



Full length article



High-resolution modelling of organic aerosol over Europe: exploring spatial and temporal variability and drivers

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ABSTRACT

Organic aerosol (OA) is a major component of atmospheric particulate matter (PM), affecting both human health and climate. However, high-resolution estimates of OA exposure needed for exposure analysis remain scarce. Here, we integrate a chemical transport model (CAMx) with a random forest (RF) machine learning approach to bias-correct and downscale daily OA concentrations across Europe. CAMx OA simulations at ~15 km resolution show moderate agreement with observations ($r = 0.55$). By combining these outputs with high-resolution land-use data and training the RF model on ~48,000 daily OA measurements from 137 sites, prediction accuracy improved ($r = 0.65$), with ~15% reduction in root mean square error. The resulting maps provide European daily OA concentrations at ~250 m resolution for alternate years from 2011 to 2019. The model captures key spatial features, including elevated OA in the Po Valley, Southeastern, and Central Europe, as well as intracity variations due to local hotspots. Seasonal analysis reveals higher concentrations in winter, while long-term trends indicate a general decline in OA levels. Exposure estimates show that half of the European population experiences OA levels above $3 \mu\text{g}/\text{m}^3$, and ~50 million people are exposed to more than $5 \mu\text{g}/\text{m}^3$, which is the current guideline level recommended by the world health organization for total $\text{PM}_{2.5}$. These high-resolution OA maps offer vital critical support for epidemiological research and air quality policy.

1. Introduction

Atmospheric aerosols have adverse effects on both human health and the environment. These aerosols have varying chemical composition, morphology, and size, which determine their impacts on health and climate. Organic aerosol (OA) contributes 20–90% of particulate matter (PM) mass, while its proportion varies by regions and seasons (Jimenez et al., 2009). In Europe, OA accounts for approximately 50% of total

aerosol mass (Bressi et al., 2021), with higher contribution in winter compared to summer (Chen et al., 2022). OA can be directly emitted from combustion activities like biomass burning (BB) and fossil fuel combustion or formed in the atmosphere as secondary aerosol through gas-to-particle conversion (Kanakidou et al., 2005). BB is the largest source of OA in Europe in winter, while biogenic emissions represent a major OA source in summer (Jiang et al., 2019). The contribution of vehicular sources is low at the regional scale, yet much higher at urban locations with high traffic density (Chen et al., 2022). Stricter air quality standards that target specific aerosol components are becoming increasingly important from a regulatory perspective. Given the

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different toxicity of various aerosol constituents, it is crucial to prioritize regulations on the most harmful components, such as organic compounds.

Epidemiological studies have firmly established the link between air pollution and a range of adverse health effects, including respiratory, cardiovascular and cerebrovascular diseases. PM alone is estimated to cause 4.2 million premature deaths annually (Cohen et al., 2017). Among PM components, Organic carbon (OC) or OA have been specifically associated with adverse health effects. Elevated OC levels have been linked to increased hospital admissions for myocardial infarction and diabetes (Zanobetti and Schwartz, 2009). Secondary organic aerosol (SOA) shows an even stronger association with cardiorespiratory mortality than total PM_{2.5}, with mortality rates up to 6.5 times higher attributed to SOA exposure in the U.S. (Pye et al., 2021). In the UK, studies have also linked carbonaceous particles particularly those from the wood burning to non-accidental and respiratory deaths (Piper et al., 2024). While these findings underscore the health risks of OA, most focus on short-term effects or are limited to the U.S. context. There remains a critical need for large-scale, long-term, high-resolution OA data across other regions to better assess its chronic health impacts.

OA levels are typically estimated using ground-based observations, chemical transport models (CTM), and satellite-derived products. Ground observations offer high accuracy but limited spatial coverage, while satellite products often lack sufficient temporal and chemical resolution, limiting their direct use in exposure assessments and epidemiological studies (El Haddad et al., 2024). Satellite data can provide spatially continuous data but retrieving aerosol composition from satellites remains challenging due to algorithmic limitations, though ongoing efforts are improving these methods (Chen et al., 2024; van Donkelaar et al., 2019). CTMs provide continuous, continent-scale OA fields, yet their accuracy is constrained by uncertainties in input data, emissions, and parameterizations, and their coarse resolution is insufficient to capture local primary emissions. Hybrid approaches that integrate CTM outputs with local land-use information and satellite products have successfully improved high-resolution predictions for regulated and emerging pollutants, including PM, NO₂, O₃, and ultrafine particles (Gariazzo et al., 2020; Jianyao et al., 2025; Li et al., 2024; Shen et al., 2022; Upadhyay et al., 2024). However, extending these approaches to predict specific PM components, such as OA, remains limited. Several studies in North America have generated OA maps at regional scales, demonstrating the potential of these hybrid methods (Chen et al., 2020; Weagle et al., 2018). In Europe, recent applications combining CTMs and machine learning have shown promising results for mapping trace metals like copper, a tracer of brake-wear emissions (Upadhyay et al., 2025). Yet, comprehensive, high-resolution maps of OA across Europe using hybrid modeling approaches are still lacking. In a recent study, we applied a machine-learning framework to estimate OA components (HOA, BBOA, and OOA) at dense OC monitoring networks, achieving prediction accuracies with R² values of 0.58–0.75 and expanding the source-apportioned OA database by a factor of four (Jouanny et al., 2025). That approach combined CTM-simulated OA components, land-use variables, and on-site OC concentrations, and was trained on source-apportioned OA from aerosol mass spectrometer (AMS) and aerosol chemical speciation monitor (ACSM) observations. While highly valuable for providing source information at sites where only total OC is routinely measured, the predictions are limited to monitoring locations and do not yield gridded OA fields suitable for large-scale exposure assessment.

In the present study, our goal is to develop spatially continuous, high-resolution, daily total OA maps across Europe for alternate years between 2011 and 2019 (2011, 2013, 2015, 2017, and 2019). To achieve this, we integrate CTM predictions for OA and co-emitted pollutants, and land-use features in a machine learning framework trained on an extensive database of ground-based OC and online OA measurements to predict European OA maps. The resulting OA fields provide continent-wide coverage at a resolution and temporal frequency appropriate for

epidemiological and exposure analyses from regional to local scales.

2. Datasets and methodology

2.1. Organic aerosol observation database

We compiled a comprehensive OA observation database to ensure broad spatial and temporal coverage across Europe. Data were sourced from in-house measurements, research collaborators, open-access datasets (e.g., EBAS: <https://ebas-data.nilu.no/>; last access: December 20, 2025), and federal monitoring agencies. The dataset includes OC measured via offline filter-based thermal-optical analysis, primarily using Sunset Lab instruments and the EUSAAR₂ protocol (Cavalli et al., 2010), as well as OA measured using Aerosol Mass Spectrometers (AMS) and Aerosol Chemical Speciation Monitors (ACSM) (DeCarlo et al., 2006; Fröhlich et al., 2013; Jayne et al., 2000; Ng et al., 2011). We converted OC to OA using a factor of 1.5. Uncertainties associated with this conversion are addressed in Section 2.4.

