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Fine particles at a background site in Central France: Chemical compositions, seasonal variations and pollution events

Lin HE1,2, Hui CHEN1, Jérôme RANGOGNIO3, Abderrazak YAHYAOU13, Patrice COLIN3, Jinhe WANG4, Véronique DAËLE1, Abdelwahid MELLOUKI1,2,*

1 Institut de Combustion, Aérothermique, Réactivité et Environnement, CNRS, Orléans, France
2 School of Environmental Science and Engineering, Shandong University, Jinan, People’s Republic of China
3 Lig’Air, Réseau de Surveillance de la Qualité de l’Air en Région Centre-Val de Loire, Saint-Cyr-en-Val, France
4 School of Municipal and Environmental Engineering, Shandong Jianzhu University, Jinan 250101, and Shanghai Key Laboratory of Atmospheric Particle Pollution and Prevention (LAP3), Department of Environmental Science & Engineering, Fudan University, Shanghai 200433, People’s Republic of China

Abstract: To expand our knowledge of regional fine particles in Central France (Centre-Val de Loire region), a field observation study of PM$_{2.5}$ was carried out at Verneuil site (46.81467N, 2.61012E, 180 m.a.s.l.) from 2011 to 2014. The mass concentrations of water-soluble inorganic ions (WSIIs), organic carbon (OC), elemental carbon (EC) and biomass burning tracer (Levoglucosan) in PM$_{2.5}$ were measured. Annual average PM$_{2.5}$ mass concentrations were 11.8, 9.5, 12.6 and 10.2 μg m$^{-3}$ in 2011, 2012, 2013 and 2014, respectively, three of four higher than the WHO guideline of 10 μg m$^{-3}$. Secondary inorganic aerosol (SIA) and organic matter (OM) appeared to be the major components in PM$_{2.5}$ in Verneuil, contributing 30.1 - 41.8% and 36.9 - 46.3%, respectively. Main chemical species were observed in the following order: winter ≥ spring> autumn > summer. Backward atmospheric trajectories were performed using Hysplit model and suggested that the PM$_{2.5}$ pollutants caused by atmospheric transport were mainly originated from European inland, mainly east to north-east areas. During the observation period, five pollution events were reported and indicated that not only the polluted air masses from central Europe but also the biomass burning from East Europe significantly influenced the air quality in Verneuil site.

Keyword: PM$_{2.5}$, Chemical characteristics, Backward trajectory, Pollution events, Biomass burning, Central France
*Corresponding author. E-mail: mellouki@cnrs-orleans.fr
1. Introduction

Airborne particles have attracted intense scientific and public attention in recent years for their adverse effect on human health, visibility and climate (Koçak et al. 2007, Pateraki et al. 2014). In particular, fine particles (PM$_{2.5}$, particulate matter with aerodynamic diameter (AD) $< 2.5$ µm) can remain in suspension much longer than coarse particles and can penetrate more deeply into the respiratory tract causing many kinds of reactions (oxidative stress, local pulmonary, systemic inflammatory responses, etc.) (Amodio et al. 2010, Maté et al. 2010). Furthermore, they have a clear impact on climate, causing direct (absorbing or reflecting the radiation, modifying the scattering), semi-direct (heating may cause the burn-off of clouds) and indirect (favoring or preventing the formation of clouds, their albedo and life-times) effects on the global radiative budget (Barker et al. 2007). Additionally, other effects are evident on visibility, building materials and ecosystems (Razak et al. 2015).

PM$_{2.5}$ is a mixture of many species, loading complex information of air pollution. It consists mainly of water-soluble inorganic ions (WSIIs), carbonaceous species, crustal elements and trace elements. WSIIs, dominated by secondary ionic aerosols (SIA, including NO$_3^-$, SO$_4^{2-}$ and NH$_4^+$), are directly related to their precursor gases SO$_2$, NO$_x$ and NH$_3$, with rather well known sources mostly related to fossil fuel burning (SO$_2$, NO$_x$) and agriculture (NH$_3$), and can directly affect the acidity of precipitation as well as atmospheric visibility (Pathak et al. 2011, Zhang et al. 2015). Organic carbon (OC), which comprises thousands of organic compounds, originates from both natural and
anthropogenic sources and contains many toxic substances, such as polycyclic aromatic hydrocarbons (PAHs) and normal alkanes (Li et al. 2013). Element carbon (EC), existing in an inert state, is directly emitted from primary combustion and plays an important role on radiative forcing and climate system (Srinivas and Sarin 2014). The crustal elements (Mg, Al, K, Ca and Fe), mainly from soil or construction dust, are also important factors leading to degradation of visibility (Tian et al. 2016). The annual, seasonal and daily tendencies of these chemical compositions have been widely used to identify the sources responsible for the PM pollution in the atmosphere.

Europe, as the most developed continent, has experienced serious air pollution in the history and keeps launching thematic strategy on air pollution since then. The most recent wave of policy was launched in 2005 including control on PM. In order to understand the impact of human activities on the atmospheric environment and assess air pollution over a regional area, it is important to monitor air quality in the regional/background area in addition to that in the urban/polluted area.

In Europe, studies on fine particles at background sites in recent years have been reported for a number of areas such as Veneto (Italy), Diabla Góra (Poland), Montseny (Spain), Melpitz (Germany) (Bressi et al. 2012, Khan et al. 2016, Pey et al. 2009, Rogula-Kozłowska et al. 2014). These sites were far away from cities and industrial areas and PM$_{2.5}$ mass concentrations at these background sites showed similar seasonal variation trends, lower in the summer and higher in the winter and spring. Based on the chemical composition and trajectory analysis, natural sources were considered as the main contributor to fine particles in the summer, while mid- or long-range transport
anthropogenic sources such as fossil fuel combustion and biomass burning contributed more in the winter and spring. Additionally, oceanic wind played a clean-up role for those coastal sites near the Atlantic.

In France, studies were mainly found at urban and suburban measurement sites around large cities (Cazier et al. 2016, Dimitriou and Kassomenos, El Haddad et al. 2011, Fortems-Cheiney et al. 2016, Kfoury et al. 2016). Particle pollution in these areas were regarded as a complex consequence of industrial, domestic and traffic (shipping for dock cities) emissions. As we were aware of, there is only study in Rubrouck, North France, where PM$_{2.5}$ were mainly from natural emissions in the summer and PM$_{2.5}$ concentration peaks recorded in spring with high SIA contributions was highly related to the accumulation phenomena and long-range transport from the Eastern and Central Europe origins (Cazier et al. 2016).

