

Assessment of vertically-resolved PM₁₀ from mobile lidar observations

J.-C. Raut^{1,*} and P. Chazette²

¹Laboratoire de Météorologie Dynamique, Ecole Polytechnique, 91128 Palaiseau, France

²Laboratoire des Sciences du Climat et de l'Environnement, Laboratoire mixte CEA-CNRS-UVSQ, CEA Saclay, 91191 Gif-sur-Yvette, France

*now at: Laboratoire Atmosphères, Milieux et Observations Spatiales, Laboratoire mixte CNRS-UVSQ-UPMC, Université Paris 6, 75252 Paris, France

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Abstract. We investigate in this study the vertical PM₁₀ distributions from mobile measurements carried out from locations along the Paris Peripherique (highly trafficked beltway around Paris), examine distinctions in terms of aerosol concentrations between the outlying regions of Paris and the inner city and eventually discuss the influence of aerosol sources, meteorology, and dynamics on the retrieved PM₁₀ distributions. To achieve these purposes, we combine in situ surface measurements with active remote sensing observations obtained from a great number of research programs in Paris area since 1999. Two approaches, devoted to the conversion of vertical profiles of lidar-derived extinction coefficients into PM₁₀, have been set up. A very good agreement is found between the theoretical and empirical methods with a discrepancy of 3%. Hence, specific extinction cross-sections at 355 nm are provided with a reasonable relative uncertainty lower than 12% for urban (4.5 m² g⁻¹) and periurban (5.9 m² g⁻¹) aerosols, lower than 26% for rural (7.1 m² g⁻¹) aerosols, biomass burning (2.6 m² g⁻¹) and dust (1.1 m² g⁻¹) aerosols. The high spatial and temporal resolutions of the mobile lidar (respectively 1.5 m and 1 min) enable to follow the spatiotemporal variability of various layers trapping aerosols in the troposphere. Appropriate specific extinction cross-sections are applied in each layer detected in the vertical heterogeneities from the lidar profiles. The standard deviation (rms) between lidar-derived PM₁₀ at 200 m above ground and surface network stations measurements

was ~14 μg m⁻³. This difference is particularly ascribed to a decorrelation of mass concentrations in the first meters of the boundary layer, as highlighted through multiangular lidar observations. Lidar signals can be used to follow mass concentrations with an uncertainty lower than 25% above urban areas and provide useful information on PM₁₀ peak forecasting that affect air quality.

1 Introduction

Aerosol studies have experienced a revival of interest since human activities tend to increase their concentrations in the atmosphere. The anthropogenic aerosols currently account for about 10% of the total mass concentration of aerosols over the globe and this amount is associated with a high degree of regional variation (IPCC, 2007). During the last decades, epidemiological studies have identified a link between pollution by airborne particulate matter (PM) and health hazards such as respiratory (allergies, asthma, altered lung function) and cardiovascular diseases (Dockery et al., 1993). Particle toxicity depends on its concentration and chemical composition, but also and foremost on its size since the smallest aerosols are recognized to be the most harmful given that they can reach human breathing apparatus down to the pulmonary cells (Donaldson et al., 1998). At urban scale, issues addressing atmospheric pollution concern pollution peaks forecasting and their corresponding factors, both influencing the air quality and its impact on public health. This becomes an important research area in megacities, which are in full expansion and whose number is expected to increase



Correspondence to: J.-C. Raut
(jean-christophe.raut@latmos.ipsl.fr)

during the 21st century. In such large urban centers, local experimental studies have already been devoted to the microphysical, chemical and optical characterization of urban aerosols: Athens (Kambezidis et al., 1995), Los Angeles (e.g. Lurmann et al., 1997), Sao Paulo (e.g. Landulfo et al., 2003), Marseilles (e.g. Mestayer et al., 2005) or Paris (e.g. Randriamiarisoa et al., 2006). In such an industrialised region, the main source of photo-oxidant pollution turns out to be automobile traffic (e.g. Menut et al., 2000; Chazette et al., 2005b). Air quality over Paris is continuously surveyed by a dedicated surface network carrying out measurements of critical pollutant concentrations, aerosols in particular (AIR-PARIF, <http://www.airparif.asso.fr/>).

However, surface measurements are not sufficient to fully understand the pollutants dynamics and chemistry. Forecasting pollution peaks are mainly constrained using ground-based observations (e.g. Hodzic et al., 2006; Tombette et al., 2008). The improvement of PM prediction, required for driving emission reduction strategies, needs a thorough understanding of the processes affecting aerosol concentrations as well as vertically-resolved measurements in the atmospheric column. This can now be achieved thanks to the new generation of portable lidar systems developed in the past five years (e.g. Raut and Chazette, 2007; Chazette and al., 2007). Such active remote sensing instruments document the mid and lower troposphere by means of aerosol optical properties. Yet, health standards defined in Europe and especially in France (European directive no. 1999/30/CE of 22 April 1999) for particulate pollution impose a monitoring from surface measurements of aerosol mass concentrations in terms of aerodynamic diameter lower than 10 μm (PM₁₀). Such a boundary has been reduced to 2.5 μm (PM_{2.5}) in United-States (e.g. Parkhurst et al., 1999). As an alternative to gravimetric measurements, one direction is to use the optical properties of the aerosols to estimate their abundance. As a consequence, the challenge is to convert vertically-resolved optical measurements into mass concentrations and this requires instruments with a fine vertical resolution.

We focus in this study on the retrieval of PM₁₀ from ground-based and mobile lidar systems over the Paris megalopolis area (~12 million inhabitants). The choice of this agglomeration has been driven by the need of a sufficiently representative database to establish a reliable relation between mass concentration and optical properties of pollution aerosol. It is the case in Paris area, where a great number of research programs have been carried out from 1998 with the ESQUIF project (Vautard et al., 2003). Hence, this study relies on observations performed during ESQUIF in 1999 (Chazette et al., 2005b; Raut and Chazette, 2008a), MEAUVE (Modélisation des Effets des Aérosols en Ultra Violet et Expérimentation) in 2001 (Lavigne et al., 2005), LISAIR (Lidar pour la Surveillance de l'Air) in 2005 (Raut and Chazette, 2007) and ParisFog in 2007 (Elias et al., 2009; Haeffelin et al., 2009) campaigns.

This paper investigates for the first time the vertical PM₁₀ distributions from mobile measurements carried out from locations along the Paris Peripherique (highly trafficked beltway around Paris) and examines the horizontal gradient of pollution between Paris centre and its remote suburbs. Section 2 presents the lidar systems involved in the experiment, as well as in situ instrumentation. Section 3 describes the methodology to derive mass from aerosol optical parameters. The corresponding results obtained during the different campaigns are presented in Sect. 4. Observations of the different aerosol layers converted in PM₁₀ are described in Sect. 5. In Sect. 6 we discuss the relationships between the surface mass concentration measurements performed by the operational air quality network and the lidar-derived mass concentration profiles. We conclude in a seventh section.

2 Experimental set-up

2.1 Location sites

Specific experiments involving both lidar systems and ground-based in situ instruments were performed in Paris area (UTC+2h) from 1999 to 2007. In this study, we consider four experimental sites corresponding to urban, periurban and rural locations. Two experiments were performed in Paris (48°51'24" N, 2°21'07" E): the ESQUIF campaign in July 1999 (Chazette et al., 2005b; Randriamiarisoa et al., 2006) and the LISAIR experiment in May 2005 (Raut and Chazette, 2007). The periurban sites are located at ~15 km in the south-west of Paris: Saclay (48°43'51" N, 2°10'21" E) in the framework of the ESQUIF project in July 1999 (Randriamiarisoa et al., 2006) and Palaiseau (48°42'52" N, 2°14'45" E) (Elias et al., 2009; Haeffelin et al., 2009) during the 6-months long ParisFog campaign carried out from October 2006 to March 2007. The rural site is Brétigny (48°36'41" N, 2°18'21" E), ~30 km far from Paris, where MEAUVE experiment was conducted in March 2001 (Lavigne et al., 2005).

2.2 Instruments

The *Lidar Aérosol UltraViolet (LAUV-EZ lidar[®])* system is a homemade prototype backscatter lidar emitting in the ultraviolet developed by the Commissariat à l'Energie Atomique (CEA) and the Centre National de la Recherche Scientifique (CNRS) (Chazette et al., 2007). It is now commercialized by the LEOSPHERE Company under the name of EZ Lidar[®] (www.leosphere.com). It is designed to monitor the aerosol dispersion in the low and middle troposphere. It operates with a Nd:Yag laser at the wavelength of 355 nm. It is light, compact, eye-safe and suitable for a mobile platform. The resolution along the line of sight is 1.5 m. For this experiment, it was operated onboard a small personal vehicle. The advantage of such a small car is its ability to follow small

atmospheric features due to its maneuverability. The lidar measurement is associated with an overlap factor close to 1 at ~ 150 m above the ground level (agl).

The *Lidar pour l'Etude et le Suivi de l'Aérosol Atmosphérique (LESAA)* was developed by the Commissariat à l'Energie Atomique (CEA) to measure the atmospheric reflectivity at 355 or 532 nm in the lower troposphere over polluted areas. It is a part of the instrumental payload of the Mobile Aerosol Station (SAM; Chazette et al., 2005a). LESAA uses aerosol backscattering to examine the lower troposphere structure with a vertical resolution of 7.5 m (Chazette et al., 2005a). The sky background radiance is measured from the lidar signal at high altitude (45 to 55 km) where the laser beam is considered to be negligible. The lidar measurement is associated with an overlap factor close to 1 at ~ 200 m above the ground level (a.g.l.). Both LESAA and LAUV lidars worked at 20 Hz. Averaging thousand lidar shots every 50 s provided lidar profiles every minute.

A three-wavelength (450, 550 and 700 nm) nephelometer (manufactured by TSI[®]) Model 3563 was used onboard SAM to measure the aerosol scattering coefficient of particles with a diameter larger than $0.05 \mu\text{m}$ in a $7\text{--}170^\circ$ range of scattering angle (Bodhaine et al., 1991) every minute at a flow rate of 20 L min^{-1} . Calibration procedure is done by the constructor before each intensive experiment using two span gases and comparing the results between air (low span) and CO₂ (high span). The three wavelength instrument scattering chamber was maintained at about 35–40% relative humidity. To take into account the non-observed scattering angles, a correction factor has been assessed, from Mie computations, to be close to 1.03 for urban aerosols. Relative uncertainty on the measurements is close to 5%.

The particle measurements were performed using automatic *TEOM (Tapered Element Oscillating Microbalance, Rupprecht and Pataschnik)* Model 1400 A instruments equipped with a PM₁₀ inlet at a flow rate of $1 \text{ m}^3 \text{ h}^{-1}$. The air is heated inside the instruments to 50°C in order to remove any water on the particles. Mass concentrations are obtained every minute with a maximum absolute uncertainty of $5 \mu\text{g m}^{-3}$. TEOM instruments belong to the operational air quality network AIRPARIF or are included in the SAM payload.

The *Electrical Low Pressure Impactor (ELPI)* is a real-time particle size spectrometer designed at the Tampere University of Technology (Delkati, Ltd., <http://dekati.com/cms/elpi>) for real-time monitoring of aerosol particle size distribution (Keskinen et al., 1992). It has been recently included in the instrumental payload of SAM. The ELPI measures airborne particle size distribution from 0.028 to $10.03 \mu\text{m}$ within 12 channels every minute with a flow rate of 10 L/min . The accuracy on the aerosol number concentration measurement is about 5%.

Aerosol samples devoted to carbonaceous analyses were collected on pre-cleaned *Whatman GF/F glass-fiber filters*. The carbon mass was determined through a thermal protocol (Cachier et al., 1989). The precision of the results is estimated to be of the order of 10%. *Whatman nuclepore membranes* in polycarbonate were also mounted on stack filters unit. Those filters were used for measuring the major water soluble (WS) inorganic cations (Na^+ , NH_4^+ , K^+ , Mg^{2+} , Ca^{2+}) and anions (Cl^- , NO_3^- , SO_4^{2-}) in the particle by ion chromatography (DIONEX DX600). The sensitivity of ionic analyses is ~ 0.1 ppb and blank concentrations are lower than 30 ppb for each species. The precision on ion chromatography analysis has been evaluated to be 5–10% (Chazette and Liouise, 2001).

