as long-term or real-time measurement data, to investigate the performance of machine learning algorithms.
- This study only focused on the two most popular OP assays of PM₁₀ (OP_{DTT} and OP_{AA}). However, there are actually various OP assays, such as OP_{DCFH}, OP_{OH}, OP_{FOX}, OP_{GSH}, OP_{ESR} and different sizes of PM (PM1, PM2.5, PM5). Further research should include more OP assays, which can be helpful in evaluating the performance of various regression models for different OP and different PM sizes.
- This study used the analytical uncertainty as the weighting for the weighted model. However, the weighting can be selected based on different ways, as reported by Montgomery et al. (2012): (1) Prior information from the theoretical model, (2) Using the residual extracted from the OLS model, (3) The selecting of weighting based on the uncertainty of instrument if the dependent variable measured by a different method and (4) If the dependent variable is the average of different observations, the weighting selected based on the error of these observations.

4. Conclusion

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674 The results of the OP SA marked an important milestone as they were revealed for the first time through the use of eight regression models, including OLS, WLS, PLS, GLM, Ridge, Lasso, RF and MLP. This in-depth analysis 675 676 was carried out on a complete set of data collected from six sites with different characteristics. The approach of selecting a suitable model for each site based on specific data characteristics resulted in a more consistent intrinsic 677 OP across sites, in stark contrast to the variation observed when using the basic OLS model. The revelations of the 678 679 study have provided concrete recommendations for the judicious selection of an appropriate regression model based on the unique characteristics of the dataset. These guidelines should help to improve the accuracy of OP 680 681 assessments and contribute to the refinement of air quality assessment methods. In addition, the implications of 682 this research extend to the implementation of OP monitoring as a new measure of air quality, particularly on European supersites. As this initiative aligns with the ongoing revision process of the European Directive 683 684 2008/50/CE, the study's findings assume a pivotal role in shaping the methodologies underpinning air quality 685 assessments at a broader regulatory level.

Code availability

The software code could be made available by contacting the corresponding author upon request.

Data availability

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

690 **Author contributions**

- VDNT performed the data analysis for the OP source apportionment setup. GU, JLJ mentoring, supervision, and
- validation of the methodology and results. IH, PD, and VDNT worked on the result visualization. OF, JLJ, and
- 693 GU acquired fundings for the original PM sampling and analysis. VDNT wrote the original draft. All authors
- reviewed and edited the manuscript.

Competing interests

The authors declare that they have no conflict of interest.

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