In a recent study on the origin of PM pollution in Paris, Beekmann et al. (2015) have reported that primary fossil fuel combustion emissions constituted less than 20% in winter and 40% in summer of carbonaceous fine PM, suggesting that the major part of secondary organic aerosol (SOA) is of modern origin, i.e., from biogenic precursors and from wood burning.

In our earlier study in Orléans site, by monitoring particle size distribution and aerosol extinction coefficients at a suburban measurement site in central France, it was pointed out that Atlantic Ocean air masses could clean up regional pollution in the winter. Meanwhile, this region can be influenced by air masses originating from other European areas, Atlantic Ocean, Mediterranean Sea air masses and African dust plume.
but their corresponding impacts in different seasons are not yet clear (Hu et al. 2014).

To expand our knowledge of regional fine particles in Central France, a field observation study of PM$_{2.5}$ was carried out at Verneuil site from 2011 to 2014. Herein, temporal concentration variations of PM$_{2.5}$ and their chemical compositions were reported; interrelations between different compositions were studied; primary and secondary organic sources of PM$_{2.5}$ were evaluated; backward trajectories were analyzed to identify the influence of mid- or long-range atmospheric transport; and several pollution events were reported.

2. Material and methods

2.1 Measurement site

Verneuil (46.81467N, 2.61012E, 180 m.a.s.l.) is a rural background site located in the center of France, in the southeast of the Centre-Val de Loire region and 230 km south to Paris. As illustrated in Fig. s1, the surrounding areas of Verneuil site are mainly grassland and forest, with two two-lane roads and a few residence houses nearby. The samplers and the inlets of instruments were placed on the roof, 3.5 m above the ground. Verneuil is part of MERA observatory within the framework of EMEP project (http://ce.mines-douai.fr/pages/observatoire-mera or http://www.emep.int/).

2.2 Fine aerosol particles measurement

PM$_{2.5}$ were measured by Lig’Air, the Centre-Val de Loire Region Air Quality Observatory, continuously during 2011 to 2014 by TEOM-FDMS (Thermal, TEOM
1400 with an FDMS 8500 module, TEOM: Tapered Element Oscillating Microbalance, FDMS: Filter Dynamics Measurement System). It was operated at a standard flow-rate of 1 L·min⁻¹ with a 2.5 μm cutting head. The mass loss due to volatilization is corrected by FMDS module. PM₁₀ were measured simultaneously during 2010 to 2011 using the same system but with a 10 μm cutting head and at a standard flow-rate of 1 L·min⁻¹. The filter was replaced when its loading percentage was greater than 30% to keep the flow rate. Further details of TEOM-FDMS can be found elsewhere (Patashnick and Rupprecht 1991). The PM concentrations used in this paper are hourly averages from 5-min measurements by TEOM-FDMS.

2.3 Chemical analyses

2.3.1 Aerosol sampling

Fine aerosol particles (PM₂.₅) were collected by Lig’Air for 24 hours (9:00 am to 9:00 am UT) every 7 days using a DIGITEL DA-80 high volume sampler. All the samples were collected on 47 mm quartz filters (Whatman Inc., Maidstone, UK). All quartz filters were pre-baked at 450 °C for 4 h and without binding to have very low levels of blank. After collection, samples were sent to laboratory (Laboratoire des Sciences du Climat et de l’Environnement (LSCE) et Laboratoire de Glaciologie et Géophysique de l’Environnement (LGGE)) for chemical analyses within 4 hours. A total of 224 filters have been collected and few samples have been discarded because of power failures, chemical analysis problems, etc.

2.3.2 Water Soluble Ions
The following water-soluble major ions were analyzed by Ion Chromatographs (IC) (model Dionex ICS 3000): chloride, nitrate, sulfate, sodium, ammonium, potassium, magnesium and calcium. The analytical protocol conformed to the Lig’Air technical report TR-16269 CEN / TC 264. Briefly, filter samples were extracted in 15 mL of ultrapure water during 45 min in a sonic bath. To prevent bacteria activity, 50 µL of chloroform was added in the extractions. Samples were then filtered using Acrodisc filters (Pall Gelman) with a porosity of 0.22 µm. CS16 column was used for cation analyses while AS11 HC column for anion. Calibration was performed daily from certified standard solutions. The detection limits were less than 0.05 µg m⁻³ for each ion.

2.3.3 Carbonaceous contents

The carbonaceous content of samples was analyzed for EC and OC using a Thermo-Optical Transmission (TOT) method on a Sunset Lab analyzer (Birch 1998). Recently developed EUSAAR2 temperature program was used here (Cavalli et al. 2010). Briefly, it includes temperature from 200 to 650 °C for the analysis of OC in 100% He, and from 500 to 850 °C for the analysis of EC in 98% He + 2% O₂. A detailed description concerning thermal optical methods and the relevance of the protocol chosen can be found elsewhere (Cavalli et al. 2010, Chow et al. 1993). Calibrations were performed daily. The detection limit and the uncertainty are estimated to be 0.2 µg and 5%, respectively.

2.3.4 Monosaccharides anhydrides (MA) and polyols

For the analysis of MA (Levoglucosan, mannosan and galactosan) and polyols
(arabitol, sorbitol and mannitol), filter punches (4.5 cm²) were extracted with 2 mL ultrapure water by 30 min vortex agitation. Filter extracts were analyzed by HPLC (high-performance liquid chromatography) with a PAD (pulsed amperometric detection) detector (model Dionex DX500 + ED400). Column used were Metrosep Carb 1 – 6 Guard, A Supp 15 – 150 and Carb 1 – 150. Eluent was 6.0 mmol L⁻¹ NaOH and flow rate was 1 mL min⁻¹. External calibrations were performed using standard solutions.