The final dataset comprises nearly 48,000 daily OA measurements from 137 unique sites across five years (2011, 2013, 2015, 2017, 2019). About 92% of data points are OA estimates converted from OC at 119 sites, while the remaining 8% are direct OA measurements from 18 sites (Fig. 1; site details in Table S1). The dataset spans a wide range of locations, including high- and low-concentration regions as well as urban, suburban, rural, and high-altitude sites making it one of the most comprehensive OA monitoring efforts to date. However, coverage remains limited in certain regions, including Southeastern Europe, remote areas, and high mountain zones. Despite these gaps, the time and spatial resolution of this database represent a key strength of our modeling framework, enabling improved prediction accuracy while also offering a valuable resource for future air quality and health studies.

2.2. Comprehensive air quality model with extensions (CAMx) model

We used the CAMx, version 6.5 (ENVIRON, 2018), to simulate the mass concentrations of major air pollutants across Europe for the years 2011, 2013, 2015, 2017, and 2019. The model domain spans 15°W–35°E longitude and 35°–70°N latitude, with a spatial resolution of 0.125° × 0.25° (latitude × longitude) and 14 vertical layers extending from approximately 20 m to 7000 m above sea level. Gas-phase chemistry was modeled using the Carbon Bond 6 Revision 2 (CB6r2) mechanism (Hildebrandt Ruiz and Yarwood, 2013). The ISORROPIA thermodynamic model was applied to partition inorganic aerosols (sulfate, nitrate, ammonium, sodium, and chloride) between gas and particle phases. OA was modeled using a modified volatility basis set (VBS) module, which distinguishes primary and secondary components from biomass burning, road traffic, other anthropogenic activities, and biogenic sources, estimating primary and secondary biomass burning, hydrocarbon-like, anthropogenic and biogenic OA components. Details on the modified VBS scheme and its parameterization are available in Jiang et al. (2019).

Anthropogenic emissions of major gaseous and particulate pollutants, including POA and precursors of oxygenated OA (OOA), were obtained from the high-resolution European emission inventory CAMS-REG-v4 (Copernicus Atmosphere Monitoring Service-Regional inventory, version 4) developed by TNO (Kuenen et al., 2022). The CAMS-REGv4 version used is referred to as “ref 2”. This version has adjusted particulate matter (PM) emissions for residential combustion to include condensable PM emissions following the approach by Denier van der Gon et al. (2015) and further discussed by Simpson et al. (2022). This inventory includes emissions from public power, industry, transport, residential combustion, agriculture, and other source categories. Non-methane volatile organic compounds (NMVOC) and PM speciation for all emission sources followed the chemical profiles proposed by TNO. Biogenic emissions (isoprene, monoterpenes, and sesquiterpenes), which are critical precursors for SOA, were estimated using the PSI

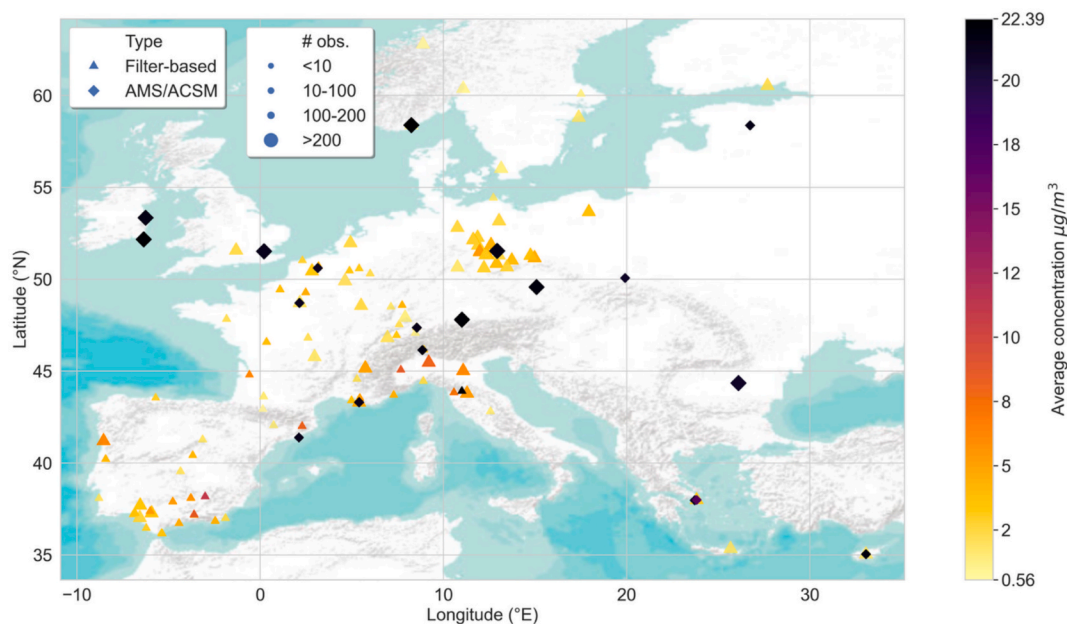


Fig. 1. Locations of organic aerosol (OA) observation sites over Europe. The triangles represent filter-based measurements, and diamonds represent AMS or ACSM (online) measurements. The marker size is proportional to the number of observations available at each location, and the color scale shows the mean OA concentration ($\mu\text{g}/\text{m}^3$) at each location for the whole measurement period. Background map shows land and water areas with terrain. .
Source: © Esri, Source: USGS, Esri, TANA, DeLorme, and NPS

model developed at the Laboratory of Atmospheric Chemistry, Paul Scherrer Institute (Andreani-aksoyoglu and Keller, 1995; Jiang et al., 2019).

Meteorological inputs for the CAMx model were generated using the Weather Research and Forecasting (WRF) model version 3.7.1 (Skamarock et al., 2008) driven by ERA5 reanalysis data from the European Centre for Medium-Range Weather Forecasts (ECMWF) (Dee et al., 2011). The WRF setup and parameterization were previously validated for the same modeling domain (Jiang et al., 2019). Chemical initial and boundary conditions were obtained from the global CAM-Chem model (Community Atmosphere Model with Chemistry) and converted to CAMx format using the MOZART2CAMx preprocessor (v3.2.1) (Emmons et al., 2010). Total column ozone data were sourced from NASA's Total Ozone Mapping Spectrometer (TOMS), and photolysis rates were calculated using the Tropospheric Ultraviolet and Visible (TUV) Radiation Model, version 4.8.

2.3. Machine learning model training

We used the Random Forest (RF) model (Breiman, 2001), a widely validated tree-based machine learning algorithm, to generate bias-corrected and downscaled OA predictions. We selected RF as the primary modeling approach after testing several algorithms, including gradient boosting (XGBoost) and bagging regressors. RF provided more stable performance across sites, slightly lower variance, and slightly higher predictive accuracy for our dataset. They are also less sensitive to noise, outliers, and heterogeneous feature response relationships, which is advantageous for OA observations collected across different monitoring environments. While gradient boosting is conceptually appealing for bias correction, achieving comparable performance required extensive hyperparameter tuning and substantially increased computational cost. The RF model used our large OA observation database as the target variables and CAMx simulated OA components and other pollutants, meteorological parameters from WRF, and high-resolution land use parameters as predictors. We processed daily OA concentration data to construct a robust target variable, filtering out sites with fewer than 30 time points and observations below $0.1 \mu\text{g}/\text{m}^3$ (considered below the detection limit). The final dataset for RF modeling included 47,800 daily

OA observations from 137 locations, spanning all months, days of the week, and five alternate years between 2011 and 2019, capturing the temporal variability present in the data. The problem is treated as a tabular prediction task; temporal correlations are not explicitly modeled. The list of RF predictors with their description is given in Table S2. The CAMx OA components (HOA, BBOA, PBIS, PAS, PBBS, PRTS) are used individually as predictors in the RF model, together with other covariates. The sum of all components provides total CAMx OA, which is used for validation against observations (Fig. 2a), and not as model input. Land use variables include land use categories (agriculture, industry, transport), road type and length, and population density. Table S2 also provides detailed names and descriptions of all the land use variables used with their source, spatial and temporal resolution, and buffer size, along with the list of other predictors. These land use parameters were obtained from various sources including satellite products and government databases. They were available at resolutions from 25 m to 1000 m for different variables and we re-gridded them to a uniform spatial resolution of ~ 250 m for modelling purposes. We also used meteorological parameters such as temperature, boundary layer height, wind speed, and atmospheric surface pressure obtained from the WRF model. The CAMx concentration fields set the background for spatial and temporal distributions of OA, while high-resolution land use inputs enabled model downscaling. We treat the problem as a tabular data prediction task, as such we do not explicitly model time correlations.