A *Vaisala meteorological probe* type PTU200 onboard SAM was used to measure the temperature (with an uncertainty of ~ 0.1 K), the relative humidity ($\sim 1\%$), and the atmospheric pressure (~ 1 hPa).

To study the influence of air masses motion in the low troposphere, a two-dimensional *SONIC anemometer* was used at the location of in situ instrumentation. It provides real-time and continuous measurements of air velocity in horizontal directions and also temperature.

3 Methodology to convert aerosol extinction coefficient into PM₁₀

3.1 General context

Studies devoted to relationships between PM concentrations and optical thicknesses retrieved from passive remote sensing instruments have received considerable attention and have underlined a good potential for aerosol optical properties to be used in air quality studies (Shinozuka et al., 2007). Chu et al. (2003) showed a fairly reasonable correlation between daily averaged values of PM₁₀ and aerosol optical thicknesses (AOT) at 550 nm derived from AERONET (Aerosol Robotic Network; Holben et al., 1998). Similarly in Alabama, Wang and Christopher (2003) linearly correlated mean hourly PM_{2.5} measurements from a TEOM and satellite MODIS (Moderate Resolution Imaging Spectroradiometer) derived AOT at 550 nm.

Those studies were however limited by the total aerosol loading over the whole atmospheric column. Pelletier et al. (2007) found that a linear model failed at explaining the data well but that the performance could be significantly improved when such a linear relationship would be conditioned on auxiliary parameters, mainly meteorological variables. The knowledge of vertical profiles of PM concentrations would be also an important step for public health related studies. Liu et al. (2004) proposed annual mean ground-level PM_{2.5} concentration maps using the Multiangle Imaging Spectroradiometer (MISR) AOT over the continuous United States but their study included vertical information from a

chemical transport model. Van Dokelaar et al. (2006) suggested that the relative vertical profile of aerosol extinction is the dominant parameter in determining the spatial variation between AOT and PM_{2.5} over North America.

Hence, complementary to the previous studies, we try to establish in this section the link between aerosol extinction coefficient and PM₁₀ concentration measured at surface. Such relationships will be then extended to vertical profiles of extinction coefficients as retrieved from lidar measurements.

3.2 Empirical linear relationship between PM₁₀ and aerosol extinction coefficient

Linear relationships between PM₁₀ and aerosol extinction coefficient at 355 nm ($\alpha_{\text{ext},355}$) have been fully investigated in Raut et al. (2009). They can be written under the form:

$$\text{PM}_{10} = k_1 \cdot \alpha_{\text{ext},355} \quad (1)$$

At the surface, the extinction coefficient is not measured directly. It is derived from the aerosol scattering coefficient at 700 nm $\alpha_{\text{scat},700}$, the scattering Angström exponent between 450 and 700 nm a , both retrieved from the nephelometer, and the single-scattering albedo at 355 nm $\omega_{0,355}$:

$$\text{PM}_{10} = \frac{k_1}{\omega_{0,355}} \cdot \left(\frac{355}{700}\right)^{-a} \cdot \alpha_{\text{scat},700} \quad (2)$$

Equation (2) assumes that the Angstrom exponent between 355 and 450 nm is the same between 450 and 700 nm. The absolute uncertainty on the Angström exponent including this effect has been assessed to be $\sim 0.05 \omega_{0,355}$ is determined from Mie calculations performed with the number size distribution and the complex refractive index. This latter can be obtained through a simple volume-linear internal mixing rule from the refractive indices of the aerosol chemical compounds measured on the filters. An alternative approach to retrieve the complex refractive index, developed by Raut and Chazette (2007, 2008a), lies on a closure study between active and remote sensing observations as well as in situ measurements. The effect of aerosol non-sphericity onto the aerosol scattering and extinction properties has been evaluated through a T-Matrix algorithm (Mishchenko et al., 1997). We considered a mixing of randomly oriented prolate and oblate particles with an aspect ratio between 1.7 and 2. We observed that the discrepancy between spherical and spheroidal particles on optical properties integrated over the whole size distribution was very weak (lower than 1% for the extinction coefficient and 2% for the scattering coefficient). This result is in agreement with the conclusions of Mishchenko et al. (1997) showing that the average of optical properties retrieved for various aspect ratios was close to the spherical case. The resulting uncertainty on the single-scattering albedo is lower than 1%. The corresponding uncertainty δ_{k_1} on k_1 is given by the respective uncertainties

δ_{ω_0} , δ_a , and δ_{C_0} on $\omega_{0,355}$, a and C_0 , the slope of the regression analysis between $\alpha_{\text{scat},700}$ and PM₁₀:

$$\frac{\delta_{k_1}}{k_1} = \left[\left(\frac{\delta_{\omega_0}}{\omega_0}\right)^2 + \left(\frac{\delta_{C_0}}{C_0}\right)^2 + \left(\delta_a \cdot \ln\left(\frac{355}{700}\right)\right)^2 \right]^{1/2} \quad (3)$$

We can notice that the uncertainties in the nephelometer measurements do not explicitly appear in Eq. (3). However, these uncertainties have been taken into account in the calculations. Indeed, the uncertainty δ_{C_0} on C_0 depends on the uncertainties in the PM₁₀ concentrations and in the aerosol scattering coefficient at 700 nm measured by the nephelometer.

3.3 Theoretical relationship between PM₁₀ and extinction coefficient

Aerosol extinction coefficient and PM₁₀ are related at the instant (i) by

$$\text{PM}_{10}^{(i)} = \rho \cdot \frac{4}{3} \pi \cdot \frac{\overline{r^3}^{(i)}}{\overline{\sigma_{\text{ext},355}}^{(i)}} \cdot \alpha_{\text{ext},355}^{(i)} \quad (4)$$

Where $\overline{r^3}^{(i)}$ is the mean cubic radius, calculated from the size distribution and $\overline{\sigma_{\text{ext},355}}^{(i)}$ is the mean extinction cross-section at 355 nm over the size range considered at the instant i , and ρ is the density of particles. In this paper, the density is estimated using the mass of ions, organic compounds and black carbon reported on the chemical filters analyses. The theoretical relationship has been established by Raut et al. (2009) as:

$$\text{PM}_{10}^{(i)} = k_2 \cdot \alpha_{\text{ext},355}^{(i)} \quad (5)$$

where k_2 , independent of time, is given by the least-squares method as:

$$k_2 = \rho \cdot \frac{4}{3} \pi \cdot \left(\overline{\sigma_{\text{ext},355}}^T \cdot \overline{\sigma_{\text{ext},355}}\right)^{-1} \cdot \overline{\sigma_{\text{ext},355}}^T \cdot \overline{r^3} \quad (6)$$

with $\overline{\sigma_{\text{ext},355}}^T$ representing transpose of the vector

$$\overline{\sigma_{\text{ext},355}} = \begin{pmatrix} \overline{\sigma_{\text{ext},355}}^{(1)} \\ \vdots \\ \overline{\sigma_{\text{ext},355}}^{(i)} \\ \vdots \\ \overline{\sigma_{\text{ext},355}}^{(n)} \end{pmatrix}.$$

The dimension of $\overline{\sigma_{\text{ext},355}}$ is n ($i=1$ to n) corresponding to the number of temporal samples. In all the experimental campaigns, measurements are recorded every minute. k_2 represents the mean value over the considered experiment.

3.4 Correction of the relative humidity influence

A common feature of anthropogenic aerosols is their ability to absorb water vapour, whose state is often represented as

a function of relative humidity (RH). As RH increases, condensation of water vapor may take place on the aerosol scatterers depending on their chemical composition (Tang and Munkelwitz, 1993). This phenomenon leads to the effects of deliquescence (hygroscopic growth of aerosols), crystallisation, hysteresis and existence of metastable droplets. Apart from the change in size, hygroscopic aerosols experience a change in their refractive index and in several key optical properties (scattering and absorption coefficients, single scattering albedo, asymmetry parameter, and aerosol optical depth). Quantification of aerosol microphysical and optical properties and their dependency on relative humidity is needed to reduce the large associated uncertainty (Penner et al., 1994). Hänel (1976) described aerosol growth in parameterising particle radius and refractive index for wet particles:

$$r_w = r \cdot \left(\frac{1 - \text{RH}}{1 - \text{RH}_{\text{ref}}} \right)^{-\varepsilon} \quad (7)$$

$$n_w = n_{\text{H}_2\text{O}} + (n - n_{\text{H}_2\text{O}}) \cdot \left(\frac{r_w}{r} \right)^{-3} \quad (8)$$

The suffix *w* refers to wet conditions and RH is the relative humidity. *r* and *n* are the radius and the refractive index of aerosol particles, respectively, at RH = RH_{ref}, and *n*_{H₂O} is the refractive index of pure water. The coefficient ε depends on the considered type of aerosol. It is determined in this study from ISORROPIA model (Nenes et al., 1998; <http://nenes.eas.gatech.edu/ISORROPIA>) since growth curves of water mass condensed around the aerosol can be parameterised versus humidity. As a consequence, ε is determined as follows (e.g. Randriamiarisoa et al., 2006):

$$\varepsilon = -\frac{1}{3} \cdot \log \left(\frac{1 + \frac{m_{\text{H}_2\text{O}} \cdot \rho_{\text{dry}}}{m_{\text{dry}}}}{1 + \frac{m_{\text{H}_2\text{O,ref}} \cdot \rho_{\text{dry}}}{m_{\text{dry}}}} \right) \cdot \left(\log \left(\frac{1 - \text{RH}}{1 - \text{RH}_{\text{ref}}} \right) \right)^{-1} \quad (9)$$

*m*_{H₂O} (respectively *m*_{H₂O,ref}) is the mass of water at RH (resp. RH_{ref}). *m*_{dry} and ρ_{dry} represent the mass and density of dry aerosol, respectively. The knowledge of *r*_w and *n*_w then allows the derivation of the wet scattering coefficient of the aerosol through Mie computations. In this study, ISORROPIA model was used to determine *m*_{H₂O} in a “reverse mode”, in which known quantities are temperature, RH and the aerosol phase concentrations of NH₃, H₂SO₄, Na, HCl and HNO₃ determined from ionic analyses on the chemical filters. Both states of aerosol were considered: the thermodynamically stable state, where salts precipitate once the aqueous phase becomes saturated, and the metastable state, where the aerosol is composed only of a supersaturated aqueous phase.

When using lidar to remotely sense properties in the boundary layer, an accurate description of this effect becomes important to avoid ambiguous interpretation of lidar backscatter data (e.g. Chazette et al., 2005b). It is important to distinguish the contribution of aerosol concentration from that of condensed water in the variations of the backscatter

coefficient. In translating remotely sensed extinction coefficient into dry aerosol mass, RH needs to be considered. In our case, the RH profile has passed the deliquescence point on the 27 May 2005 during the LISAIR campaign. Hence a correction has been applied using the increasing branch of the hysteresis (Fig. 1a) so as to exclusively consider the dry aerosol mass of the particles.

Furthermore, the use of nephelometer measurements is problematic when RH of the environment has passed the deliquescence point since particles can exist on the upper branch of the hysteresis curve. In entering the nephelometer chamber, RH decreases generally down to 30–40% and the droplets reduce their size by evaporation of water but they can exist below the deliquescence point as metastable supersaturated solutions. Such situations were encountered during experiments conducted in winter and early spring, i.e. in Brétigny (MEAUVE campaign) and Palaiseau (during Paris-Fog campaign), where the outside RH was between 70 and 95%. Thus, the hysteresis cycle of RH for those aerosols has been retrieved from ISORROPIA model (Fig. 1b and c) and the upper part of the cycle (efflorescence curve) has been used to translate scattering coefficients measured by the nephelometer into dry scattering coefficients, which could be compared to TEOM measurements (Sect. 3.2).