2.4 Chemical mass closure

Aerosol Chemical Mass Closure (CMC) consists in comparing the sum of the major aerosol chemical species (PM_{2.5-chem}) with measurements (PM_{2.5-meas}). When achieved (i.e. when PM_{2.5-chem} = PM_{2.5-meas}), CMC attests the consistency of chemical analyses, and confirms that all the major aerosol chemical species are quantified. PM_{2.5-chem} calculation is here expressed as:

\[
[PM_{2.5-chem}] = [\text{Sea Salt}] + [\text{Dust}] + [\text{Secondary Inorganic Aerosols}] + [\text{Carbonaceous Matter}]
\] (1)

[Sea salt] (ss-) concentrations are calculated from the six major ions accounting for more than 99% of the mass of salts dissolved in seawater:

\[
[\text{Sea salt}] = [\text{Na}^{+}] + [\text{Cl}^{-}] + [\text{Mg}^{2+}] + [\text{ss-K}^{+}] + [\text{ss-Ca}^{2+}] + [\text{ss-SO}_4^{2-}]
\] (2)

With \([\text{ss-K}^{+}] = 0.036 \cdot [\text{Na}^{+}]; [\text{ss-Ca}^{2+}] = 0.038 \cdot [\text{Na}^{+}]\) and \([\text{ss-SO}_4^{2-}] = 0.252 \cdot [\text{Na}^{+}]\)

Typical seawater ion ratios based on the average seawater composition are taken from Seinfeld and Pandis (Seinfeld et al. 2006).

[Dust]: The mineral dust fraction in PM_{2.5} can be quantified by different methods,
which are based on its average elemental composition from specific sites, or on specific tracers (Guieu et al. 2002, Sisler and Huffman 1994). More recently, nss-calcium (“nss-” standing for non-sea-salt) has been widely used because of its abundance (Jean-P. et al. 2003, Sciare et al. 2005). Here in this paper, 15% contribution of nss-calcium in mineral dust determined by Guinot et al. was used (Guinot et al. 2007).

$$[\text{Dust}] = \frac{[\text{nss-}\text{Ca}^{2+}]}{0.15}$$ (3)

The resulting proportion of dust in PM$_{2.5}$ during the whole campaign ranges from 1.1 to 5.2% on average from 2011 to 2014 (Fig. 4), which was in agreement with that in Paris (2% to 5%) (Bressi et al. 2012).

**[Secondary inorganic aerosols]** (SIA) are calculated as:

$$[\text{SIA}] = [\text{nss-}\text{SO}_4^{2-}] + [\text{NO}_3^-] + [\text{NH}_4^+]$$ (4)

where $[\text{nss-}\text{SO}_4^{2-}] = [\text{SO}_4^{2-}] - [\text{ss-}\text{SO}_4^{2-}]$.

**[Carbonaceous matter]** can be expressed as:

$$[\text{Carbonaceous Matter}] = [\text{EC}] + [\text{OM}]$$ (5)

[OM] is inferred from OC as:

$$[\text{OM}] = f_{\text{OC-OM}}\cdot[\text{OC}]$$ (6)

$f_{\text{OC-OM}}$ is the OC to OM conversion factor.

OC was measured by the Sunset Lab analyzer conveniently, while OM include C, O, N, H etc., which is difficult to be quantified. The varied chemical composition, highly dependent on locations, seasons and time of the day, makes the estimation of OM complex. Turpin and Lim (2001) then recommend the measurement of the average molecular weight per carbon weight in the location of interest, corresponding to a $f_{\text{OC-}}$. 
OM of 1.6 ± 0.2 and 2.1 ± 0.2 for urban and nonurban aerosols, respectively. To estimate the \( f_{\text{OC-OM}} \) from our own dataset, a method adapted from Guinot et al. (2007) who used the chemical mass closure technique as a tool to assess \( f_{\text{OC-OM}} \) from OC measurements was applied.

In this method, CMC was assumed to be achieved (\( \text{PM}_{2.5-\text{chem}} = \text{PM}_{2.5-\text{meas}} \)) and following equation could be got from equation (1):

\[
f_{\text{OC-OM}} = \frac{1}{[\text{OC}]} \cdot (\text{PM}_{2.5-\text{meas}} - ([\text{Sea Salt}] + [\text{Dust}] + [\text{SIA}] + [\text{EC}])))
\]  
(7)

to simplify equation (7), [Remaining Mass] was defined as:

\[
[\text{Remaining Mass}] = \text{PM}_{2.5-\text{meas}} - ([\text{Sea Salt}] + [\text{Dust}] + [\text{SIA}] + [\text{EC}])
\]  
(8)

and lead to:

\[
[\text{Remaining Mass}] = f_{\text{OC-OM}}[\text{OC}]
\]  
(9)

thus, \( f_{\text{OC-OM}} \) could be calculated as the slope of the linear regression from the plot of [Remaining Mass] against [OC]. Excluded days with \( f_{\text{OC-OM}} \) higher than 4 and lower than 1 which were considered as physically meaningless, \( f_{\text{OC-OM}} \) was calculated as 2.066. Finally, in order to make an easy comparison with whole chemical component data from Paris, \( f_{\text{OC-OM}} \) was chose as 2.05 (Bressi et al. 2012).

2.5 Meteorological parameters and backward trajectories

Meteorological parameters such as temperature, pressure, precipitation, wind speed and wind direction were obtained from http://www.wunderground.com/, which shares reliable and meaningful weather information with the public since 1993 by the weather company in San Francisco. Air mass backward trajectories were calculated using the
Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Stein et al. 2015) and fire spots data were obtained from NASA FIRMS (http://earthdata.nasa.gov/data/near-real-time-data/firms).

3. Result and discussion

3.1 PM$_{2.5}$ levels

The profiles of PM$_{2.5}$ and PM$_{10}$ mass concentrations in daily average from 2011 to 2014 at Verneuil site are given in Fig. 1. The annual average PM$_{2.5}$ mass concentrations from 2011 to 2014 were 11.8, 9.5, 12.6 and 10.2 μg·m$^{-3}$, respectively. These annual averages, three of four, were higher than World Health Organization guideline of 10 μg·m$^{-3}$ (WHO 2006). Compared with other rural background sites in Europe (Table 1), Verneuil site appeared to be less polluted by PM$_{2.5}$ pollutions. The differences in PM$_{2.5}$ level between those sites can probably be ascribed to the local energy structures, site locations and meteorological conditions. France and Switzerland possess larger clean energy shares such as nuclear power and hydroelectricity, leading to the relative low pollution emissions compared with the other countries of more fossil fuel consumption (OECD. 2014).