We optimized the RF model using grid search, systematically testing combinations of hyperparameters to identify those yielding the most accurate predictions. Model evaluation followed a Leave-One-Station-Out Cross-Validation (LOSO-CV) approach, using mean squared error (MSE) as the performance metric. In each iteration, one site was held out as a test set, while 20-fold group cross-validation in the inner loop optimized the RF hyperparameters and assessed performance on unseen locations. Once LOSO-CV predictions were generated, the best hyperparameters were selected using the inner loop with all training data. These were then applied to compute final LOSO predictions for model evaluation. The final model was trained using the optimal hyperparameters and the full training set. To ensure model reliability, we monitored the outer loop's CV error and verified the stability of selected

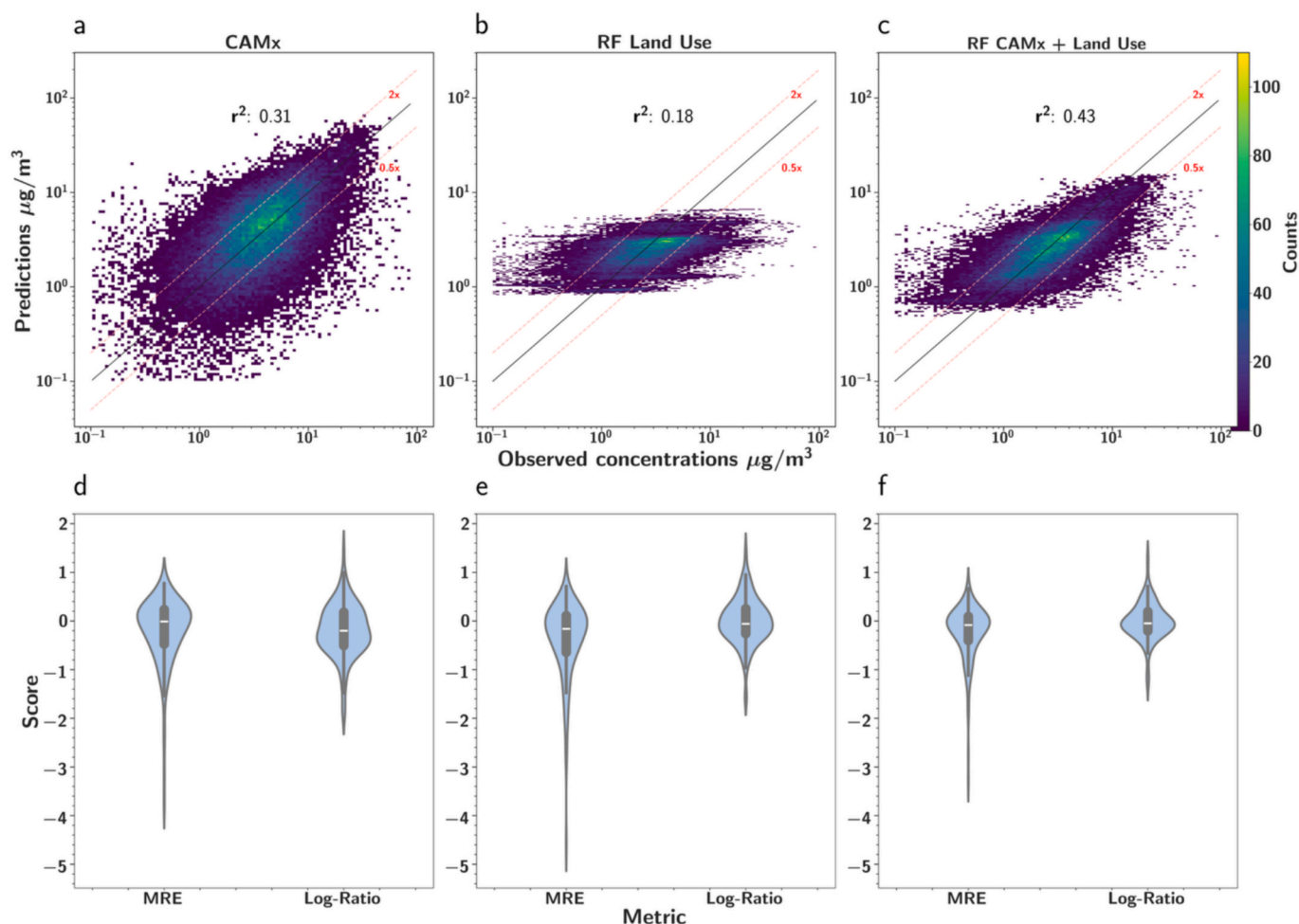


Fig. 2. The range of values of observed and predicted OA was binned to produce a histogram in 2D of the predicted values vs the observed values by the three models (a) CAMx, (b) RF_LU, (c) RF_CAMx_LU, with colors representing the number of total points in each bin. The corresponding station-stratified metrics of the Mean Relative Error (MRE) and log-ratio between observed and predicted values (Log-Ratio) are shown as boxplots in for (d) CAMx, (e) RF_LU and (f) for RF_CAMx_LU.

hyperparameters across iterations.

We trained two versions of the RF model to assess the contribution of land use and CAMx variables to OA predictions. The first model, RF_CAMx_LU, used CAMx outputs, land use, and meteorological data as predictors. The second, RF_LU, excluded CAMx outputs and relied only on land use and meteorological variables. We compared both RF models and the original CAMx predictions against observed OA concentrations. To improve prediction accuracy, we also tested log-transformed versions of the RF models (RF_CAMx_LU log-scale and RF_LU log-scale) to normalize the target variable and perform better for regression tasks whose residuals are homoscedastic in log-scale (Manning and Mullahy, 2001). All RF models were evaluated on the original scale using LOSO cross-validation. Model performance was assessed using station-stratified metrics: Mean Relative Error (MRE), which captures directional bias and is sensitive to concentration magnitude, and the Log-Ratio between observed and predicted values, which is less sensitive to outliers. To interpret the influence of individual predictors, we used permutation importance (Breiman, 2001; Pedregosa et al., 2011) and Shapley Additive Explanations (SHAP) values (Lundberg and Lee, 2017). Permutation importance quantifies the change in model error when feature values are randomly shuffled, while SHAP explains each feature's contribution to specific predictions.

2.4. Uncertainty and limitations

The uncertainties associated with OA modelling in our framework

can be broadly categorized into measurement-related and model-related uncertainties.