In Fig. 1, particles show the hysteresis effect in the growth curve. Starting at low RH, the dry particles do not change their size substantially until they reach the deliquescence point (between 45% and 65%) and a solution droplet is formed. There is some degree of water uptake prior deliquescence, which is most probably caused by water adsorption on imperfection sites of the aerosol matrix (Gysel et al., 2002). A further increase of the RH leads to particle growth by condensation of water in accordance with Köhler theory. At decreasing RH, the droplets reduce their size by evaporation of water, pass the saturation point, and continue to evaporate in the supersaturated concentration region. These results are in agreement with experiments reported in the literature, generally for ammonium sulfate aerosol (Tang and Munkelwitz, 1993; Tang and Munkelwitz, 1994; Badger et al., 2006; Gysel et al., 2004). The crystallisation point as predicted by the model is very low (RH < 15%).

In urban and periurban conditions (Fig. 1a and c, respectively), the hysteresis cycles are very similar. The deliquescence point is found a bit larger in rural conditions. The main salts formed together with the variations in RH are (NH₄)₂SO₄ and NH₄NO₃ during LISAIR (Fig. 1a) and ParisFog (Fig. 1c) campaigns. During the latter, a part of the salts formed is also constituted by Na₂SO₄. Conversely, during MEAUVE campaign in rural conditions (Fig. 1b), the main inorganic salts formed onto the aerosols are Na₂SO₄ and NH₄NO₃. According to Tang (1996), the theoretical deliquescence points of these individual salts are 80% for (NH₄)₂SO₄, 62% for NH₄NO₃ and 84% for Na₂SO₄. The predominance of Na₂SO₄ salt in the rural location with a larger deliquescence point tends to explain the discrepancies

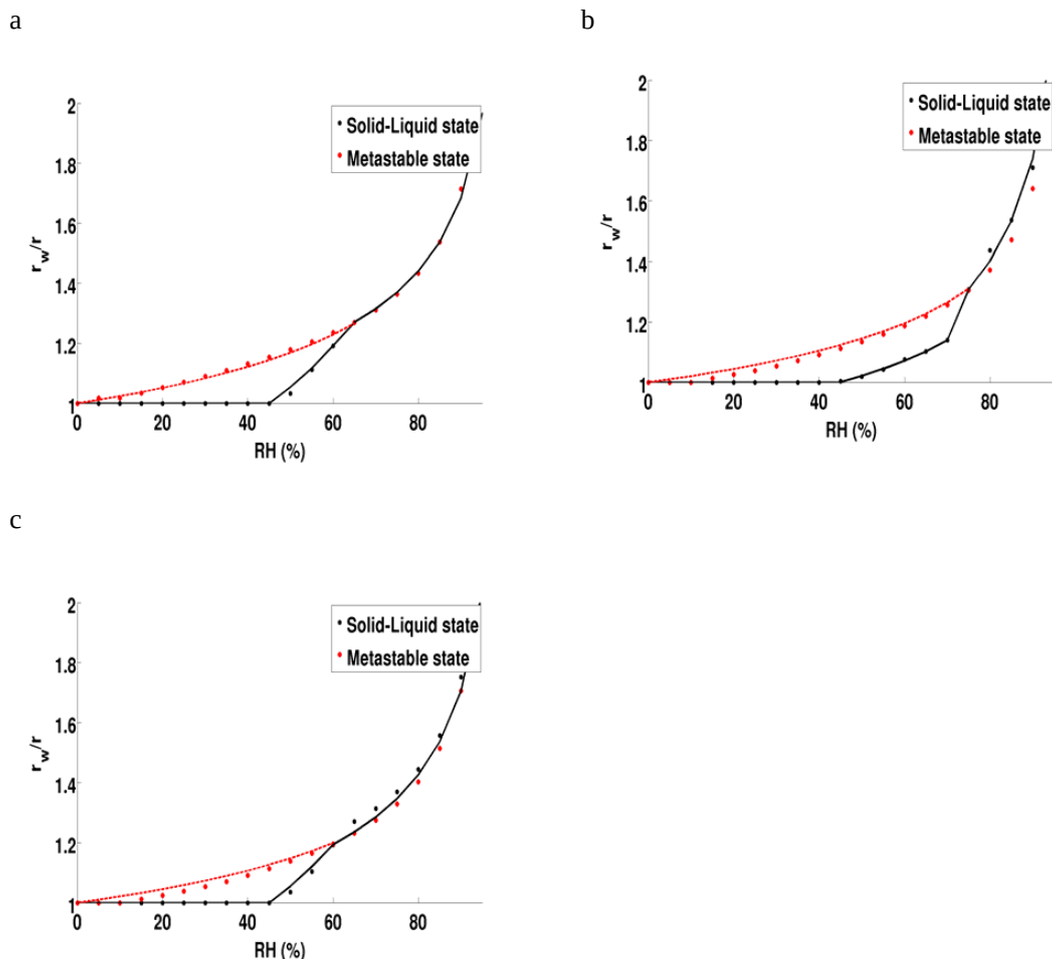


Fig. 1. Hysteresis cycle of phase transformation, growth and evaporation of aerosols derived from ISORROPIA model as a function of RH at ambient temperature in Paris during LISAIR (a), in Brétigny during MEAUVE (b) and in Palaiseau during ParisFog (c).

observed between the different situations in Fig. 1. We have to bear in mind that ISORROPIA model does not take into account organic compounds that can also be soluble products. Besides, differences can occur between summer and winter situations. A bias is therefore expected on our calculations. Because hysteresis cycles are only used in this study for deriving corrections on scattering coefficients, this effect is assumed to be low.

4 Optical to mass relationship for different aerosol types

4.1 Optical and microphysical parameters required to derive optical to mass relationships

Simple linear relationships between $\alpha_{\text{scat},700}$, measured in real time by the nephelometer and corrected from the non-observed angles and RH, and PM₁₀, measured in real time with the TEOM, have been investigated during the different

campaigns. Results are plotted in Fig. 2. Various patterns of particulate pollution in the Paris area are considered. Figure 2a and b show the correlations obtained in urban conditions, i.e. directly in Paris centre. The correlation is much better during LISAIR program (Fig. 2a) with a correlation coefficient $R \sim 0.67$ due to a considerable set of measurements than during ESQUIF ($R \sim 0.39$). Despite the limited number of samples analyzed in Fig. 2b, the slopes are in agreement: C_0 is $\sim 0.98 (\pm 0.10) \text{ g m}^{-2}$ during LISAIR (Fig. 2a) and $\sim 1.10 (\pm 0.10) \text{ g m}^{-2}$ during ESQUIF in Paris (Fig. 2b). Two cases belong to the periurban situations: ParisFog in Palaiseau and ESQUIF in Saclay (respectively Fig. 2c and d). During ParisFog campaign, a high number of real time observations were performed, which leads to a nice correlation between $\alpha_{\text{scat},700}$ and PM₁₀ ($R \sim 0.72$). The corresponding slope ($\sim 0.82 \pm 0.10 \text{ g m}^{-2}$) is lower than that retrieved in Paris. It is confirmed by the result obtained in Saclay with $C_0 \sim 0.77 \pm 0.10 \text{ g m}^{-2}$ ($R \sim 0.37$). It is worthy to notice that these campaigns were carried out during different

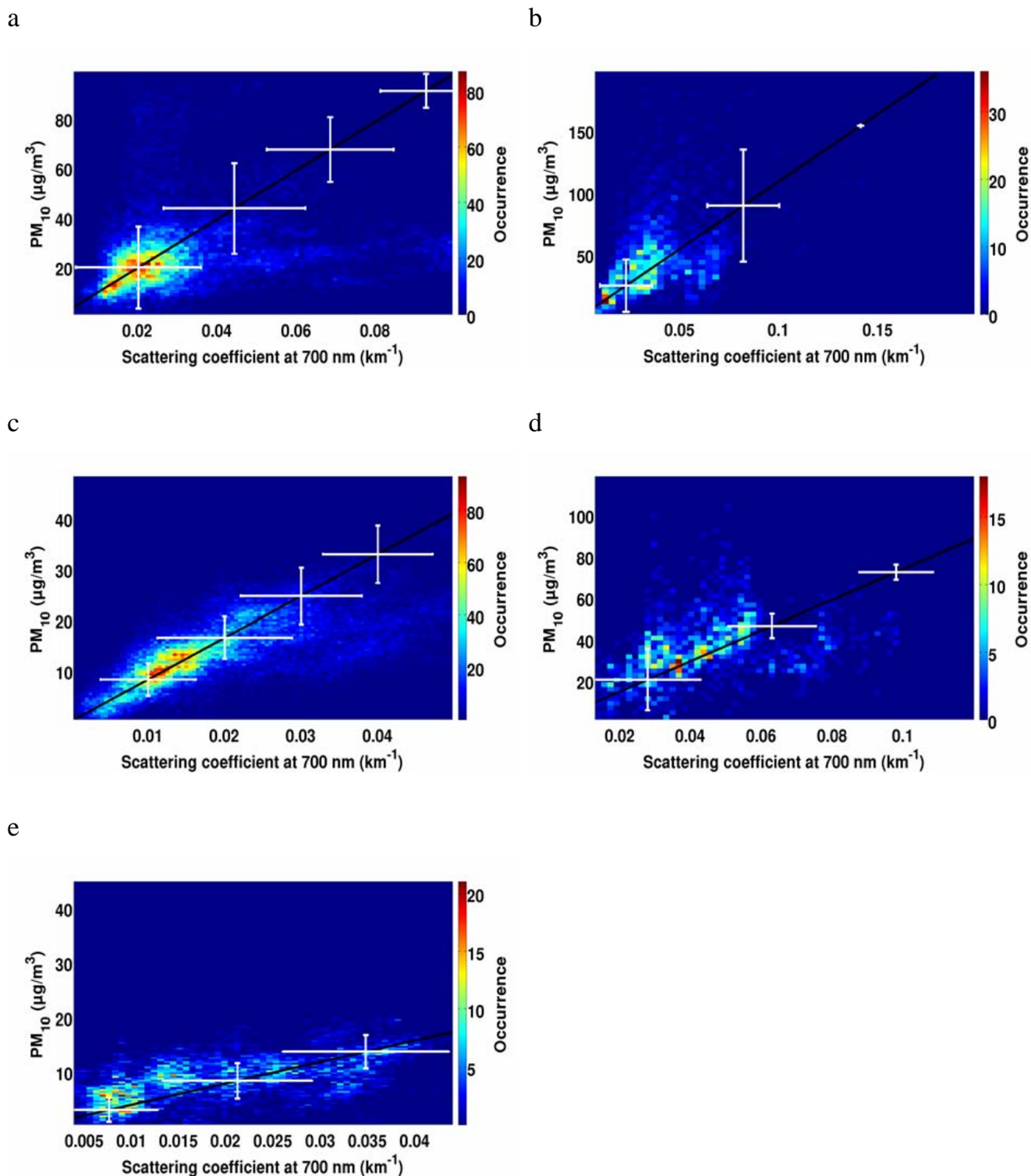


Fig. 2. Correlations between the aerosol scattering coefficient at 700 nm measured by the nephelometer and PM₁₀ concentrations measured by the TEOM in urban conditions in Paris during LISAIR (a) and ESQUIF (b), in periurban conditions in Palaiseau during ParisFog (c) and in Saclay during ESQUIF (d), and in rural conditions in Brétigny during MEAUVE (e). White horizontal and vertical bars represent the corresponding standard deviations in terms of scattering coefficient and PM₁₀, respectively.

Table 1. Optical and microphysical parameters of different types of aerosols enabling the retrieval of specific extinction cross-sections ($s_{\text{ext},355}$). k_1 coefficient is determined from the knowledge of the slope C_0 of optical to mass relationship and from the corresponding values of single scattering albedo ($\omega_{0,355}$) and Angström exponent (a). k_2 coefficient is determined from the knowledge of mean density ρ , cubic radius r^3 and extinction cross-section $\sigma_{\text{ext},355}$ of aerosols.