PM$_{2.5}$ mass concentrations showed similar seasonal pattern from 2011 to 2014 with high levels in spring (Mar, Apr and May) and winter (Dec, Jan and Feb) and low levels in summer (Jun, Jul and Aug) and autumn (Sep, Oct and Nov), except in the winter of 2014 (low PM$_{2.5}$ level might be partly explained by the absence of high
pollution events in this season). The daily average mass concentrations of PM$_{2.5}$ ranged from 1.3 to 78.8 μg·m$^{-3}$. High-pollution events, identified by daily average of PM$_{2.5}$ larger than 50 μg·m$^{-3}$, have occurred every year during February or March, as listed in Table 3, and would be further discussed in detail in section 3.4. Box plots of hourly PM$_{2.5}$ mass concentration distributions for the whole measurement period was given in Fig. s2. No evident variation of PM$_{2.5}$ was found during rush hours which indicated that Verneuil was barely influenced by vehicle emission on the roadside.

As illustrated in Fig. 1, PM$_{10}$ mass concentrations during 2011 and 2012 showed the same variation trend as PM$_{2.5}$. The PM$_{2.5}$/PM$_{10}$ ratios varied in a range of 0.28-1 with an average of 0.74. This result was also consistent with those from other European regions (annual PM$_{2.5}$/PM$_{10}$ ratio of 0.79 in Germany, 0.86 in Spain and 0.72 in Switzerland) (Pey et al. 2009). Sample number distribution of PM$_{2.5}$/PM$_{10}$ ratios against PM$_{2.5}$ mass concentrations is shown in Fig. s3. During the clean days (PM$_{2.5}$ < 10 μg·m$^{-3}$), samples mainly distributed in the areas of PM$_{2.5}$/PM$_{10}$ ratio below 0.7, whereas during pollution days (PM$_{2.5}$ > 20 μg·m$^{-3}$), the PM$_{2.5}$/PM$_{10}$ ratios were high (0.8 – 1.0). Thus, it was considered that PM$_{2.5}$ rather than PM$_{10}$ were mainly responsible for PM pollution in Verneuil during 2011 and 2012.
Fig. 1. Daily averages of PM$_{2.5}$ and PM$_{10}$ mass concentrations measured by TEOM-FDMS and ratio of PM$_{2.5}$ to PM$_{10}$ (PM$_{2.5}$ / PM$_{10}$) from 2011 to 2014. Daily averages were calculated from 0:00 am to 0:00 am.
3.2 Chemical composition

3.2.1 General description

Filter samples were collected for chemical composition analysis for 24 hours (from 9:00 am to 9:00 am UT) every 7 days. Fig. 2 shows the mass concentrations of PM$_{2.5}$ and its corresponding main species. Different from “PM$_{2.5}$” in Fig. 1, “PM$_{2.5}$ Measured” were PM$_{2.5}$ mass concentrations specially averaged from 9:00 am to 9:00 am in next day by TEOM-FDMS to get consistence with filter sample data of the other species.
Fig. 2 Daily averages of PM$_{2.5}$ mass concentrations measured by TEOM-FDMS and the mass concentrations of main chemical compositions from 2011 to 2014 in Verneuil. Daily averages of PM$_{2.5}$ measured were calculated from 9:00 am to 9:00 am. Note that different units have been used for the figures.
PM$_{2.5}$ Reconstructed was calculated from chemical species and shown in Fig. 3. The good consistency between PM$_{2.5}$ Measured and PM$_{2.5}$ Reconstructed confirmed the accuracy of our field measurements and verified the conversion factors (f$\text{OC-OM}$) used to estimate organic matter in this paper. The mass shares of the identified components in PM$_{2.5}$ were lowest in summer and highest in winter, varied in the range of 54.4% - 60.5% and 74.2% - 99.6%, respectively.

The difference between the PM$_{2.5}$ Measured and PM$_{2.5}$ Reconstructed was defined as unidentified matter (UM) including some trace elements, carbonaceous, and/or organic anions which were not covered by the analysis, or evaporated components such as water and ammonium nitrate during sampling, storage and transportation of samples (Rogula-Kozłowska et al. 2014).

The UM contributed more in the summer (18.3% - 32.9%, Table s1) and less in the winter (0% - 22.0%). This can be probably explained by that (a) in hot and dry
period, nitrates and some organic compounds might get lost by evaporation during storage and transportation of samples; (b) while in winter or spring, identified mass of PM can be overestimated due to the adsorption of SO₂ onto ammonia-rich particles or adsorption of nitric acid (HNO₃) onto mineral and salt particles (Pathak et al. 2009).

Annual and seasonal averages of PM₂.₅ mass concentrations and corresponding chemical composition shares are shown in Fig. 4 (mass concentrations are given in Table s1). Among identified components, SIA and OM appeared to be the major components in PM₂.₅ in Verneuil, contributing 30.1 - 41.8% and 36.9 - 46.3%, respectively. SIA shares varied in the reported ranges of other European rural background sites, while OM shares were close to rural background sites near Paris (40% to 47%) and much higher than other European sites (21.0% to 32.6%) (Table 1) (Bressi et al. 2012, Khan et al. 2016, Pey et al. 2009, Pirovano et al. 2015, Putaud et al. 2010). OM was reported to be originated from biomass burning, SOA formation, mid- or long-range transport, and traffic emission (Bressi et al. 2012, Khan et al. 2016). High OM shares in Verneuil was largely attributed to biomass burning and/or SOA formation in winter/spring, while solely to SOA formation favored by biogenic volatile organic compounds (VOCs) emission and solar irradiation in summer/autumn (see discussion in section 3.2.3). SIA were originated from transport from Europe inland and gas-to-particle conversion by precursors (SO₂ NOₓ, NH₃) in excess of ammonia and under favorable thermodynamic conditions (Bressi et al. 2012, Clegg et al. 1998, Hamaoui 2012) (see discussion in section. 3.2.2).
Fig. 4 Annual and seasonal average PM$_{2.5}$ mass concentration measured by TEOM-FDMS and corresponding chemical composition shares from 2011 to 2014. Daily averages of PM$_{2.5}$ Measured were calculated from 9:00 am to 9:00 am.
Table 1 Average mass concentration of PM$_{2.5}$ and its main compositions at different rural site in Europe (OM: organic matter, SIA: secondary inorganic aerosol)