Measurement uncertainties. We categorized measurement uncertainties into three main sources: (1) offline OC measurements, (2) online OA measurements, and (3) the OA/OC conversion factor used to derive OA from OC.

Offline OC measurements. Uncertainties stem from both sampling and thermal-optical analysis. Sampling-related errors include flow calibration (~3%), positive artefacts from organic vapor adsorption (+10 to +20%) and negative artefacts from OA volatilization (-15 to -5%). Measurement related-errors include calibration accuracy (~3%), detection precision (~2%) and biases associated with the thermal-optical protocol used to separate organic and elemental carbon, with OC measurements using NIOSH and EUSAAR reported to differ by up to 10%. When propagated, these sources yield a total uncertainty of -10% to +21%, with an average positive bias of ~5% (El Haddad et al., 2011a,b).

Online OA measurements. OA data from ACSM/AMS instruments are affected by detection precision (<1% for 5 µg/m³ and daily averages), ionization efficiency calibration (~10%), collection efficiency correction (~25%), and the assumed relative ionization efficiency (set to 1.4, but ranges from 1.4 to 1.8). Propagating these errors results in an uncertainty range of -30% to +54%, with an average positive bias of ~12% (El Haddad et al., 2011a,b).

OC to OA conversion. Two factors influence the OC-to-OA conversion: (1) the transmission efficiency of the AMS/ACSM for PM_{2.5},

which is generally high (~80–90%) due to OA being predominantly in the accumulation mode, and (2) the OM/OC ratio, which varies spatially and temporally from 1.5 to 2.0 (variability ~14%), with higher values in rural areas. Considering an average OM/OC ratio of 1.75 and a transmission efficiency of 85%, we calculate an average OA/OC ratio of 1.5 used to convert OC measurements to OA. This ratio is consistent with that determined based on simultaneous online OA and offline OC measurements at few sites. We fitted a model with a conversion factor of 1.8 and compared with the factor 1.5, as detailed in the following section.

Implications. 1. Although most of our OA dataset is derived from converted OC measurements – which dominate the performance of the RF model, values are anchored to AMS/ACSM data through the constant OA/OC ratio of 1.5. If users want to anchor the measurements to offline OA, they should use a factor of 1.75 instead. 2. Using a fixed conversion factor introduces uncertainty to offline OA estimates but also enables a straightforward inference of OC concentrations from OA values. 3. Considering uncertainties from both OC measurements and the OA/OC ratio, we estimate an uncertainty of –19% to +26% for offline-derived OA, compared to –30% to +54% for online OA measurements.

Modelling uncertainties. Errors are primarily associated with the components of the chemical transport model, including errors in meteorological inputs, emission inventories, and the representation of atmospheric chemical and physical processes. CAMx biases are not strictly linear, as they result from multiple interacting sources, including meteorology, emissions, and chemical processes. This non-linearity can increase uncertainty, particularly in regions with sparse observational data where the machine learning model has limited guidance for bias correction. For machine learning when observations fall outside the range covered by the training set, predictions may be biased with unwarranted confidence. Techniques such as conformal prediction (Vovk et al., 2022) may help address these concerns and will be investigated in future work.

3. Results

3.1. Model performance and sensitivity

The observational dataset shows a mean observed OA concentration of 5.0 $\mu\text{g}/\text{m}^3$ across Europe, with a standard deviation of 5.1 $\mu\text{g}/\text{m}^3$, reflecting high regional variability (Fig. 1). Montale (Italy), Kraków (Poland), and Villanueva del Arzobispo (Spain) were the three monitoring locations with the highest mean OA concentrations of 33.6, 18.3, and 16.5 $\mu\text{g}/\text{m}^3$, respectively. We used this observational dataset as the target variable to train the RF model.

We evaluated model performance on a per-left-out-station basis. Table S3 reports the mean performance metrics across all sites, reflecting per-site prediction accuracy. In contrast, Fig. 2 presents the overall metrics calculated from all LOSO predictions combined for each of the models. Both RF based models performed better with log-transformed predictors, and we used those for further analysis and predictions. Fig. 2 compares the performance of CAMx, RF_CAMx_LU log-scale (from now on denoted only as RF_CAMx_LU), and RF_LU on a logarithmic scale through scatter density plots. CAMx achieved a coefficient of determination (R^2) of 0.31 (Pearson's correlation, $r = 0.55$), capturing the observed value range but exhibiting significant under- and over-predictions at lower concentrations (Fig. 2a). The LOSO predictions for RF_LU (Fig. 2b) had an $R^2 = 0.19$ ($r = 0.43$), with values concentrated mainly between 1 and 10, indicating that land-use modeling alone is insufficient to predict regional pollutants like OA. In contrast, RF_CAMx_LU (Fig. 2c) achieved the highest R^2 of 0.43 ($r = 0.65$), slightly over-predicting lower values but covering the full range of observed OA concentrations. The RMSE decreases from 4.26 $\mu\text{g}/\text{m}^3$ for CAMx to 3.63 $\mu\text{g}/\text{m}^3$ for RF_CAMx_LU, corresponding to a 15% improvement (Table S3). Analysis of MRE and log ratio shows that all models have slightly negative biases. CAMx (Fig. 2d) had a lower median MRE but a wider distribution. In contrast, RF_CAMx_LU (Fig. 2f) had a more

constrained MRE distribution. Among the three models, RF_CAMx_LU had the smallest Log-Ratio values, indicating higher accuracy across all observation magnitudes. Additional performance statistics are presented in a per-station stratification in Table S3. The sensitivity analysis of model performance to OA/OC ratios of 1.5 and 1.8 (Table S4) shows only small differences, indicating that the models are weakly sensitive to the choice of OA/OC ratio within this range.

While the land-use model (RF_LU) was capable of high-resolution predictions, it lacks information on pollution sources, chemical reactions, and transport dynamics. By integrating CAMx outputs with land-use variables in RF_CAMx_LU, we leveraged both types of information to produce downscaled predictions of OA levels in Europe. Combining the strengths of both approaches RF_CAMx_LU model showed the highest global R^2 and the lowest median Log-Ratio. Despite slightly higher median MRE than CAMx, it is compensated with less spread of the error bars and outliers for this metric (Fig. 2). We performed Wilcoxon paired tests on the LOSO results, using the null hypothesis that the mean performance metrics of the best-performing model do not differ from those of the other models. The tests confirmed that all previously discussed improvements in overall performance (e.g., R^2 , RMSE) are statistically significant. We also applied the Wilcoxon test to model bias (true – predicted), for which no statistically significant differences were found. This indicates that bias alone does not capture the full improvement in predictive skill and highlights the need to consider multiple performance metrics when evaluating model quality.