Aerosol type	C_0 ($10^{-1} \times$ g m^{-2})	$\omega_{0,355}$	a	k_1 ($10^{-1} \times$ g m^{-2})	ρ (g cm^{-3})	r^3 (μm^3)	$\sigma_{\text{ext},355}$ (cm^2)	k_2 ($10^{-1} \times$ g m^{-2})	$s_{\text{ext},355}$ ($\text{m}^2 \text{g}^{-1}$)
Urban	9.81	0.89	2.07	2.17	1.77	1.18×10^{-4}	3.96×10^{-11}	2.20	4.5
Periurban	8.21	0.93	2.15	1.77	1.68	1.69×10^{-4}	6.98×10^{-11}	1.71	5.9
Rural	3.86	0.91	1.36	1.40	1.67	3.62×10^{-4}	1.83×10^{-10}	1.38	7.1
Biomass burning	–	0.77	1.74	–	~ 2.00	1.64×10^{-3}	3.54×10^{-10}	3.90	2.6
Dust	–	0.94	~ 0.8	–	~ 2.00	7.03×10^{-3}	6.72×10^{-10}	8.76	1.1

seasons: in winter and early spring for ParisFog and in summer for ESQUIF. Regardless of the season, and therefore of the type of organic matter present in the particles, the similarities in C_0 for polluted aerosols is remarkable given the possible discrepancies in aerosol size distributions and chemical composition. This result is in agreement with the conclusions of Carrico et al. (2003). Only one case (Brétigny during MEAUVE) involved a rural situation influenced by pollution in the Paris area (Fig. 2e). The slope C_0 is smaller than in the previous cases (0.39 g m^{-2}) but was accompanied with a larger uncertainty (25%) and a lower correlation coefficient ($R \sim 0.62$) than for the regression analyses performed for LISAIR and ParisFog experiments (Fig. 2a and c). This is because the experiment performed in Brétigny was done during winter 2001. Owing to rainy weather, only scarce measurements were available and the corresponding aerosol loading in the vertical column was very variable.

The slopes C_0 of optical to mass relationships obtained for the described situations encountered are given in Table 1, together with the corresponding values of single scattering albedo and Angström exponent. Urban aerosols are smaller ($a \sim 2.02$) and more absorbing ($\omega_{0,355} \sim 0.89$) than those presenting a rural origin. This is most likely due to the high proportion of small black carbon particles in Paris streets because of the importance of vehicle traffic and combustion processes. When getting away from Paris in direction of the suburbs, aerosols have time to evolve and to gather chemical components and the developed coating shell increases their sizes. This result is also suggested by the values of r^3 and $\sigma_{\text{ext},355}$ averaged over the whole campaigns and required to solve Eq. (4). r^3 and $\sigma_{\text{ext},355}$ have been computed from the knowledge of the size distribution and the complex refractive index. During LISAIR, Raut and Chazette (2007) have found a mean size distribution with nucleation and accumulation modes. No coarse mode was observed. The modal radii were 0.03 and 0.08 μm , both having a geometric deviation of 1.5. 94% of particles were in the fine mode. During ESQUIF, Chazette et al. (2005b) reported in the mixed layer

a size distribution with two modes with corresponding modal radii of 0.03 and 0.07 μm , both having a standard geometric deviation of 1.5. The first mode represented 83% of the total number of particles. The corresponding complex refractive indices, needed to determine $\sigma_{\text{ext},355}$ and $\omega_{0,355}$ in Eqs. (3) and (4), have been derived by Raut and Chazette (2008a) through a synergy between lidar and in situ measurements for aerosol located inside (resp. outside) Paris plume: $1.510-0.017i$ (resp. $1.55-0.013i$). In Brétigny, the size distribution looked like that retrieved during ESQUIF experiment but with a larger second modal radius (0.1 μm). The complex refractive index has been computed from chemical analysis and Lorentz-Lorenz formula (Lorentz, 1880; Lorenz, 1880): $1.55-0.019i$.

4.2 Specific cross-section following the two methods

Coefficients k_1 and k_2 have been calculated according to the two approaches described in Sects. 3.2 and 3.3. The value of the mean density ρ of aerosols in the boundary layer, predominantly of urban origin, was required for the calculation of k_2 . Biomass burning and dust events, discussed in Sect. 5.4, were mostly found in the free troposphere during LISAIR experiment, except on 26 and 27 May 2005. Thus, the limited influence of biomass burning and dust plumes on regional pollution of Paris area, hence on the mass-scattering relationship in the boundary layer, justifies the use of a single density value for the size distributions in the boundary layer. We estimated mass of ions, organic compounds and black carbon using chemical analyses during the different campaigns. Retrieved density values are reported in Table 1, present a small variability ($\sim 5\%$) and fall within the density range of urban pollution samples ($1.54-1.77 \text{ g cm}^{-3}$; e.g. McMurry et al., 2002).

The comparison of both methods provides excellent results within a discrepancy of 3%. According to Eq. (3), the relative uncertainty on k parameter depends on the uncertainties on the single-scattering albedo (3%), on the Angström

Table 2. Specific scattering cross-sections for different types of aerosols reported in the literature.

Aerosol type	Specific scattering cross-section (m ² g ⁻¹)	Location	References
Fine PM, Urban	3.1±0.2	New York City (USA)	Waggoner et al. (1981)
	3.5	Houston (USA)	Dzubay et al. (1982)
	2.4–2.5	Southwest USA	White et al. (1994)
	3.2	NE Atlantic	Hegg et al. (1995)
	4.0±0.7	NE Atlantic	Clarke et al. (1996)
	3.2	Ontario, Canada	Hoff et al. (1996)
	2.8	Mid-atlantic coast, USA	Hegg et al. (1997)
	2.3–6	Beijing, China	Bergin et al. (2001)
	2.65	Tessaloniki, Grece	Chazette and Liousse (2001)
	4.10–4.57	Bondville, USA	Koloutsou-Vakakis et al. (2001)
Fine PM, remote site	3.7±0.6	Negev desert, Israël	Ichoku et al. (1999)
	5.8±0.2	Negev desert, Israël	Andreae et al. (2002)
Biomass burning	5.6–5.9	Eastern Mediterranean	Vrekoussis et al. (2005)
	4.06	Simulation (Mie)	Trentmann et al. (2002)
	0.2–3.3	Baltimore, Maryland, USA	Adam et al. (2004)
	2.8–4.2	Fresh aerosol (review)	
	3.5–4.2	Aged aerosol (review)	Reid et al. (2005)
Dust	2.48	South Western Iberia Peninsula	Pereira et al. (2008)
	0.34–0.45	Southwest, USA	White et al. (1994)
	1.1	Aged dust, Atlantic Ocean	Clarke et al. (1996)
	0.83	Aged dust, Barbados	Li et al. (1996)
	0.71±0.04	Negev desert, Israël	Andreae et al. (2002)
	1.05±0.13	Transported dust, China	Alfaro et al. (2003)
	0.21–0.96	Eastern Mediterranean	Vrekoussis et al. (2005)
	0.97	South Western Iberia Peninsula	Pereira et al. (2008)

exponent (3%) and the slope C_0 of the regression analysis between the scattering coefficient and PM₁₀ concentration. The latter varies with the type of aerosol. According to Fig. 2, it has been assessed to ~10% for urban and periurban aerosols and ~25% for rural aerosols. As a consequence, k_1 and k_2 coefficients are provided with a reasonable uncertainty for aerosols of urban and periurban origins (~12%). This confirms that both methods are equivalent and that the simple linear empirical relationship defined in Sect. 3.2 is appropriate to retrieve PM from extinction coefficients retrieved from lidar profiles. In the rural location (Brétigny), the uncertainty on k_1 and k_2 coefficients was larger: 26% (according to Eq. 3). The results of this situation will not be used for optical to mass conversion purposes in Sect. 5: lidar measurements were not performed over such an area.

The inverse of k_1 and k_2 constants is called the specific extinction cross-section at 355 nm: $s_{\text{ext},355}$. Mean values for the different cases have also been mentioned in Table 1. $s_{\text{ext},355}$ was found to be ~4.5 m² g⁻¹ for urban aerosols, and a bit larger for particles observed on a more rural area

5.9–7.1 m² g⁻¹. This is in agreement with results described in many studies (Table 2). They generally exhibit values in the range from 3.5 to 4.5 m² g⁻¹ for fine urban aerosol and between 4 and 6 m² g⁻¹ for fine anthropogenic aerosols in a remote site. Notwithstanding, those results are generally expressed in terms of scattering specific cross-sections in the mid-visible, which can explain, via $\omega_{0,355}$ and a , that our retrievals present slightly larger values. Another bias can be ascribed to the fact that the frequently used cut-off diameter in USA studies is 2.5 μm instead of 10 μm. Nevertheless, the specific cross-sections are very similar regardless of the measurement site and period. As a consequence, k_1 and k_2 values will be used in the next section to convert aerosol extinction coefficient retrieved from lidar signals inversion into PM₁₀ profiles. Indeed, extinction coefficients derived from nephelometer measurements are in very good agreement with those retrieved by the lidar, as shown in Fig. 3. Such an agreement was found on 13 and 14 March 2007, in the framework of the ParisFog campaign, when the lidar was shooting horizontally at the same altitude of the

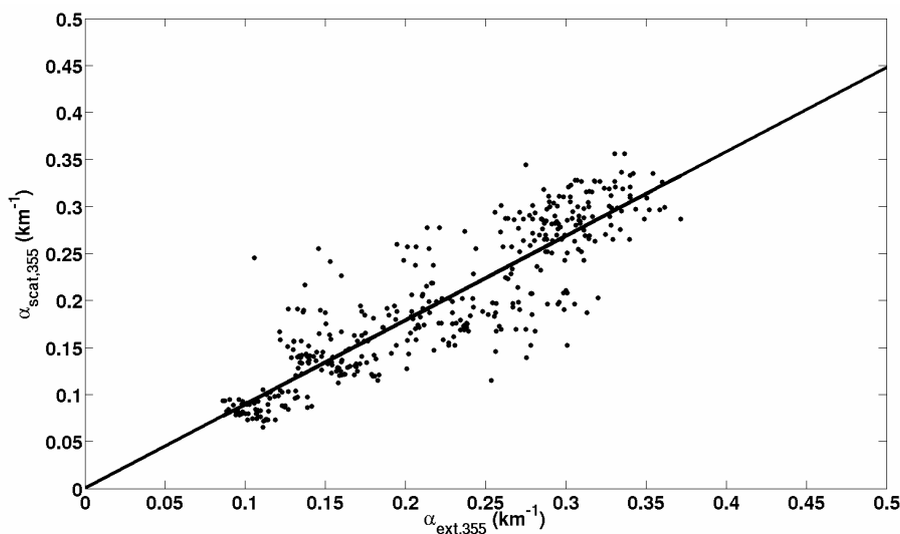


Fig. 3. Correlation between the extinction coefficient at 355 nm retrieved by the lidar when shooting horizontally at ~ 4 m above ground level and the scattering coefficient at the same wavelength derived from nephelometer measurements on 13 and 14 March 2007, during ParisFog campaign.

nephelometer (~ 4 m above ground level). The correlation coefficient is ~ 0.88 and the slope of the regression analysis (~ 0.92) corresponds to the single-scattering albedo of typical periurban aerosols, as shown in Table 1. The approach based on k_1 and k_2 to retrieve PM₁₀ is validated by the fact that the aerosol extinction coefficient derived from the lidar is in accordance with nephelometer measurements (Sect. 4.2 and Fig. 3), which are themselves in agreement with TEOM observations (Sect. 4.1 and Fig. 2). As a consequence the relationship between the aerosol extinction coefficient and the aerosol concentration (PM₁₀) is linear for a given type of aerosol (Table 1).

5 Spatiotemporal variability of PM₁₀ in the troposphere

5.1 Analysis of lidar vertical profiles

As mentioned in Sect. 2.2, the LAUVA system has been operated onboard a small personal vehicle, which permits a regional study of anthropogenic particles in urban environment. Two kinds of experiments have been performed during the LISAIR program in May 2005. The first one has been devoted to the study of the particulate gradient between the large southern suburb of Paris (Saclay or Palaiseau) and Paris centre (city hall of Paris). For the second application, observations have been carried out along the Paris Peripherique, so as to identify its role in the production of anthropogenic aerosols.