<table>
<thead>
<tr>
<th>location (country)</th>
<th>Type</th>
<th>Measurement period</th>
<th>PM$_{2.5}$ (μg m$^{-3}$)</th>
<th>OM %</th>
<th>SIA %</th>
<th>Data source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Verneuil</td>
<td>Rural</td>
<td>2011-2014</td>
<td>9.5-12.6</td>
<td>36.9-46.3</td>
<td>30.1-41.8</td>
<td>This study</td>
</tr>
<tr>
<td>South Paris (France)</td>
<td>Rural</td>
<td>09/2009-09/2010</td>
<td>10.8</td>
<td>40.0</td>
<td>48.0</td>
<td>M. Bressi et al. 2012</td>
</tr>
<tr>
<td>Northeast Paris (France)</td>
<td>Rural</td>
<td>09/2009-09/2010</td>
<td>12.6</td>
<td>47.0</td>
<td>42.0</td>
<td>M. Bressi et al. 2012</td>
</tr>
<tr>
<td>Chaumont (Switzerland)</td>
<td>Rural</td>
<td>1998</td>
<td>8.1</td>
<td>21.0</td>
<td>43.1</td>
<td>J.-P. Putaud et al. 2010</td>
</tr>
<tr>
<td>Diabla Góra (Poland)</td>
<td>Rural</td>
<td>2010</td>
<td>15.0</td>
<td>21.2</td>
<td>36.3</td>
<td>W. Rogula-Kozlowska et al. 2014</td>
</tr>
<tr>
<td>Streithofen (Austria)</td>
<td>Rural</td>
<td>1999-2000</td>
<td>17.5</td>
<td>25.7</td>
<td>35.4</td>
<td>J.-P. Putaud et al. 2010</td>
</tr>
<tr>
<td>Bosco Fontana (Italy)</td>
<td>Rural</td>
<td>2004-2007</td>
<td>34.7</td>
<td>-</td>
<td>-</td>
<td>G. Pirovano et al. 2015</td>
</tr>
<tr>
<td>Montseny (Spain)</td>
<td>Rural</td>
<td>2003</td>
<td>13.6</td>
<td>27.0</td>
<td>36.0</td>
<td>J. Pey et al. 2009</td>
</tr>
<tr>
<td>Königsplatz (German)</td>
<td>Kerbside</td>
<td>11/2007-03/2008</td>
<td>43.0</td>
<td>-</td>
<td>-</td>
<td>R.M. Qadir et al. 2014</td>
</tr>
<tr>
<td>Katowice (Poland)</td>
<td>Urban</td>
<td>2010</td>
<td>43.0</td>
<td>25.8</td>
<td>21.7</td>
<td>W. Rogula-Kozlowska et al. 2014</td>
</tr>
<tr>
<td>Zuerich (Switzerland)</td>
<td>Urban</td>
<td>1998</td>
<td>19.0</td>
<td>23.2</td>
<td>41.0</td>
<td>J.-P. Putaud et al. 2010</td>
</tr>
<tr>
<td>Milano (Italy)</td>
<td>Urban</td>
<td>2004-2007</td>
<td>31.5</td>
<td>-</td>
<td>-</td>
<td>G. Pirovano et al. 2015</td>
</tr>
<tr>
<td>Venice (Italy)</td>
<td>Residence</td>
<td>2012-2013</td>
<td>25.3</td>
<td>32.6</td>
<td>-</td>
<td>Md. Badiuzzaman Khan et al. 2016</td>
</tr>
<tr>
<td>Barcelona (Spain)</td>
<td>Urban</td>
<td>2003-2006</td>
<td>29.0</td>
<td>31.0</td>
<td>29.7</td>
<td>N. Perez et al. 2008</td>
</tr>
</tbody>
</table>
The validity of the water soluble inorganic ions could be ascertained through the balance of anions and cations (Chow et al. 1994). The anion and cation microequivalents for the PM$_{2.5}$ samples were calculated as follows:

\[
A(\text{anion microequivalents}) = \frac{\text{Cl}^-}{35.5} + \frac{\text{SO}_4^{2-}}{45} + \frac{\text{NO}_3^-}{62}
\]

\[
C(\text{cation microequivalents}) = \frac{\text{Na}^+}{23} + \frac{\text{NH}_4^+}{18} + \frac{\text{K}^+}{39} + \frac{\text{Mg}^{2+}}{12} + \frac{\text{Ca}^{2+}}{20}
\]

As illustrated in Fig. 5, the average ratio of the molar equivalents (eq m$^{-3}$) of cations (Na$^+$, NH$_4^+$, K$^+$, Mg$^{2+}$ and Ca$^{2+}$) to anions (Cl$^-$, NO$_3^-$ and SO$_4^{2-}$) was close to unity, i.e. 1.01 ± 0.01 in Verneuil, suggesting that almost all the ions had been quantified in this study, although bicarbonate and carbonate were not measured in this work. The strong correlation ($R^2 = 0.97$) between the anion and cation equivalents indicated that the relationship between the measured cations and anions remained constant during neutralization. Most of the samples collected were positioned close to the 1:1 cation:anion (C:A) line, implying that they were generally neutral.
Fig. 5 The correlations between the anion and cation equivalents in the four seasons in Verneuil from 2011 to 2014. 1:1 dash line indicates the complete neutralization. Linear regressions for each season were calculated and equations were given.
3.2.2 Secondary inorganic aerosol

SIA (NO$_3^-$, nss-SO$_4^{2-}$ and NH$_4^+$) shares showed clear seasonal tendencies in the order of spring > winter >> autumn > summer (Fig. 4). High SIA contributions found in the spring/winter can be highly related to the higher emissions of SIA precursors (NO$_x$ and SO$_2$), accumulation phenomena and mid- or long-range transport. Seasonal average mass concentration of NO$_3^-$, nss-SO$_4^{2-}$ and NH$_4^+$ are given in Table s1. Average of NO$_3^-$ mass concentration was 3.24 μg·m$^{-3}$ in spring, which was 36 times higher than that in summer (0.09 μg·m$^{-3}$). Whereas average of nss-SO$_4^{2-}$ mass concentration was 1.68 μg·m$^{-3}$ in spring and 1.34 μg·m$^{-3}$ in summer. Although NO$_3^-$ and nss-SO$_4^{2-}$ showed same seasonal variation trends with highest concentrations in spring/winter, the difference between the spring/winter and summer nss-SO$_4^{2-}$ concentrations was much smaller compared to that for NO$_3^-$.