3.2. Drivers of organic aerosol concentrations

Ambient OA concentrations are shaped by their emissions, precursor sources, physical and chemical transformation, meteorological conditions, and land-use factors. These processes are further emphasized in geographical basins (Salma et al., 2020). The two methods, permutation importance and SHAP, identified biomass burning and other anthropogenic sources as key predictors, with urban fabric, natural green areas, and road networks emerging as the most influential land-use variables (Fig. S1 and S2), it's worth noting that while the results are suggestive, correlations among predictor variables may distort the true importance of these variables (Fig. S3) (Silva and Keller, 2024). We analyzed the relationship between land-use characteristics and high-resolution OA concentrations simulated by the RF_CAMx_LU model using binned boxplots (Fig. 3). OA levels generally decline with increasing natural green cover (Fig. 3a) but rise slightly with larger urbanization (Fig. 3b) and higher density of residential roads (road class 3). Although forests and other green areas contribute to biogenic SOA, our results for the European scale show, these regions tend to have lower OA concentrations than urban areas, with mean OA decreases by ~20% when natural green coverage increases by 50%. The decrease in OA with natural green is steeper in winter season than the summer season, where mean OA decreases by 27% in winter compared to 14% decrease in summer for a 50% increase in green coverage (Fig. S4). In contrast, grid cells with over 30% urban cover or more than 450 m of road class 3 within a 1 km buffer show elevated OA levels. While OA is typically viewed as a regional pollutant, these results highlight a modest but notable local-scale enhancement due to urbanization and road networks. To further evaluate the realism of the downscaled of fields following land-use parameters, we assessed model validation across three urbanization regimes categorized using IMD data. Fig. S6 shows predicted and observed OA comparison as a function of urbanization level and across the full temporal range. The RF_LU_CAMx model captures observed OA concentrations across urbanization levels and temporal scales. These results indicate that the RF_LU_CAMx model resolves sub-grid-scale variability and reproduces key spatial structures across the datasets. The influence of meteorological parameters on OA concentrations over Europe is shown in Fig. S5. OA concentrations increase with temperature, indicating the influence of secondary organic aerosol (SOA) formation (Fig. S5a). However, elevated OA concentrations are also observed at

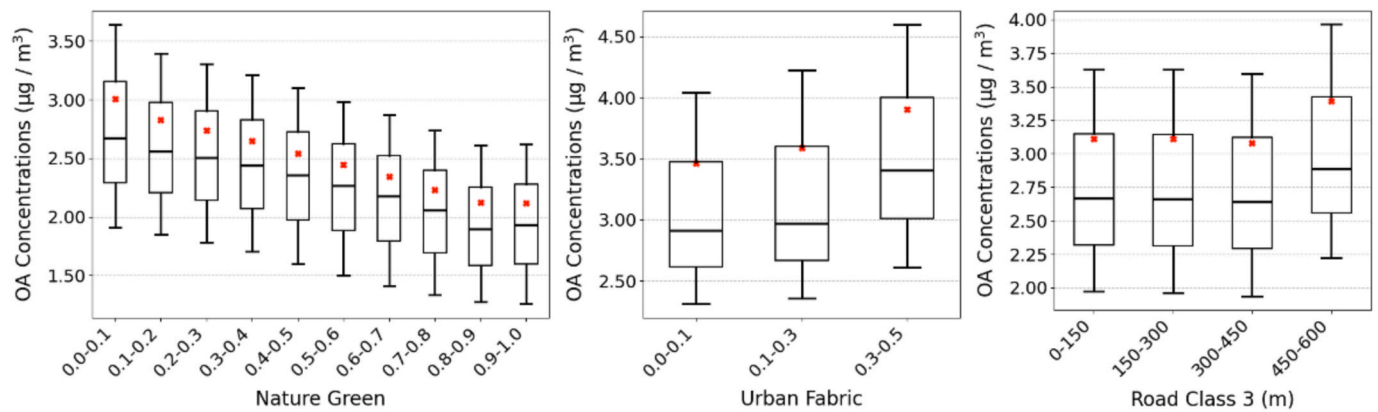


Fig. 3. Box-and-whisker plots showing the relationship between OA concentrations and binned land use parameters: (a) Natural Green, (b) Urban Fabric, and (c) Road Class. Boxes represent the interquartile range (IQR), central lines denote medians, whiskers extend to the 5th and 95th percentiles, and red squares indicate mean OA concentrations for each bin. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

lower temperatures, reflecting the impact of biomass burning during winter. Figs. S5(b, c) illustrate that high OA concentrations are associated with low relative humidity (RH) and low planetary boundary layer height (PBLH).

3.3. Spatial distribution of organic aerosol in Europe

We used the RF_CAMx_LU model, trained with the full dataset and optimal hyperparameters, to predict OA concentrations at a fine spatial resolution of $0.005^\circ \times 0.0025^\circ$ (~ 250 m) across Europe for the years 2011, 2013, 2015, 2017, and 2019. These high-resolution OA predictions over Europe consist of approximately 70 million grids for a single day, totaling 128 billion values for the entire dataset. Fig. 4 illustrates the mean predicted OA concentrations for the selected years across Europe. The results highlight several regions with high concentrations, including the Po Valley, Central Europe (Czechia, southern Poland, Slovakia, and Hungary), and Southeastern Europe (Croatia, Serbia, Bosnia, and Romania). In these areas, yearly mean OA concentrations exceed $10 \mu\text{g}/\text{m}^3$, which is more than double the revised annual $\text{PM}_{2.5}$ guideline level set by the WHO (WHO, 2021). Additionally, highly urbanized areas such as the Benelux region (Belgium, the Netherlands, and Luxembourg), Central Spain, Paris, Warsaw, among others, also showed high OA concentrations ($> 4 \mu\text{g}/\text{m}^3$).

Fig. 5 examines OA distribution across four regions in Europe: the Po

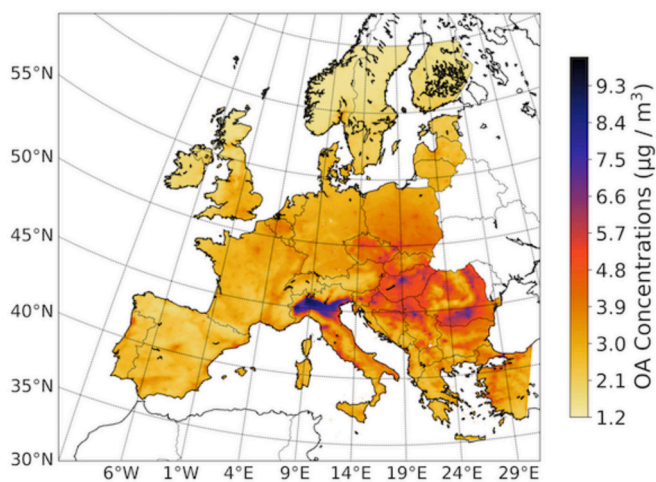


Fig. 4. Average OA concentration ($\mu\text{g}/\text{m}^3$) predicted by RF_CAMx_LU for years 2011, 2013, 2015, 2017, and 2019 at the spatial resolution of $0.0025^\circ \times 0.005^\circ$ (~ 250 m).