5.1.1 Lidar-derived aerosol extinction coefficient

In both cases, the lidar signals have been calibrated, range-corrected, corrected from the overlap factor and inverted using a well-known method based on Bernoulli's differential form of the propagation equation (Klett, 1981). This approach relies on the assumption of a constant BER (backscatter-to-extinction) in the tropospheric column, that has been assessed to be $\sim 0.011 \text{ sr}^{-1}$ ($\pm 0.002 \text{ sr}^{-1}$) at 355 nm through an iterative method converging when the optical thickness retrieved by the lidar is equal to that of the sun-photometer (Chazette, 2003). This column-averaged BER has been determined for both urban aerosols during LISAIR (Raut and Chazette, 2007) and periurban aerosols (Chazette et al., 2005; Raut and Chazette, 2008a) during ESQUIF. Sun-photometer data were actually taken from a stationary site located in Paris or in Palaiseau depending on the considered campaign. The dependence of the BER on RH has been fully investigated in a previous paper from the authors (Raut and Chazette, 2007). BER has been computed as a function of RH through a Mie code using the complex refractive index and the size distribution, both dependent on RH. The result of this previous study performed during LISAIR experiment highlighted a variability of the BER at 532 nm lower than 10% when RH increased from 20% to 70%. But the variability of BER over the same range of relative humidity decreased down to 0.5% at 355 nm. It is due to a compensation between the changes in the single-scattering albedo and in the backscattering phase function with RH.

The retrieved extinction coefficients have been adjusted from hygroscopic growth when the RH profile has exceeded the deliquescence point: it was the case during the night between 26 and 27 May 2005: RH was larger than 65% at the surface and in a layer located between 2.5 and 3 km height. The corresponding vertical profiles of RH were derived from radiosounding measurements performed twice a day in Paris area (Trappes station, 48°46′39″ N, 2°00′09″ E). An appropriate correction factor in each layer detected has been calculated using the increasing branch of the hysteresis (Sect. 3.3) in order to deal only with the dry aerosol mass of the particles. On the 26 and 27 May 2005, the scattering growth factor has been assessed to be ~ 0.69 in the layer located between 2.5 and 3 km height, which leads to a ratio of 0.73 between the wet and the dry aerosol scattering coefficients in that layer. According to the previous discussion, the extinction profiles computed using the constant BER estimate do not have any systematic errors due to differences in water uptake of aerosols in the profile.

5.1.2 Lidar-derived vertical structures

Lidar profiles have generally been obtained during evening or night periods, allowing the detection of multiple layers: boundary layer, nocturnal layer, residual layers and elevated layers in the free troposphere (between 500 m and 1 km thick). The distinction between the different scattering layers in terms of vertical structures has been done through an algorithm sensitive to vertical heterogeneities in particulate concentrations derived from lidar profiles. On each profile, the minimum of the vertical gradient in aerosol extinction coefficient is detected (Dupont et al., 1994; Flamant et al., 1997). This has been done through an analytic derivation of the second order polynomial function retrieved from least mean squares method applied onto the extinction coefficient. Calculations are realized in a sliding window, whose size is close to the thickness of the analysed transition area. The accuracy on the layer altitudes is close to 30 m. This algorithm is then applied to the whole temporal series of measured lidar profiles.

5.2 Lidar-derived PM₁₀ profiles

Once the various layers have been identified, an appropriate specific cross-section is attributed for each of them, according to the study performed in Sect. 4.1. Paris turns out to be one of the greatest urbanized areas in Europe, located far from other big cities so that the signatures and origins of pollution are easier to determine. At a given location, the ground-based instruments have recorded the same slope represented in Fig. 2 whatever the air masses origins have been. This indicates that k_1 and k_2 parameters obtained from the linear relationships are not statistically dependent on wind directions. The discrepancies observed among the different

sites are more likely due to the proximity of particles sources and to the aerosol ageing.

A typical coefficient $k_1=0.217 \text{ g m}^{-2}$ for urban aerosols is applied in the boundary layer or nocturnal layer for pollutants emitted on the urban area or Paris Peripherique. Indeed, the pathway followed by the car when travelling from the urban to the periurban location is only highway. k values are principally governed by the traffic influence, which is a source of “urban” particles on our transects. A typical coefficient $k_1=0.177 \text{ g m}^{-2}$ for periurban aerosols is applied in the residual layers. This is suggested by the fact that those layers are mainly composed with aged aerosols emitted on the previous day and trapped in altitude during the erosion of the boundary layer in the evening. Nevertheless, k coefficient does not linearly vary as a function of altitude since layer structures are well disjoint (Sect. 5.3). The coefficient obtained in the rural site (30 km far from Paris) has not been applied to lidar data because the lidar transects have been performed up to 15 km far from Paris (periurban location).

The uncertainty on the lidar-derived PM₁₀ concentrations is due to the uncertainty on the retrieved aerosol extinction coefficient, the uncertainty on k_1 in traffic (urban) or no traffic (periurban) conditions and the uncertainty in the assumed aerosol type (urban or periurban). The latter is directly related to the difference between k_1 in urban and periurban locations, which is of order of 20% and can be considered as low since error bars on k_1 (12%) overlap. As a consequence, the choice of k_1 parameter does not add an uncertainty larger than 5% in the retrieved PM₁₀ concentrations. The uncertainty on the retrieved aerosol extinction coefficient depends on the lidar signal variability, the choice of the altitude of normalization and the BER assumption. It has been assessed to be of the order of $\sim 10\%$. Uncertainties on the relative humidity corrections are not taken into account in the computation of k_1 and k_2 constants since Eqs. (1) and (5) are determined for dry aerosols. But these uncertainties influence the uncertainties on the dry PM₁₀ concentrations retrieved from the lidar-derived aerosol extinction coefficients that have been adjusted from RH. Finally, the global uncertainty on the lidar-derived PM₁₀ is $\sim 25\%$.

5.3 Local contributions in the boundary and residual layers

Two transects between the southern suburbs (Palaiseau) and Paris centre have been carried out on the 24 and 25 May in nighttime conditions before the morning traffic starts (Figs. 4 and 5, respectively). Both cases highlight the presence of stable layers corresponding to the inversion layer trapping pollutants at low altitudes and the residual layer in altitude. The top of the stable nocturnal layer is between 300 and 400 m on the 24 May (Fig. 4b) and lower than 500 m on the 25 May (Fig. 5b). The altitude of the residual layer was higher on the 24 May (0.8–1.5 km) than on the 25 May (0.4–0.9 km). This can be ascribed to the higher temperatures observed on

Table 3. Mean mass concentrations in terms of PM₁₀ values (layer and time-averaged values) and associated standard deviations in layers detected from the mobile lidar during the LISAIR campaign over the Paris area. Mean altitudes (in km) and thicknesses (in km) of these layers are also specified between brackets.

Date	Diurnal/ Nocturnal Conditions	Figure	1st layer PM ₁₀ ($\mu\text{g m}^{-3}$)	2nd layer PM ₁₀ ($\mu\text{g m}^{-3}$)	3rd layer PM ₁₀ ($\mu\text{g m}^{-3}$)	4th layer PM ₁₀ ($\mu\text{g m}^{-3}$)
24 May	Nighttime	Fig. 4	8±3	10±11	–	–
Palaiseau-Paris	Before traffic		{0.25 km/0.5}	{1.1 km/0.8}		
25 May	Nighttime	Fig. 5	24±5	11±10	–	–
Palaiseau-Paris	Before traffic		{0.17 km/0.35}	{0.6 km/0.4}		
25 May	Nighttime	Fig. 6	32±16	10±11	–	–
Paris	Before traffic		{0.2 km/0.4}	{0.75 km/0.3}		
25 May	Nighttime	Fig. 7	46±41	17±12	–	–
Paris	Traffic conditions		{0.2 km/0.4}	{0.7 km/0.35}		
25 May	Daytime	–	24±16	–	–	–
Paris-Saclay	Boundary layer well developed		{0.3 km/0.6}			
26 May	Nighttime	–	8±6	–	–	–
Paris-Saclay	Before traffic		{0.65 km/1.3}			
26 May	Daytime	–	18±6	3±3	9±14	17±26
Paris	Afternoon		{0.7 km/1.4}	{2 km/0.8}	{2.75 km/0.5}	{3.75 km/0.5}
26 May	Daytime	Fig. 8	13±6	6±3	41±32	13±18
Palaiseau-Paris	Evening		{0.45 km/0.9}	{1.5 km/1}	{2.8 km/0.8}	{3.7 km/0.6}
27 May	Nighttime	Fig. 9	33±25	11±5	28±20	10±15
Paris	Before traffic		{0.5 km/1}	{1.9 km/1.1}	{2.9 km/0.7}	{3.7 km/0.6}

the 23 May ($\sim 23^\circ\text{C}$) than on the 25 May ($\sim 18^\circ\text{C}$) enabling a stronger development of the convective layer during the day and therefore a residual layer trapped at higher altitude in the evening. A weaker large scale subsidence may have also led to a higher residual layer on 24 May. In these layers, mass concentrations are rather constant. PM₁₀ values are reported in Table 3. Concentrations are very similar in the residual layer but important differences can be noticed in the nocturnal layer. PM₁₀ values (layer and time-averaged values) are $\sim 8 \mu\text{g m}^{-3}$ on the 24 May, which is very similar to those derived from a transect carried out on the 26 May between Paris and Saclay (southern suburb of Paris) before traffic in nighttime conditions (Table 3). PM₁₀ values are $\sim 24 \mu\text{g m}^{-3}$ on the 25 May. This is principally due to the lower height of the surface nocturnal layer, preventing dilution on the 25 May. Wind measurements performed from the sonic anemometer indicate very low winds on the 25 May ($\sim 0.5 \text{ m s}^{-1}$), also suggesting weak dilution processes. Such a mass concentration is close to that obtained on the 25 May in a well-developed boundary layer during the transect Paris-Saclay (Table 3).

Two situations around the Paris ring before and during the morning traffic period on the 25 May are illustrated in Figs. 6 and 7 respectively. Together with the increasing activity of the automobile traffic on the morning, surface concentrations significantly rise by $\sim 50\%$ between 05:00 and 08:00 UTC (Table 3). The slow development of the boundary layer, due

to a low surface temperature ($\sim 17^\circ\text{C}$), leads to a mixing of the layers containing residual pollutants with those recently emitted on the surface. Considering the doubling of the mass concentrations in the centre of Paris (from 30 to $60 \mu\text{g m}^{-3}$), the impact on air quality is clearly noticeable.

5.4 Role of the long-range transport

Such an increase of particulate concentrations in the lowest layers of the troposphere has also been observed on the 26 May in the evening (Fig. 8b). This phenomenon occurs after the formation of the nocturnal inversion trapping aerosols that are still emitted at this period in the centre of Paris (St. Michel district). We can also notice a plume in altitude aloft A6 highway at between 500 m and 900 m in the South of Paris linked to particles emitted from automobile traffic a few hours before. This plume appears after the temperature inversion close to the ground and is associated with the nightfall. In addition to the nocturnal inversion layer and the residual layers, observations performed on 26 and 27 May (Figs. 8b and 9b) also highlight elevated layers from a different origin. Indeed, Paris agglomeration has a rather flat topography and is then also strongly influenced by synoptic meteorological conditions. Measurements performed in the vicinity of the city hall of Paris by LESAA lidar confirm those observations. Figure 10a presents the temporal evolution of these layers on the 26 May. The classical development

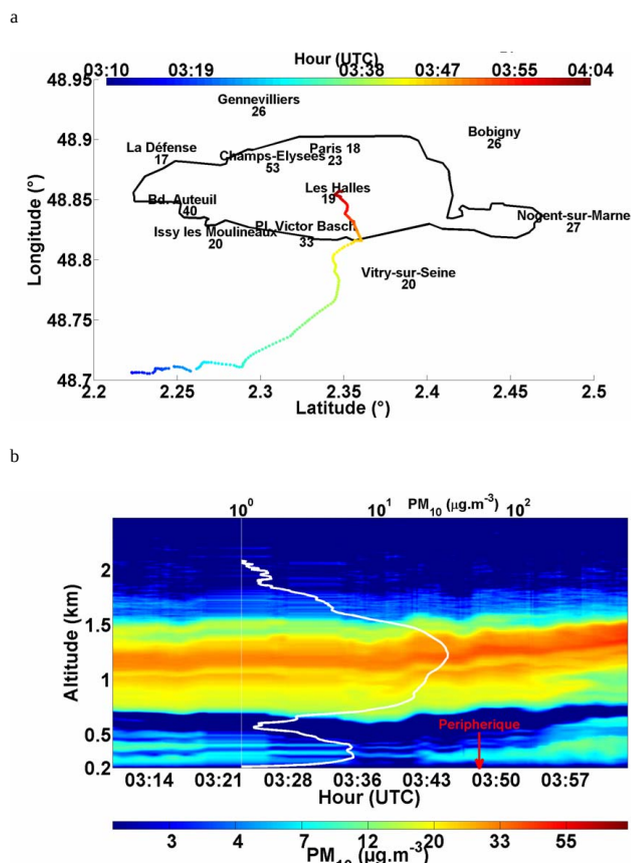


Fig. 4. Route followed by the small vehicle embarking the lidar on the 24 May 2005 from Palaiseau to Paris (a), and corresponding 2-D colour-coded temporal evolution of lidar-derived PM₁₀ concentrations (b). Hourly averaged PM₁₀ values measured by AIRPARIF network stations are reported in panel (a). The mean PM₁₀ profile retrieved from lidar signals is shown in white in panel (b). The black line circling Paris on the map (a) is the geographic demarcation of Paris city. The red arrow in (b) indicates where the lidar track intersected the Peripherique.

of the boundary layer, its mixing with the elevated residual layer at ~12:00 UTC, its erosion in the evening, the development of the nocturnal layer and the corresponding accumulation of pollutants close to the surface are clearly visible on those lidar profiles. We can also notice that the top height of the planetary boundary layer (PBL) is maintained at a fairly constant level in the afternoon. It is due to an anticyclone located over central Europe than maintains high mean sea-level pressure over the Paris area (1015–1018 hPa) and hinders the development of the PBL.