The low temperature favored the conversion of gaseous HNO$_3$ to aerosol particles as NH$_4$NO$_3$, leading to higher contributions of SIA in spring and winter (Tao et al. 2012); and possibly, the formation of SO$_4^{2-}$ was enhanced by the increased photochemical reactivity in the summer (Choi et al. 2012).

The ratio of [NH$_4^+$] to [SO$_4^{2-}$] is a common factor used to estimate the neutralization of all the three SIA species, which was defined as

$$\frac{[\text{NH}_4^+]}{[\text{SO}_4^{2-}]} = \frac{[\text{NH}_4^+/18]}{[\text{SO}_4^{2-}/96]}$$

Usually, area with $[\text{NH}_4^+]/[\text{SO}_4^{2-}] > 1.5$ was regarded as ammonium-rich, and acidic sulfate is considered to be fully neutralized at $[\text{NH}_4^+]/[\text{SO}_4^{2-}] \geq 2$ (Huang et al. 2011).

Annual averages of $[\text{NH}_4^+]/[\text{SO}_4^{2-}]$ were calculated from Table s1, which were 4.36,
3.12, 3.94 and 2.49 in 2011 to 2014, respectively. All of them were significantly higher than 2. Meanwhile, \( \text{NO}_3^- + \text{SO}_4^{2-} \) and \( \text{NH}_4^+ \) microequivalents were calculated as follows,

\[
\text{NO}_3^- + \text{SO}_4^{2-} \text{ microequivalents} = \text{SO}_4^{2-}/48 + \text{NO}_3^-/62
\]

\[
\text{NH}_4^+ \text{ microequivalents} = \text{NH}_4^+/18
\]

and their correlations in different seasons were plotted in Fig. s4. The slopes close to the unity also indicated that most of \( \text{NO}_3^- \) and \( \text{SO}_4^{2-} \) were neutralized with \( \text{NH}_4^+ \) to form \( (\text{NH}_4)_2\text{SO}_4 \) and \( \text{NH}_4\text{NO}_3 \). To summarize, atmosphere in this area was regarded as ammonium saturated.

Bressi et al. (2012) reported ammonium-saturated SIA in Paris region with excess ammonia. In Verneuil, nitric acid can be easily converted to nitrate thanks to the great availability of ammonia probably originated from agricultural activity (Hamaoui 2012, Pey et al. 2009).

3.2.3 Carbonaceous aerosols

Carbonaceous aerosols, include organic carbon (OC, primarily or secondarily produced by gas-to-particle conversion from volatile organic compounds) and elemental carbon (EC, primarily from incomplete combustion). The annual averages of OC and EC mass concentrations in PM\(_{2.5}\) ranged from 1.83 to 2.67 \( \mu \text{g} \cdot \text{m}^{-3} \) and 0.20 to 0.28 \( \mu \text{g} \cdot \text{m}^{-3} \) in Verneuil site, respectively (Table s1). The low EC concentration indicated that there were no significant combustion emission sources nearby (Samara et al. 2013), in consistence with the absence of PM\(_{2.5}\) rush-hour peaks. The seasonally averaged OC and EC mass concentrations ranked in the descending order of winter >
autumn ≈ spring > summer as the PM$_{2.5}$ mass concentrations. The relatively high levels of OC and EC in winter implying the aerosols were highly influenced by fossil fuel and/or wood burning for energy production and domestic heating.

3.2.3.1 OC/EC ratios

Because carbonaceous aerosol represented a variety of emission sources, the ratio of OC to EC can be used to distinguish source characteristics. As shown in Fig. s5, the medians of OC/EC ratios at Verneuil were 11.2, 8.4, 8.3 and 9.3 from 2011 to 2014, respectively, which was comparable with those measured at forest sites in Europe (11.5 in western Germany) (Bressi et al. 2012, Kourtchev et al. 2008), about 1.5 times of those measured at rural background sites close to Paris (5.8), but much higher than those measured at European urban sites (2.6–3.1 in Barcelona, 2.8–4.7 in Amsterdam, and 3.5–4.4 in Ghent) (Kim et al. 2000, Na et al. 2004). Previous studies have shown that the OC/EC ratio from biomass burning is very high (with a lower limit of 6.0 in Central and Southern Europe) (Gelencsér et al. 2007), whereas fossil-fuel emissions tend to have relatively low OC/EC ratios in range of 2.5–5.0 (Chen et al. 2006, Schauer et al. 2002). The OC/EC ratios with a median of 9.3 (10% as 5.7 and 90% as 15.0) indicated that biomass burning and/or SOA formation rather than primary traffic emission contributed strongly to the organic aerosols in Verneuil. In the absence of active biomass burning in summer/autumn, OC/EC always exhibited high values except for 2012, indicating strong SOA contribution in Verneuil, in consistent with other Europeans rural sites (Bressi et al. 2012, Khan et al. 2016, Pey et al. 2009). Taking the
meteorological conditions into account, high average temperature and efficient solar irradiation (Table s2) resulted in strong biogenic VOC emission and active photochemical reaction (Liu et al. 2009, Peñuelas and Llusià 2003), which can partly explain significantly high OC/EC ratios in 2011 compared with 2012-2014.

3.2.3.2 Primary organic carbon (POC) and secondary organic carbon (SOC)

OC could be divided into two parts: primary organic carbon (POC) and secondary organic carbon (SOC). POC is emitted directly in the particle-phase and SOC is formed from gas-to-particle conversion in the air (Choi et al. 2012). Although the presence of SOC in aerosols can be identified by the OC/EC ratio when the ratio is higher than 2.0, it is still difficult to directly quantify the POC and SOC by chemical analysis technology (Zhang et al. 2012). The minimum OC/EC ratio method has been widely used to estimate the POC and SOC (Zhang et al. 2015), which was described as follows,

\[
POC = EC \times (OC/EC)_{\text{min}} \\
SOC = OC - POC
\]

The minimum OC/EC ratio in this study was 3.08, defined by the mean of the lowest 5% OC/EC values. Seasonal averages of POC and SOC mass concentration and SOC/OC ratios are given in Table 2. Annual averages of POC and SOC mass concentration were 0.69 and 1.98 μg·m\(^{-3}\), 0.67 and 1.42 μg·m\(^{-3}\), 0.86 and 1.67 μg·m\(^{-3}\), 0.61 and 1.22 μg·m\(^{-3}\) in 2011 to 2014, respectively. Except for the winter in 2014, POC and SOC always exhibited highest in winter and lowest in summer. The latter can be explained by blowing-away effect from Atlantic Ocean air masses in summer (Hu et al.}
The seeming high SOC masses in winter, however, could be overestimated using minimum-OC/EC method under biomass burning influence (Ding et al. 2012). On the other hand, increasing humidity in Verneuil in winter provided favorable meteorological conditions for SOC formation (Zhang et al. 2015). Moreover, a lower boundary layer height (BLH) in winter also favored SOC precursor stagnation and SOC formation (Duan et al. 2005).