Valley, the Iberian Peninsula, Southeastern Europe (Serbia, Hungary, Slovakia, Romania, and Bulgaria), and the Benelux region as predicted by both CAMx and RF_CAMx_LU models. Fig. 5 further illustrates regional differences in OA predictions between RF_CAMx_LU and CAMx. While both models identify major high-concentration zones, CAMx produces broader, less distinct areas of elevated OA, whereas RF_CAMx_LU captures more localized distributions. The refined mapping provided by RF_CAMx_LU highlights the influence of land-use parameters in governing fine-scale spatial variations, with high OA concentrations closely aligning with urbanization and traffic density. For example, in the Benelux region, RF_CAMx_LU shows sharper distinctions between polluted and less polluted areas compared to CAMx, here downscaling enhances urban hotspots while reducing OA in surrounding areas, indicating that CAMx smooths sub-regional variability. Over Eastern Europe, CAMx produces relatively smooth and elevated background OA, whereas RF introduces stronger spatial heterogeneity and generally lower values, particularly in rural areas. Similarly, over Po-Valley RF is enhancing the sub-grid scale heterogeneity. Over the Iberian Peninsula, RF reducing CAMx background concentrations across much of the interior while preserving localized enhancements near urban and coastal regions. These results underscore the importance of high-resolution OA modeling for more accurately characterizing local air quality dynamics and identifying key drivers of OA variability. The difference between OA predicted by RF_CAMx_LU and CAMx over Europe (Fig. S7) indicates that the downscaled and bias-corrected OA generally reduces concentrations in regions where CAMx predicts higher values, while increasing concentrations in areas where CAMx predicts lower values. The zoomed difference maps in Fig. 5 further show that RF_CAMx_LU tends to apply positive corrections along road networks and in smaller urban areas, whereas it applies negative corrections over major cities. In these densely populated urban grids, CAMx's coarse resolution produces uniformly elevated OA concentrations, which are subsequently reduced by the RF_CAMx_LU downscaling.

3.4. Temporal distribution of OA concentrations

Fig. 6 presents monthly mean OA concentrations across Europe for the simulated years, using RF_CAMx_LU. The maps reveal a distinct seasonal cycle, with higher OA levels in winter and lower concentrations in summer, consistent with previous regional and local studies (Jiang et al., 2019). The seasonal differences are more clearly illustrated by the winter and summer OA means shown in Fig. S8, which highlights a greater regional contrast in concentrations during winter compared to summer. In high-concentration areas like the Po Valley, and Southeastern Europe, OA exceeds $10 \mu\text{g}/\text{m}^3$ in winter but typically remains below $5 \mu\text{g}/\text{m}^3$ in summer. This seasonal variation is largely driven by

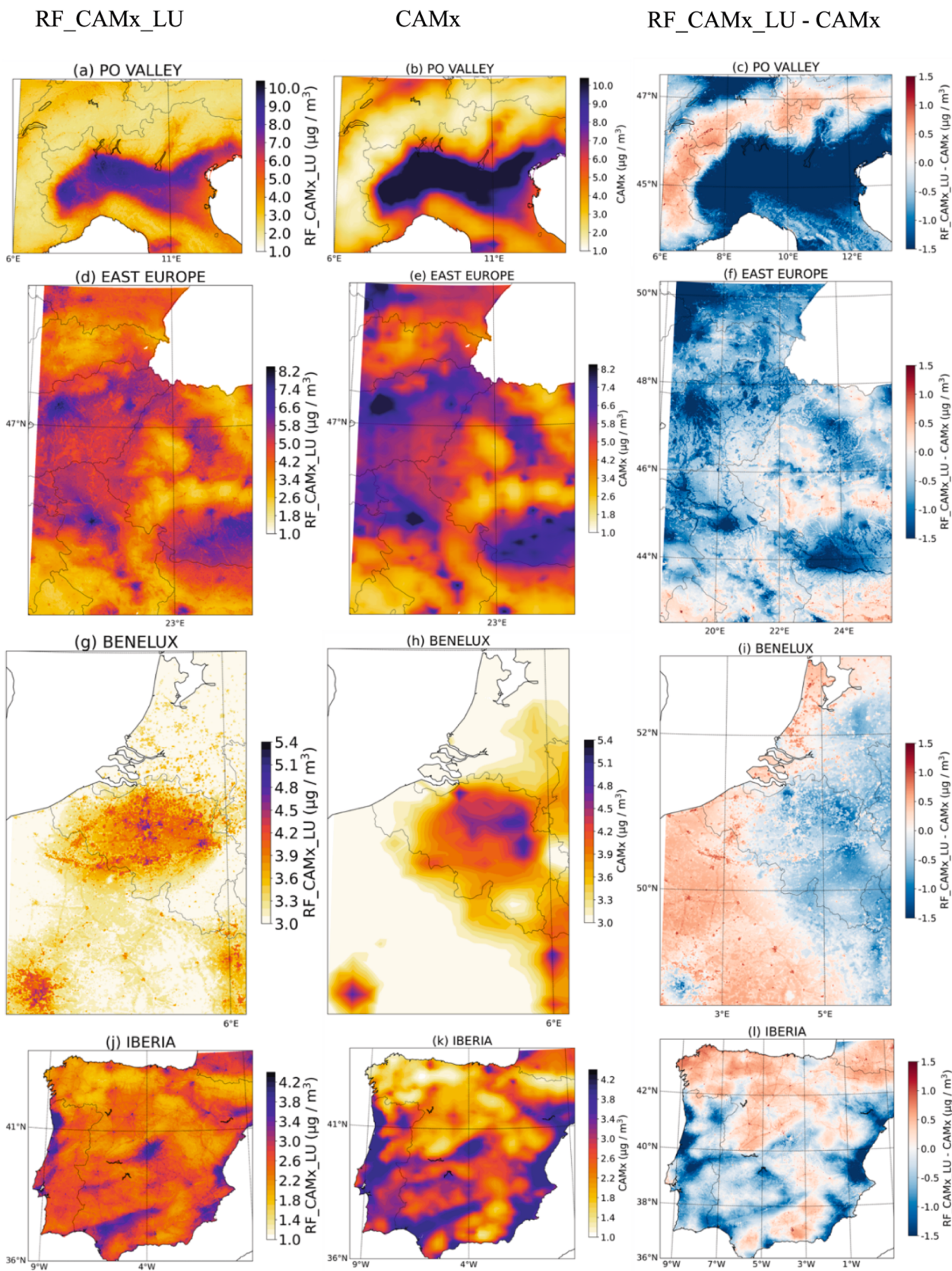


Fig. 5. Yearly mean OA concentrations predicted using RF_CAMx_LU (a, d, g, j), CAMx (b, e, h, k), and their difference (RF_CAMx_LU – CAMx) (c, f, i, l) across four regions: the Po Valley (a–c), Southeastern Europe (Slovakia, Hungary, Serbia, Romania, Bulgaria) (d–f), Benelux (Belgium, the Netherlands, Luxembourg) (g–i), and the Iberian Peninsula (j–l).