The aerosol extinction coefficient nevertheless shows unexpected large values in the free troposphere all day long and higher values at ~3 km from 15:00 UTC. At this altitude, the aerosols are not characterized by a significant depolarization ratio, whereas the particles located in the highest layer present stronger total depolarization ratios (~0.05), twice

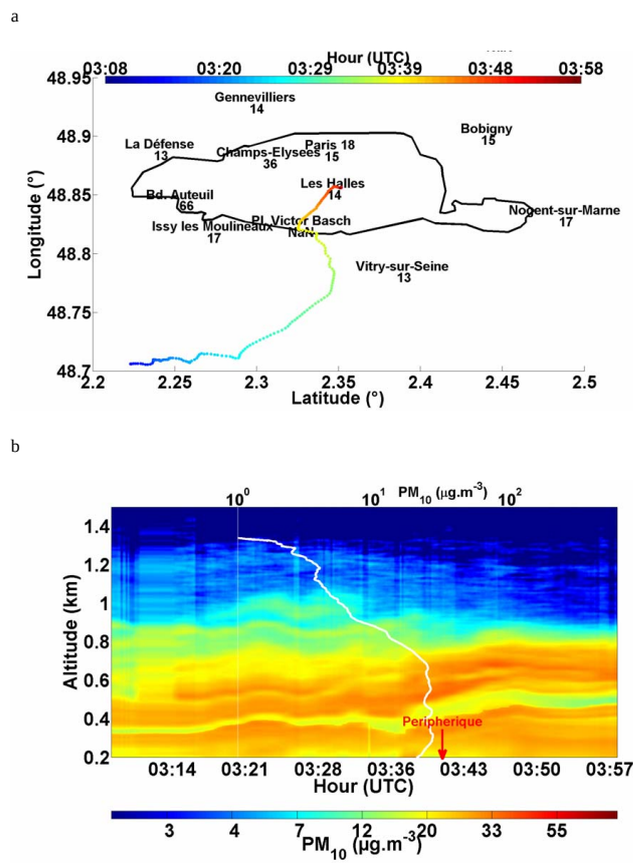


Fig. 5. Route followed by the small vehicle embarking the lidar on the 25 May 2005 from Palaiseau to Paris (a), and corresponding 2-D colour-coded temporal evolution of lidar-derived PM₁₀ concentrations (b). Hourly averaged PM₁₀ values measured by AIRPARIF network stations are reported in panel (a). The mean PM₁₀ profile retrieved from lidar signals is shown in white in panel (b). The black line circling Paris on the map (a) is the geographic demarcation of Paris city. The red arrow in (b) indicates where the lidar track intersected the Peripherique.

larger than the background level. Hence, aerosols within the highest layer could include a mineral contribution associated to non spherical particles whereas the aerosols located in the PBL and in the upper layer appear rather spherical. Such observations are in accordance with the fact that aerosols at ~3 km are rather biomass burning particles and those at ~4 km are dust particles.

The sources of biomass burning aerosols are likely forest fires in Portugal or Spain, accordingly to backtrajectories arriving over Paris on the 26 May at 20:00 UTC (Fig. 10c and d). These five-day backtrajectories of air masses were computed using the Hysplit model (courtesy of NOAA Air Resources Laboratory; <http://www.arl.noaa.gov>). This model uses the meteorological data to compute advection and dispersion of air parcels. Fire locations are given by MODIS (Moderate Resolution Imaging Spectroradiometer)

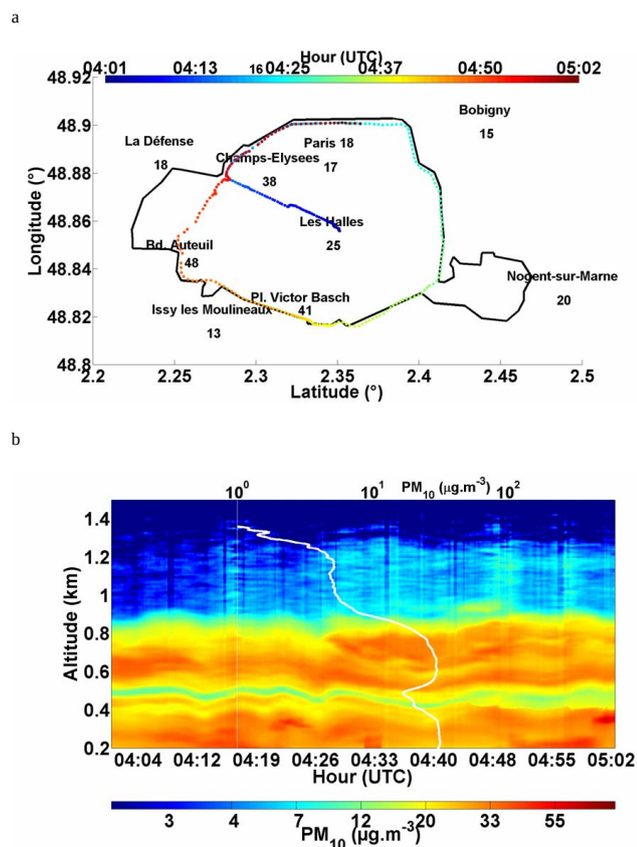


Fig. 6. Route followed by the small vehicle embarking the lidar on the 25 May 2005 the Paris Peripherique before traffic (a), and corresponding 2-D colour-coded temporal evolution of lidar-derived PM₁₀ concentrations (b). Hourly averaged PM₁₀ values measured by AIRPARIF network stations are reported in panel (a). The mean PM₁₀ profile retrieved from lidar signals is shown in white in panel (b). The black line circling Paris on the map (a) is the geographic demarcation of Paris city.

fire products obtained from 21 to 24 May (Roy et al., 2005). Dry convective events can inject such primary aerosols of biomass smoke into the free troposphere especially during summer in Europe. Backtrajectories reaching Paris at 4 km suggest that aerosols in the highest layer may be nonspherical mineral dust coming from the Sahara and transported over the Atlantic Ocean. The simulations performed using the DREAM (Dust REgional Atmospheric Model) model (<http://www.bsc.es/projects/earthscience/DREAM/>) have suggested that dust was actually over the Atlantic Ocean.

These aerosols coming from Portugal, Spain or from the Sahara have different optical and microphysical properties than those retrieved in the vicinity of the surface from in situ measurements. In particular, lidar data inversion with the assumption of a constant BER would have led to erroneous extinction coefficients, and therefore mass concentrations, in the dust layer. The BER profile has been retrieved in this elevated layer to be 0.020 sr^{-1} , while maintaining a constant

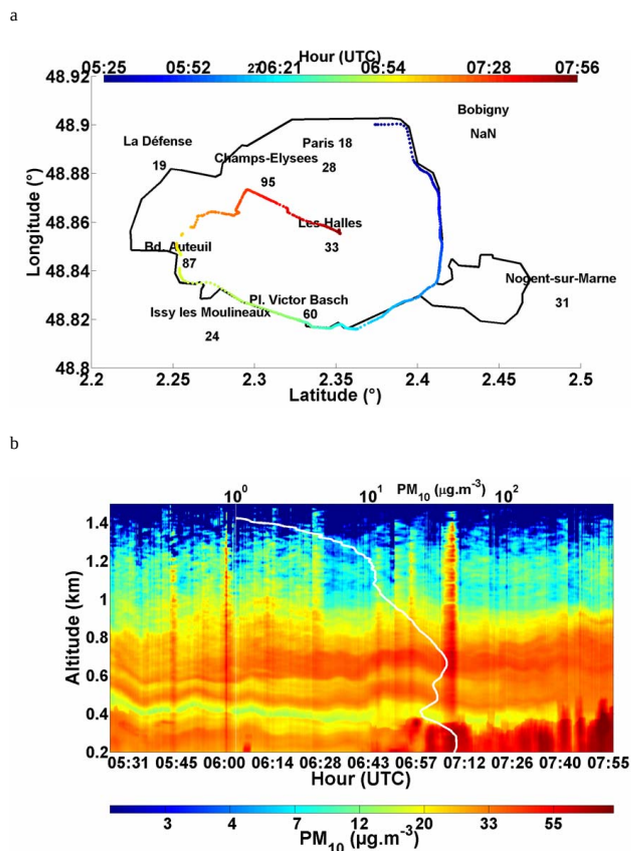


Fig. 7. Route followed by the small vehicle embarking the lidar on the 25 May 2005 the Paris Peripherique in traffic conditions (a), and corresponding 2-D colour-coded temporal evolution of lidar-derived PM₁₀ concentrations (b). Hourly averaged PM₁₀ values measured by AIRPARIF network stations are reported in panel (a). The mean PM₁₀ profile retrieved from lidar signals is shown in white in panel (b). The black line circling Paris on the map (a) is the geographic demarcation of Paris city.

BER value in the lowest layers (0.011 sr^{-1}) and using an iterative method converging when the total optical thickness retrieved by the lidar was equal to that measured by the sun-photometer.

Given that there were not any direct mass to optics relationship available for biomass-burning and dust aerosols, the determination of PM₁₀ concentrations in these layers has required the determination of k_2 , through the knowledge of their corresponding size distributions and complex refractive indices. Concerning biomass-burning aerosols, a typical size distribution for particles observed in the Sahelian region has been retrieved from AERONET measurements performed in 2006 and 2007 in Banizoumbou (Raut and Chazette, 2008b). The number size distribution presents two modes with corresponding modal radii of 0.07 and $0.7 \mu\text{m}$ with standard deviations of 1.5 and 1.9 respectively. The first mode represents 99% of the total number of particles. The corresponding refractive index has been retrieved by Raut and

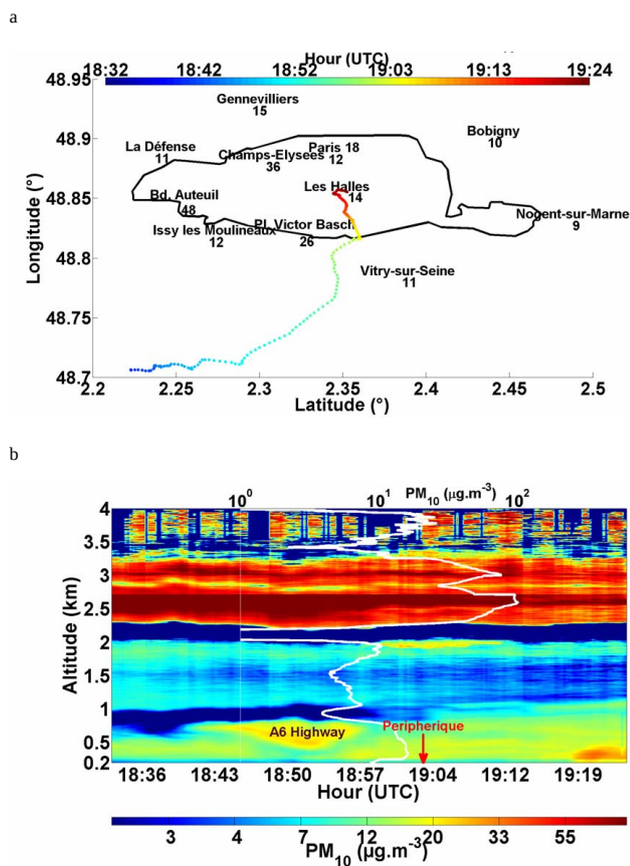


Fig. 8. Route followed by the small vehicle embarking the lidar on the 26 May 2005 from Palaiseau to Paris (a), and corresponding 2-D colour-coded temporal evolution of lidar-derived PM₁₀ concentrations (b). Hourly averaged PM₁₀ values measured by AIRPARIF network stations are reported in panel (a). The mean PM₁₀ profile retrieved from lidar signals is shown in white in panel (b). The black line circling Paris on the map (a) is the geographic demarcation of Paris city. The red arrow in (b) indicates where the lidar track intersected the Peripherique.