The medians of SOC mass fractions (SOC/OC) was presented in Table 2. The significant high SOC/OC in 2011 was in consistent with OC/EC as discussed in section 3.2.3.1. The seeming seasonal tendency of SOC/OC in 2012 and 2013—high in winter/spring and low in summer/autumn, however, might be caused by the overestimation using minimum-OC/EC method under biomass burning influence of in winter/spring (Ding et al. 2012).
Table 2 Seasonal POC and SOC concentrations and SOC/OC ratios (in median) in 2011 to 2014

<table>
<thead>
<tr>
<th></th>
<th>2011</th>
<th>2012</th>
<th>2013</th>
<th>2014</th>
</tr>
</thead>
<tbody>
<tr>
<td>Winter</td>
<td>1.19</td>
<td>3.50</td>
<td>0.92</td>
<td>2.58</td>
</tr>
<tr>
<td>Spring</td>
<td>0.60</td>
<td>1.64</td>
<td>0.41</td>
<td>1.27</td>
</tr>
<tr>
<td>Summer</td>
<td>0.38</td>
<td>1.28</td>
<td>0.37</td>
<td>0.67</td>
</tr>
<tr>
<td>Autumn</td>
<td>0.80</td>
<td>2.24</td>
<td>0.80</td>
<td>1.14</td>
</tr>
<tr>
<td>Annual</td>
<td>0.69</td>
<td>1.98</td>
<td>0.67</td>
<td>1.42</td>
</tr>
</tbody>
</table>
3.2.4 Levoglucosan and K⁺

Levoglucosan and K⁺ have been considered as tracer for biomass burning (BB). Recent reports pointed out that using levoglucosan as BB tracer was less interfered since there can be other sources for K⁺ in addition to BB (Caseiro et al. 2009). However, due to the absence of levoglucosan measurement in 2011 and 2012, consistency between K⁺ and levoglucosan was analyzed. Mass concentrations of K⁺ and levoglucosan are given in Fig. 2 (e), which were strongly correlated with a $R^2 = 0.79$ (Fig. s6). This correlation coefficient was in consistency with previous study ($R^2 = 0.81$) for typical biomass burning (Urban et al. 2012). However, it should be noticed that Levoglucosan occasionally appeared low (<100 ng m⁻³) even if K⁺ exceeded 50 ng m⁻³ (background concentration), implying K⁺ might originate from other sources such as agricultural activities rather than biomass burning. On the other hand, Levoglucosan always appeared high (>300 ng m⁻³) when K⁺ exceeded 100 ng m⁻³, reliable for identifying biomass burning events. The surroundings in Verneuil was mainly vegetations and didn’t change in 2011-2014; possible K⁺ sources such as soil resuspension, fertilizers, seawater and waste incinerator were limited. Took them into account and compared with studies from other five European countries (Jedynska et al. 2014, Reche et al. 2012), K⁺ could be regarded as a reliable biomass burning tracer at Verneuil when higher than 100 ng m⁻³ for 2011 and 2012 in the absence of Levoglucosan measurement. Taking event 1 as an example, the high K⁺ concentration of 556 ng·m⁻³ on Jan 31, 2011 indicated that this pollution event was significantly
influenced by biomass burning, in consistence with a high OC/EC ratio of 15.4. Further discussion on biomass burning tracers in section 3.2.4 implied that biomass burning had remarkable impact on PM$_{2.5}$ in Verneuil.

3.3 Cluster analysis

Backward trajectory cluster analysis was utilized to identify the PM$_{2.5}$ sources that influenced Verneuil site. 48-hour back trajectories at 100 m a.g.l. (meter above ground level) altitude with time resolution of 3 hours were clustered into 4 main trajectories for seasons from 2011 to 2014 as showed in Fig. 6.

As illustrated in Fig. 6, air masses arriving at Verneuil came from north to northeast, west and south directions. Fastest air flows came from west were always with lower PM$_{2.5}$ concentrations which indicated that the Atlantic flows brought clean air to the center France. Southerly air flows came from mountainous areas and the west Mediterranean Sea, which had PM$_{2.5}$ concentrations higher than those of westerly flows but lower than those of northerly flows. Northeasterly air flows crossed central European regions such as Belgium, Netherland and west Germany, which accounted for 20% to 30% in different seasons. And average PM$_{2.5}$ concentrations of air masses came from these regions were 23.8, 23.1, 10.4 and 12.0 µg m$^{-3}$ for winter, spring, summer and autumn, which were the highest ones among those of westerly and southerly flows in the same season. Thus, it was suggested that the pollutants caused by atmospheric transport were mainly came from regions and countries located in northeast of Verneuil.
Fig. 6 Seasonal 48-hours backward trajectory clusters at 100 m a.g.l. (above ground level) altitude with corresponding statistic PM$_{2.5}$ mass concentrations (small interior figures) from 2011 to 2014. A solid triangle denotes the average. The horizontal line across the box is the median, the lower and upper hinges represent the 25th and 75th percentiles, respectively, while the lower and upper cross represent the 1st and 99th percentiles, respectively. The bottom and top line represent the minimum and maximum, respectively.
<table>
<thead>
<tr>
<th>Event</th>
<th>Period</th>
<th>Highest PM$_{2.5}$ mass concentration (μg m$^{-3}$)</th>
<th>SIA (μg m$^{-3}$)$^b$</th>
<th>OC/EC$^b$</th>
<th>Levoglucosan/K$^+$ (ng m$^{-3}$)$^{a,b}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>29/01/2011 - 02/02/2011</td>
<td>55.7 (31/01/2011)</td>
<td>33.0</td>
<td>15.4</td>
<td>NA/556.2</td>
</tr>
<tr>
<td>2</td>
<td>01/03/2011 - 07/03/2011</td>
<td>78.8 (05/03/2011)</td>
<td>25.6</td>
<td>10.2</td>
<td>NA/151.6</td>
</tr>
<tr>
<td>3</td>
<td>01/02/2012 – 13/02/2012</td>
<td>53.3 (13/02/2012)</td>
<td>20.3</td>
<td>12.5</td>
<td>NA/257.8</td>
</tr>
<tr>
<td>4</td>
<td>25/03/2013 – 07/04/2013</td>
<td>63.8 (31/03/2013)</td>
<td>30.6</td>
<td>13.3</td>
<td>583.7/165.4</td>
</tr>
<tr>
<td>5</td>
<td>11/03/2014 – 15/03/2014</td>
<td>62.0 (15/03/2014)</td>
<td>25.4</td>
<td>8.5</td>
<td>234.0/181.2</td>
</tr>
</tbody>
</table>