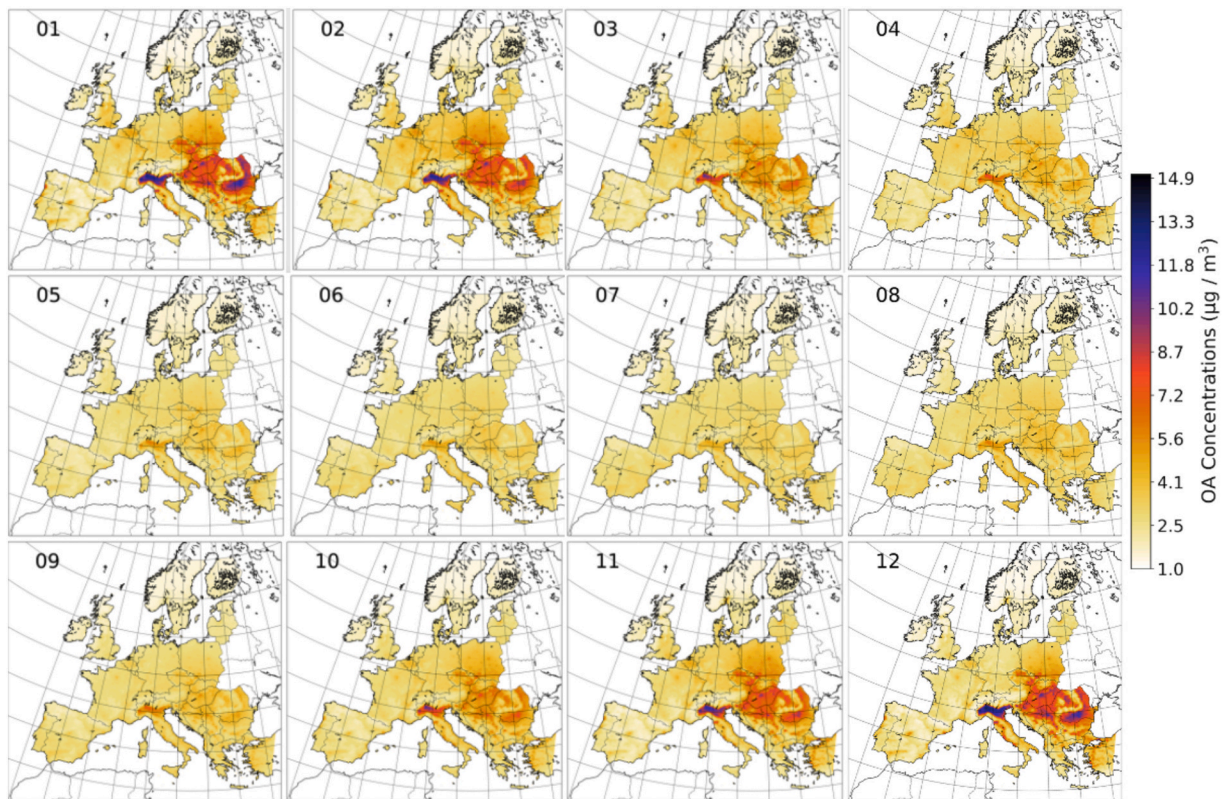


Fig. 6. Mean OA concentrations for each month (labels from 1 to 12 represent months from January to December) averaged over years 2011, 2013, 2015, 2017 and 2019 predicted at ~ 250 m resolution using the RF_CAMx_LU model.

increased residential biomass burning emissions, and low-dispersion conditions, such as low planetary boundary layer height in winter (Chen et al., 2022; Sandrini et al., 2014). While total OA follows this general trend, individual components may behave differently; For example, biogenic secondary organic aerosol production peaks in summer (Aksoyoglu et al., 2011; Jiang et al., 2019).

Fig. 7 presents temporal trends in OA concentrations across the Po Valley, Southeastern Europe, Benelux, and the Iberian Peninsula. Fig. 7 (a–d) In the Po Valley and Southeastern Europe, OA distributions show a bimodal pattern, indicating two distinct pollution zones. Mean OA concentrations declined from 2011 to 2019, likely reflecting mitigation efforts in residential heating, transport, and fuel combustion. Benelux exhibited a similar but narrower distribution, while the Iberian Peninsula showed a unimodal pattern with occasional peaks up to $12 \mu\text{g}/\text{m}^3$. Emission inventories report a 5–10% reduction in OC emissions across Eastern and Western Europe from 2012 to 2023, primarily due to decreased residential emissions (Soulie et al., 2024). In the UK, carbonaceous aerosol levels from residential wood combustion dropped by 2.5–5.5% annually between 2015 and 2021 at three monitoring sites, attributed to the replacement of old fireplaces with more efficient wood stoves (Font et al., 2022).

Monthly variations (Fig. 7e–h, middle row) reveal a winter-dominated bimodal OA distribution in the Po Valley and Southeastern Europe, with lower and more uniform concentrations in summer. Throughout the year, most of the Po Valley and Central Europe experienced OA levels between $2\text{--}6 \mu\text{g}/\text{m}^3$. However, in winter, concentrations showed a broader range, with significant areas reaching $6\text{--}14 \mu\text{g}/\text{m}^3$ likely driven by increased local emissions from residential heating and atmospheric stagnation. In contrast, summer concentrations were more evenly distributed, likely due to enhanced atmospheric mixing and regional SOA formation. The winter shift toward higher concentrations in source regions and lower concentrations elsewhere underscores the importance of high-resolution maps in capturing this

spatial variability. Benelux and the Iberian Peninsula exhibited similar seasonal patterns, with greater spatial heterogeneity in winter. Notably, Benelux showed wider concentration variability in summer compared to other regions, likely because primary emissions in this region are less linked to residential heating and more to transport-related activities, including ports and airports. Fig. S9 shows monthly OA variation and distribution over whole Europe, here monthly mean markers indicate show winter maxima, early-summer minima, and a stronger secondary peak in late summer. Spread is greatest in winter and late summer, reflecting higher spatial variability during elevated OA periods.

The weekly OA cycle, depicted by a distribution for each day of the week, (Fig. 7i to 7l, bottom row) shows higher concentrations on weekdays, peaking on Wednesdays and Thursdays, with a decline over the weekend. However, this variation is less pronounced than seasonal changes, suggesting that while traffic and industrial activity contribute to OA levels, dominant sources such as biomass burning, and secondary aerosol formation do not follow a strong weekly trend.

The temporal patterns of OA (annual, seasonal, weekly) in the RF_CAMx_LU model largely reflect the underlying CAMx drivers: year-specific emissions control annual trends, meteorology drives seasonal variability, and weekly emission profiles generate weekly cycles. Land-use and population variables vary only at annual resolution and have limited influence on temporal trends. While the temporal dynamics remain similar, the RF model provides bias correction and high-resolution spatial downscaling, which refine the concentration distributions shown in Fig. 7. For comparison, Fig. S10 shows the corresponding distributions from CAMx alone, showing qualitatively similar trends but with different distributions.

3.5. Population exposure to OA

We estimated the European population's exposure to OA using the high-spatial resolution annual mean OA concentrations and gridded