Chazette (2008b) from a synergy between remote sensing and in situ measurements: 1.53–0.047i. Concerning mineral dust particles transported from the Sahara to Western Europe, the size distribution is shifted towards the small radii during the transport. Thus, AERONET measurements performed between 1999 and 2008 over the stations of Paris, Palaiseau and Fontainebleau (48°24′21″ N, 2°42′07″ E) have been used to extract typical size distributions for dust episodes observed over the Paris area selected by an Angström exponent (between 400 and 670 nm) lower than 0.8 and a volume median radius larger than 1.6 μm . The resulting size distribution presents three modes. The fine mode containing 99.5% of the number of aerosols has a modal radius of 0.07 μm and a standard deviation of 1.5. The second mode is located at 0.63 μm with a geometric standard deviation of 1.4 and representing 0.4% of the number of aerosols. The third mode

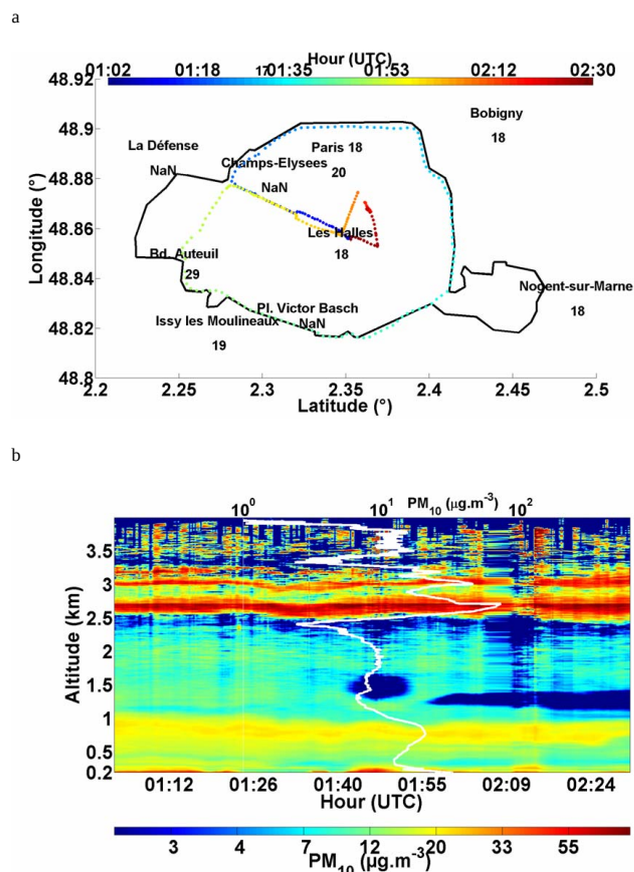


Fig. 9. Route followed by the small vehicle embarking the lidar on the 27 May 2005 over the Paris ring (a), and corresponding 2-D colour-coded temporal evolution of lidar-derived PM₁₀ concentrations (b). Hourly averaged PM₁₀ values measured by AIRPARIF network stations are reported in panel (a). The mean PM₁₀ profile retrieved from lidar signals is shown in white in panel (b). The black line circling Paris on the map (a) is the geographic demarcation of Paris city.

at 1.96 μm with a standard deviation of 1.7 only stands for 0.1% in number concentration but represents a larger volume contribution. The complex refractive index of dust particles has been taken equal to 1.52–0.007i from the results obtained over Niamey by Raut and Chazette (2008b).

The corresponding extinction specific cross-sections $s_{\text{ext},355}$ are lower than those obtained for anthropogenic aerosols (Table 1): 2.6 $\text{m}^2 \text{g}^{-1}$ for biomass-burning particles and 1.1 $\text{m}^2 \text{g}^{-1}$ for mineral dust. The result for biomass-burning aerosols is in the range of values reported in the literature (Table 2) for fresh combustion aerosols (Reid et al., 2005) and over the South Western Iberia Peninsula (Pereira et al., 2008). The value of $s_{\text{ext},355}$ retrieved for dust particles exported from Africa to Western Europe in this study is close to that found for aged dust over the Atlantic Ocean (1.1 $\text{m}^2 \text{g}^{-1}$; Clarke et al., 1996), for dust in the South Western Iberia Peninsula (0.97 $\text{m}^2 \text{g}^{-1}$; Pereira et al., 2008), and

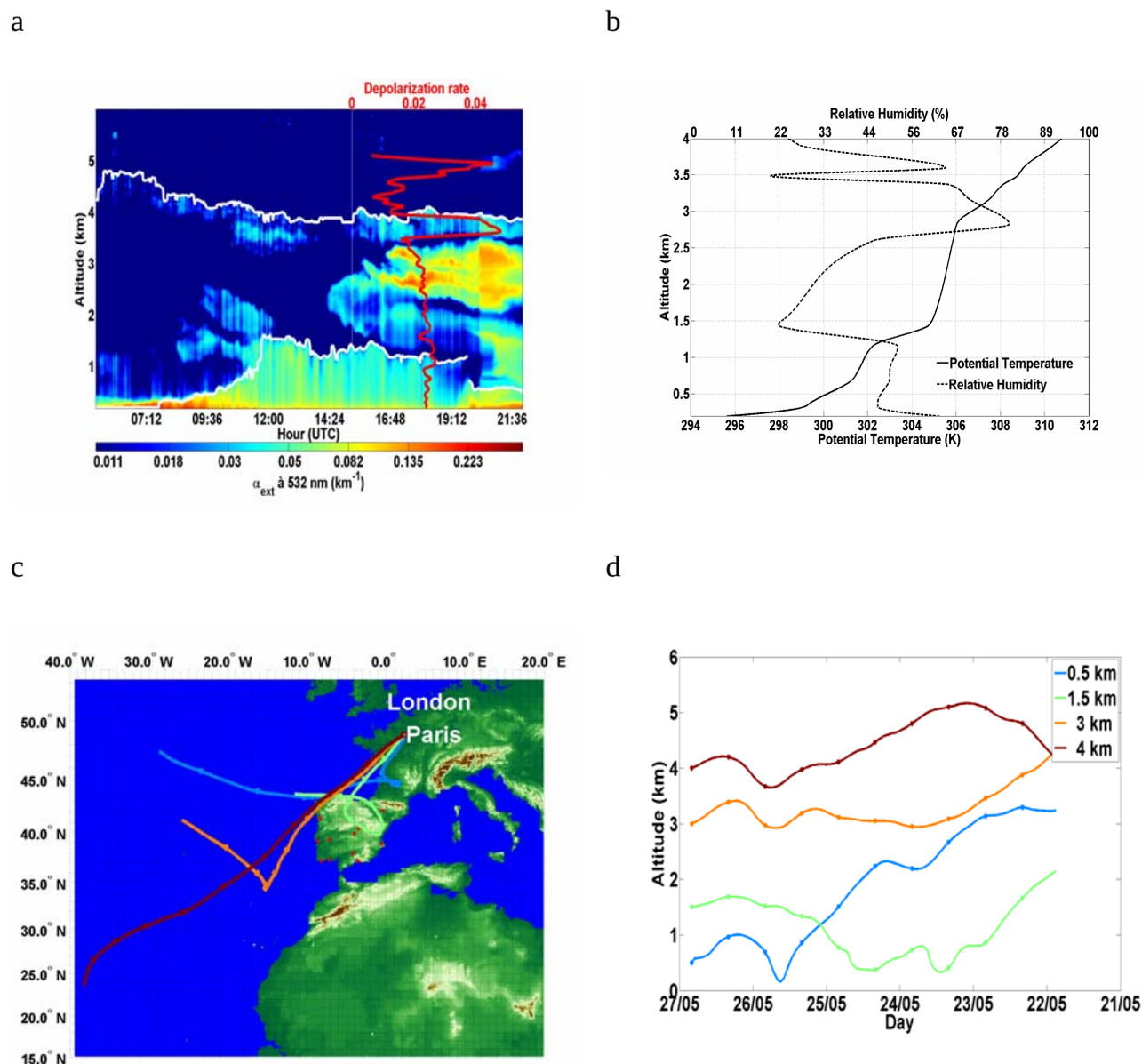


Fig. 10. Temporal evolution of lidar vertical profiles of extinction coefficient at 532 nm measured over Paris city hall on the 26 May 2005 by LESAA lidar (a). The associated vertical profile of the depolarization rate retrieved at 20:00 UTC is shown in red. The white lines represent the results of the detection of lidar-derived vertical structures: top of the dust layer and top of the boundary layer. The mean vertical profiles of RH and potential temperature are represented in (b). Back trajectories for 5-day periods ending over Paris at 20:00 UTC on 26 May at 0.5, 1.5, 3 and 4 km m.s.l. (courtesy of NOAA Air Resources Laboratory <http://www.arl.noaa.gov>) are reported in (c). The triangles give the 12-h spacing. The locations of the fires detected by MODIS products from 21 to 24 May are given by red dots over Spain and Portugal. The location of the air mass for each day against the altitude is given in (d).

for dust transported over China ($1.05 \pm 0.13 \text{ m}^2 \text{ g}^{-1}$; Alfaro et al., 2003). Notwithstanding, values are often found lower for dust (Table 2). This result indicates that aged dust particles have larger values of $s_{\text{ext},355}$. This is most likely because, in the case of aged dust aerosols, the large particles have settling thereby shifting the particle size distribution, resulting in a smaller scattering cross-section (as r^2) but also in a smaller

mass (as r^3), and therefore in a higher scattering efficiency (stronger $s_{\text{ext},355}$).

The conversion of aerosol extinction coefficients into PM₁₀ through the parameter k_2 requires the detection of the top and bottom altitudes of layers transporting biomass-burning and dust aerosols. This has been done thanks to the algorithm based on the strongest gradient in aerosol

concentrations, mentioned in Sect. 5.1. These altitudes also fit well to the variations observed in the mean profile of potential temperature drawn in Fig. 10b. The RH profile also presents significant variations along the vertical tropospheric column. RH is $\sim 50\%$ in the atmospheric boundary layer (up to 1.2 km) but decreases down to 22% in the residual layer (up to 2.5 km), then sharply increases up to 80% in the more humid biomass-burning layer (2.5–3.2 km). The highest layer (3.5–4 km) containing mineral dust is relatively dry (RH < 55%). It is possible that this layer is composed by a mixing of dust and biomass-burning particles, but backtrajectories alone are inconclusive.

The resulting temporal evolutions of PM₁₀ vertical profiles are shown in Fig. 8b during the Palaiseau-Paris transect on the 26 May in the evening and in Fig. 9b over the Paris Peripherique during the night from 26 to 27 May. Aerosol concentrations are greater on the 27 May ($\sim 33 \mu\text{g m}^{-3}$) due to the low inversion layer preventing vertical dilution than on the 26 May in the afternoon ($\sim 13 \mu\text{g m}^{-3}$). PM₁₀ in the residual layer are almost similar ($6\text{--}11 \mu\text{g m}^{-3}$), although slightly larger in nighttime conditions. This approach also enables us to follow the spatiotemporal evolution on the biomass-burning aerosols mass concentration transported over Paris area from the South Western Iberia Peninsula. This episode occurs on the 26 May in the afternoon (Table 3) with low aerosol mass concentrations ($\sim 9 \mu\text{g m}^{-3}$). In the evening, the aerosol loading is much more important ($\sim 41 \mu\text{g m}^{-3}$) and turns out to be the predominant source of pollutants in mass in the troposphere (Fig. 8b). During the night from 26 to 27 May (Fig. 9b), this plume appears more diluted ($\sim 28 \mu\text{g m}^{-3}$). Conversely, the dust layer stays very constant in terms of shape and aerosol loading during its transport over the Paris area (Table 3).