$^a$ NA means data Not Available

$^b$ Values were the averages during corresponding event
3.4 Identification of pollution events

As listed in Table 3, five heavy pollution events occurred during the four years. The period of event was defined from the first day when PM$_{2.5}$ mass concentration became higher than 25 μg·m$^{-3}$ in continuous 3 days, and until the first day when it became lower than 25 μg·m$^{-3}$ in continuous 3 days.

Fig. s7 illustrated the 5-days backward trajectories as well as fire spots distribution during the pollution periods and clean days. Evidently, these events had strong relationship with air mass transport. Before and after the pollution periods, air masses mainly originated from the Atlantic with low PM$_{2.5}$ levels. As illustrated in Fig. s9, sea salt contributed more during these clean days. While during the heavy pollution periods, air masses mainly came from central European countries.

It should be noted that backward trajectories of Event 5 traced to area with intensive wild fire spots in East Europe regions. This event was recorded by the sample collected on March 13$^{th}$, 2014. 5-days backward trajectory clusters, fire spots and mass concentrations for PM$_{2.5}$ and its main components are illustrated in Fig. 7. The high levoglucosan mass concentration (424.5 ng·m$^{-3}$) and OC/EC ratio (8.5) could be indicators of the impact of biomass burning (Cao et al. 2009). And high SIA contribution was related to long-range atmospheric transport. To summarize, event 5 was mainly caused by long-rang transport biomass burning plumes originated from East Europe.
Fig. 7 5-days backward trajectory clusters at 100 m a.g.l. altitude with corresponding wild fire spots distribution and mass concentrations of PM$_{2.5}$, levoglucosan and main components during the period of Event 5. Single trajectories were calculated per hour. Units of main components were in $\mu$g m$^{-3}$. 
Mass concentrations of main composition in PM$_{2.5}$ for pollution days during events 1 to 4 and clean days after or before these events are given in Fig. s9. High concentrations of biomass burning tracers (K$^+$/levoglucosan) were found during the periods of events 1 to 4, but fire spots were not evidently observed here. These events were influenced by biomass burning but slightly from fire spots we got. Compared to other events in the spring, event 1 and 3 happened in January and early February with the highest K$^+$ concentrations (556 and 215 ng·m$^{-3}$) and OM contributions (26.8 and 15.0 %), which means more biomass burnings in winter. And according to other researches in France, they were mainly from domestic heating (Kfoury et al. 2016). SIA, which increased from 11.19, 7.02, 3.84 and 0.79 μg·m$^{-3}$ during clean days to 32.98, 25.57, 24.57 and 33.88 μg·m$^{-3}$ during pollution days, accounted for the most part of PM$_{2.5}$ in these events, especially for Event 2 and 4 which appeared in spring. As a rural measurement site, without local emission sources, pollutants were mainly imported from regions and countries located east of Verneuil. Meantime, these countries in central Europe used more fossil fuels for the energy production and more woods for residence heating during cold days, which would contribute a lot to PM$_{2.5}$ and lead to high levels of K$^+$ and levoglucosan, as well as high levels of SIA precursors, NO$_x$ and SO$_2$ (Rogula-Kozłowska et al. 2014). And with mid- or long-range atmospheric transport and complex reactions, these SIA precursors led to a high level of SIA.

Meteorological data in Fig. s8 also showed that these events occurred in the coldest and temperature dropping days when there would be more fossil fuel and wood burnings. Thus, it was suggested that Events 1 to 4 were mainly impacted by aerosols
from fossil fuel combustion and biomass burning in central European countries during cold days, and these aerosols were brought to Verneuil site by the mid- or long-range transport. Additionally, BLH was usually low in winter, which may lead to pollutant accumulations (Salmond 2005).

4. Conclusions

In this study, PM$_{2.5}$ and its main chemical species were investigated at a rural background site in central France (Val-de-Loire region) from 2011 to 2014. PM$_{2.5}$ contributed predominately to particle pollution in Verneuil site as indicated by high PM$_{2.5}$/PM$_{10}$ ratios. Annual average mass concentrations of PM$_{2.5}$ were 1.8, 9.5, 12.6 and 10.2 μg·m$^{-3}$ in 2011 to 2014, respectively. PM$_{2.5}$ concentrations showed similar seasonal pattern from 2011 to 2014 with high levels in spring and winter and low levels in summer and autumn. OM and SIA, which accounted for 30.1% ~ 41.8% and 36.9% ~ 46.3%, respectively, were major components in PM$_{2.5}$. OM contributed more in the winter while SIA (NO$_3^-$, nss-SO$_4^{2-}$ and NH$_4^+$) shares showed clear seasonal tendency in the order of spring > winter > autumn > summer. High OM shares in Verneuil was largely attributed to biomass burning and/or SOA formation in winter/spring, while solely to SOA formation in summer/autumn. SIA, characterized to be ammonium-saturated, originated from transport from Europe inland and gas-to-particle conversion by precursors (SO$_2$, NO$_x$, NH$_3$) in excess of ammonia under favorable thermodynamic conditions. OC/EC ratios indicated that the influence of biomass burning and SOC contributed strongly to the organic aerosols in Verneuil site. Meteorological conditions
appeared Mass concentrations of levoglucosan and K\(^+\) also proved the impact of biomass burning on the particle pollution in Verneuil. Backward trajectory cluster analysis implied the air masses with high PM\(_{2.5}\) levels had strong relationship with atmospheric transport from central Europe. Finally, five heavy pollution events were identified and results showed not only the polluted air masses from Central Europe but also the biomass burning from East Europe will influence the air quality in Verneuil.

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