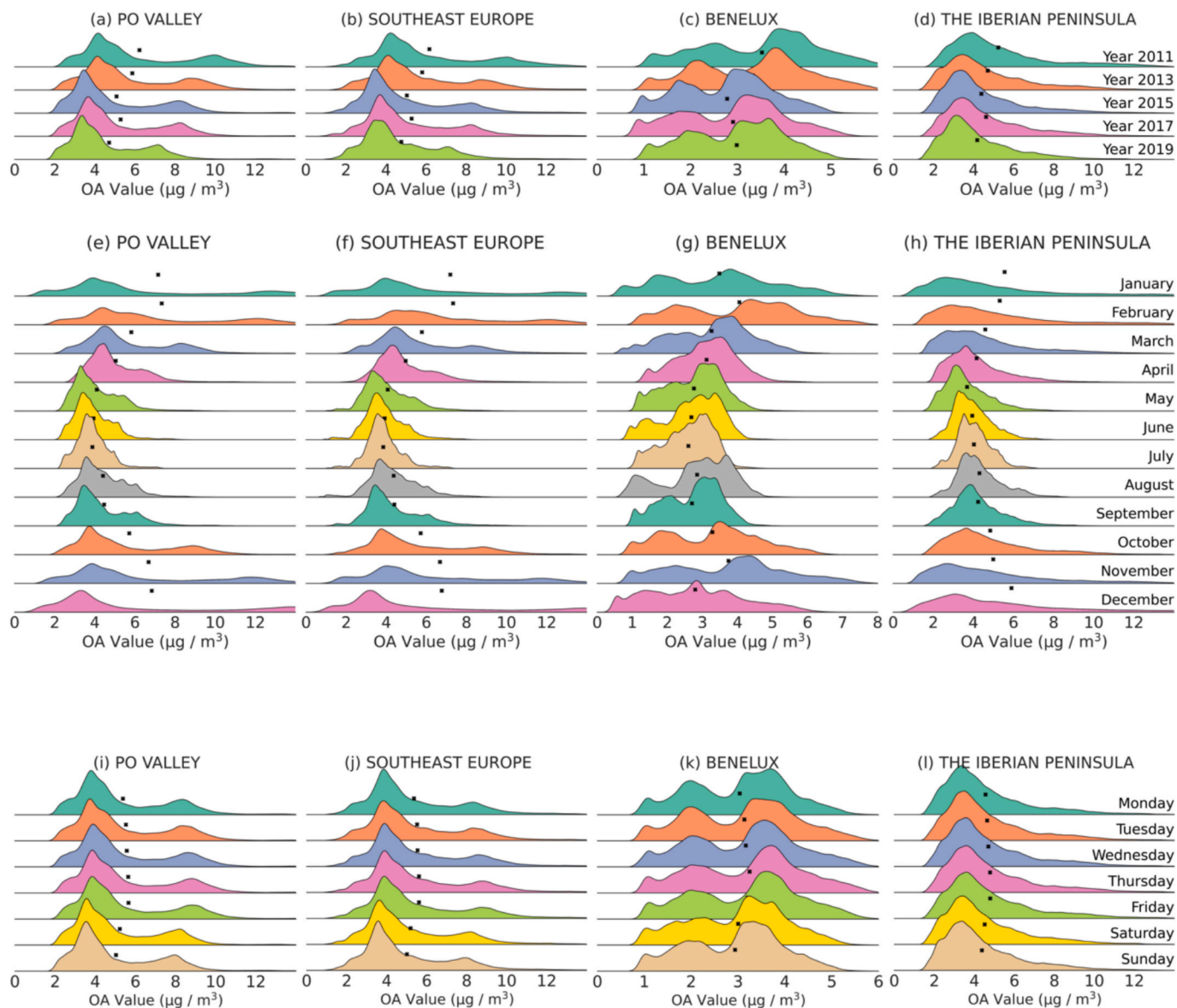


Fig. 7. Distribution of organic aerosol (OA) concentrations using all simulated years for the four regions of interest: (a–d) annual variation in the Po Valley, Southeastern Europe, Benelux, and the Iberian Peninsula, respectively; (e–h) corresponding monthly variations; and (i–l) corresponding weekly (day-of-week) variations for each region. The black dot represents the mean OA concentration.

population data. OA levels were grouped into relevant concentration bins and analyzed alongside population density and total population per bin (Fig. 8). The results show that while ~35% (~190 million) of the European population experienced annual mean OA levels below $3 \mu\text{g}/\text{m}^3$, 47.65 million people were exposed to concentrations exceeding $5 \mu\text{g}/\text{m}^3$, which is higher than the revised WHO guideline level for $\text{PM}_{2.5}$. Areas with higher population density had elevated OA levels, amplifying health risks. Notably, areas with OA concentrations above $8 \mu\text{g}/\text{m}^3$ have a median population density exceeding 500 inhabitants/ km^2 . For reference, the mean population density for the European Union is ~109 inhabitants/ km^2 (ranging from ~18 to 1700 inhabitants/ km^2 among EU countries). Population exposure derived from the downscaled OA fields is more skewed than that based on CAMx OA (Fig. S11). The high-resolution RF_CAMx_LU fields shift exposure toward lower OA bins and introduce greater variability in population density within the high-OA categories, reflecting the fine-scale urban hotspots that CAMx cannot resolve. Comparing Fig. 8 with Fig. S11 shows that downscaling sharpens spatial contrasts and reduces averaging effects, leading to more realistic population-weighted exposure estimates.

4. Conclusions

The integration of CTM with machine learning, using a uniquely extensive OA observation database with wide spatial and temporal coverage, has significantly enhanced prediction accuracy and resolution of daily OA concentration. This approach reveals key OA hotspots and long-term trends, demonstrating the importance of targeted mitigation measures, such as addressing biomass burning and vehicular emissions. OA concentrations in regions such as the Po Valley and Southeastern Europe often exceed twice the WHO annual $\text{PM}_{2.5}$ guideline, underscoring the critical need for higher resolution, source-specific OA maps to meet WHO guidelines. Expanding the observation network to Eastern Europe, improving emission inventories, and refining chemical parameterizations will further enhance predictions. This methodology, applicable to other pollutants such as secondary inorganic aerosols, offers a promising pathway for better understanding regional aerosol pollutants. Generated OA maps are suited for epidemiological studies aiming at understanding the health impacts of OA.

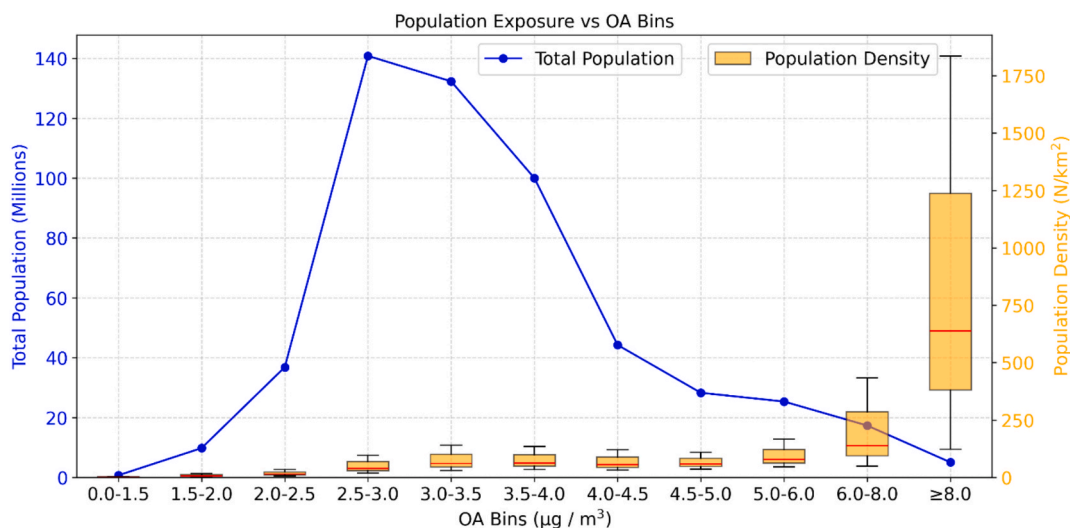


Fig. 8. Total population exposed (blue line) and population density (box-whiskers) for bins of OA concentrations ($\mu\text{g}/\text{m}^3$) over Europe. Orange boxes represent the interquartile range, whiskers represent the 5th–95th percentile, and horizontal red lines denote the median of population density. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

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Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.envint.2026.110143>.

Data availability

High-resolution annual mean organic aerosol (OA) concentrations over Europe, predicted using integrated CAMx and random forest modelling, are available on Zenodo at the following <https://doi.org/10.5281/zenodo.18516697>, for the years 2011, 2013, 2015, 2017, and 2019. The observed OA concentration used to compile the OA database cannot be shared directly; however, it may be obtained upon request from the respective data owners or the corresponding authors.

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