The knowledge of the vertical distribution of PM₁₀ in the various tropospheric layers is particularly crucial when surface concentrations are influenced by the subsidence of air masses from the elevated layers aloft the boundary layer. This phenomenon occurred on the 26 May at $\sim 20:00$ UTC. After the evening rush hour linked to the automobile traffic jam ($\sim 17:00\text{--}18:00$ UTC), PM₁₀ concentrations measured at the surface by a TEOM of Paris air quality network (AIRPARIF, Les Halles station) were maintained at a fairly constant level ($15\text{--}20 \mu\text{g m}^{-3}$). Between 19:00 and 20:00 UTC, PM₁₀ strongly increase by a factor 3 up to a level of $45 \mu\text{g m}^{-3}$. This sharp increase cannot be explained by the erosion of the boundary layer, which only decreases by a factor 1.8. A more plausible scenario is the subsidence of aerosols from the residual layers, followed by the incorporation of those particles in the boundary layer during the erosion of this latter. The lidar profiles on Fig. 10a tend to illustrate this likely hypothesis. An accurate knowledge of the vertical distribution of PM₁₀ would therefore be required in air quality models so as to more precisely follow the variations of aerosol concentrations at the surface.

6 Discussion on retrieved mass concentrations

We evaluated the concentrations derived from mobile lidar measurements on AIRPARIF air quality network stations. Figures 4a, 5a, 6a, 7a, 8a and 9a present the spatial distribution of the stations considered in this study together with PM₁₀ concentrations reported from TEOM measurements at these stations. AIRPARIF observations are given each hour and are provided in this work at the nearest hour of lidar profiles. Lidar signals produce a sequence of analyzed states, which are thereafter compared to hourly observations through a statistical parameter: the statistical measure to evaluate the results is the Root Mean Square Error (RMSE), expressed in $\mu\text{g m}^{-3}$, defined by the geometric average of the differences between lidar-derived PM₁₀ at 200 m above ground level and PM₁₀ observations performed by AIRPARIF air quality network stations at the surface. In a general manner, a strong variability is observed in the concentrations measured by the lidar in the lowest layers of PBL and nocturnal layers. The corresponding RMSE has been found to be $\sim 14 \mu\text{g m}^{-3}$, which is of the order of half the concentrations observed in the Paris area. We have to bear in mind that the natural temporal fluctuations of the PM₁₀ values in Paris are characterized by a standard deviation of $5 \mu\text{g m}^{-3}$ on a one-hour basis (retrieved from our continuous TEOM measurements). Besides, AIRPARIF stations are not exactly localized on the path used by the small vehicle embarking LAUVA lidar. The spatial variability of mass concentrations thus contributes to the global RMSE.

There is furthermore a possible discrepancy between the concentrations derived from the lidar at 200 m above ground level and concentrations reported by stations directly at the surface. Most of the ground sites, where continuous measurements are performed, are considered as “urban stations”, indicating that they are mostly representative of Paris background. Only three stations (Bd. Auteuil, Pl. Victor Basch and Champs-Élysées) turn out to be traffic stations. It is worth noting that lidar-derived mass concentrations generally overestimate PM₁₀ values reported on Paris background stations, but underestimate measurements obtained in traffic stations. This can be explain by the fact that lidar measurements are performed over Paris highways and in traffic conditions but, due to the overlap factor, only values above 200 m are given in this study. At this altitude, pollutants have had time to be slightly diluted along the vertical and their concentrations have therefore decreased. In regional chemistry-transport models, the assumption of a well-mixed lowest layer is generally made down to the surface, enabling a direct comparison of surface concentrations and PM₁₀ derived from remote sensing applications a few hundred meters above. This coarse hypothesis has been tested in this study from lidar measurements performed on the 12 July 2005 over Palaiseau under high aerosol loading: the optical thickness at 380 nm given by AERONET sunphotometer in Palaiseau was ~ 0.9 and the corresponding Angström exponent between

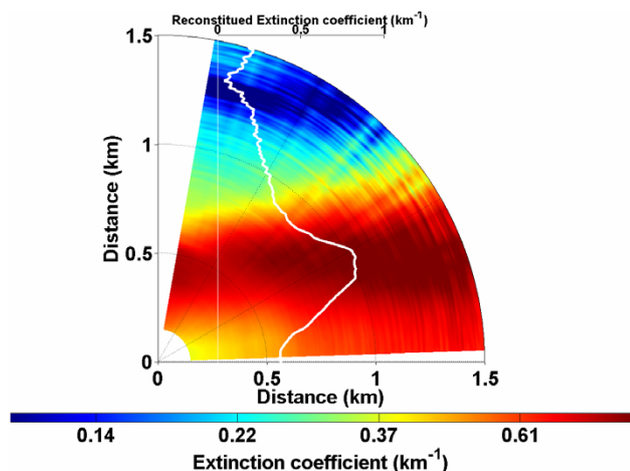


Fig. 11. Lidar scanning tomography between 0° and 90° performed on the 12 July 2005 over Palaiseau at ~21:00 UTC and represented in aerosol extinction coefficient at 355 nm. The white line gives the reconstituted extinction coefficient from multiangular profiles down to the surface.

440 and 670 nm was 1.6, indicating the presence of small aerosols. Multiangle lidar measurements were performed in the evening after 19:00 UTC in order to follow the erosion of the atmospheric boundary layer and the possible vertical particulate gradient close to the surface. Lidar measurements identified above 1.5 km a region deemed to be of low aerosol loading (so-called Rayleigh zone). The lidar was calibrated to estimated molecular returns in this region, which allowed the inversion of lidar signals with a BER of 0.011 sr⁻¹, except for profiles captured with a large zenithal angle (>68°): corresponding lidar signals indeed reach the aerosol-free zone for emitter-scatterers distances of the order of 4 km, i.e. at distances where signal-to-noise ratio is too weak to permit an inversion with a satisfying accuracy. As a consequence, lidar profiles with large zenithal angles have been inverted through a Klett method, choosing for reference aerosol extinction coefficients retrieved at 1 km altitude from lidar signals measured at lower zenithal angles. This approach assumes that the atmosphere is horizontally homogeneous over the explored angular range (Sicard et al., 2002). The reconstitution of the aerosol extinction coefficient has been done down to 5 m above the ground (zenithal angle of 88° and overlap factor at 150 m, Fig. 11).

This tomographic approach has enabled to follow the decreasing height of the aerosol layer against time: the top altitude of the layer was 1.2 km at 19:00 UTC, decreased down to 0.8 at 20:00 UTC and reached 0.6 at 21:00 UTC (Fig. 11). It has also highlighted a sharp increase in the extinction coefficient profile between the ground and the top of the aerosol layer. Therefore, this approach clearly underlines a strong decorrelation between observations at the surface and measurements performed at 200 m above ground level, as also

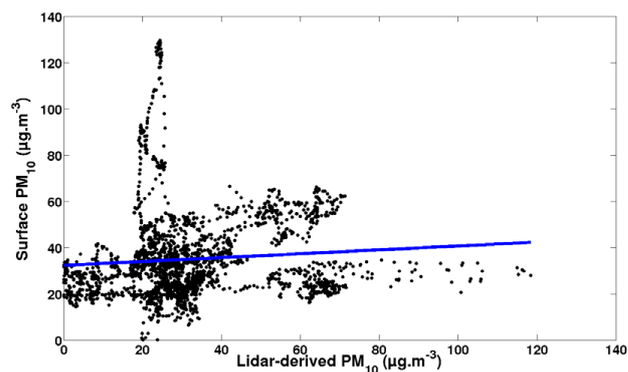


Fig. 12. Correlation between surface concentrations measurements (PM₁₀) and lidar-derived PM₁₀ at ~200 m during the LISAIR campaign when the lidar was vertically shooting on a stationary site in front of Paris City Hall. The blue line represents the result of the regression analysis with a low correlation coefficient $R \sim 0.08$.

highlighted by Chazette et al. (2005b) from airborne measurements performed over Paris area. The correlation between the ground-based PM₁₀ and the lidar-based PM₁₀ on a longer period has moreover been studied. This scatter plot uses all the data collected during the LISAIR campaign when the lidar was vertically shooting on a stationary site in front of Paris City Hall. Figure 12 highlights a lack of correlation ($R \sim 0.08$) between TEOM measurements and lidar-derived PM₁₀ at ~200 m (average on a window of 20 m). This result confirms the strong decorrelation between observations at the surface and measurements performed at 200 m above ground level that were suggested by the tomographic approach and airborne measurements (Chazette et al., 2005b) and explains the discrepancies observed between mobile lidar measurements and AIRPARIF stations observations (resulting RMSE of 14 μg m⁻³).

7 Conclusions

This study describes two approaches to convert vertical profiles of aerosol extinction coefficients retrieved from lidar measurements into mass concentrations in terms of PM₁₀ or PM_{2.5}. Simple linear relationships between the scattering coefficient at 700 nm, continuously measured by a nephelometer and corrected from the non-observed angles and RH, and PM₁₀, measured with a TEOM, have been investigated during various campaigns performed in the Paris area between 1999 and 2007. The comparison between the theoretical approach, based on the least mean squares method, and this empirical relationship provides excellent results within a discrepancy of 3%. Therefore, specific cross-sections at 355 nm are provided with a reasonable uncertainty for aerosols of urban and periurban origins (~12%). This confirms that both methods are equivalent and that the simple linear empirical

relationship is appropriate to retrieve PM from extinction coefficients retrieved by lidar profiles. In a rural location, the uncertainty was found larger (26%).

Once the various layers have been delimited through an algorithm sensitive to vertical heterogeneities derived from lidar profiles, an appropriate specific cross-section was attributed for each of them: $4.5 \text{ m}^2 \text{ g}^{-1}$ for urban aerosol, $5.9 \text{ m}^2 \text{ g}^{-1}$ in periurban conditions, $7.1 \text{ m}^2 \text{ g}^{-1}$ for rural particles. This study permits to assess the role of the Paris Peripherique in local particulate pollution and the horizontal gradient of pollution between Paris centre and its remote suburbs. In addition to the nocturnal inversion layer and the residual layers, observations performed in Paris also highlight elevated layers from a different origin biomass burning aerosols ($2.6 \text{ m}^2 \text{ g}^{-1}$) and mineral dust particles ($1.1 \text{ m}^2 \text{ g}^{-1}$). This approach enables to clearly follow the spatiotemporal evolution on the biomass-burning aerosols transported over Paris area from the South Western Iberia Peninsula, which can be the predominant source of pollutants in mass in the low troposphere. Conversely, the dust layer stays very constant in terms of shape and aerosol loading during its transport over the Paris area. The knowledge of the vertical distribution of aerosols is important in case of air masses subsidence from the elevated layers inside the boundary layer.

Multangular lidar measurements have highlighted a possible discrepancy between PM₁₀ derived from the lidar at 200 m above ground level and concentrations reported by stations directly at the surface. The resulting RMSE between lidar-derived PM₁₀ at 200 m above ground and surface network stations measurements was $\sim 14 \mu\text{g m}^{-3}$.

The respective agreements between lidar and nephelometer measurements and between nephelometer and TEOM measurements validate the approach. Hence, lidar observations could be used to validate air quality models in terms of particulate pollution. Vertically resolved measurements in the atmospheric column are indeed required to get reliable forecasts. This paper enables to envisage improved decision-support tools based on assimilation approaches or ensemble analyses. On the long run, an assimilation of vertical profiles of mass concentrations provided by a lidar network could be considered to improve the description of the vertical mixing processes in the atmospheric column. This could be done by a sequential assimilation using an ensemble Kalman filter.

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