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LOAC: a small aerosol optical counter/sizer for ground-based and balloon measurements of the size distribution and nature of atmospheric particles – Part 1: Principle of measurements and instrument evaluation

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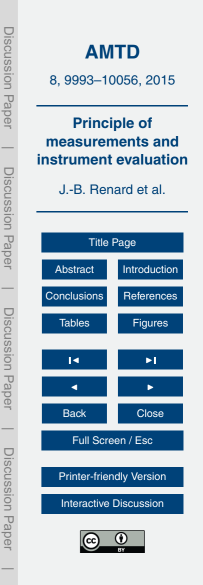
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Abstract

The study of aerosols in the troposphere and in the stratosphere is of major importance both for climate and air quality studies. Among the numerous instruments available, aerosol particles counters provide the size distribution in diameter range from

- few hundreds of nm to few tens of μm. Most of them are very sensitive to the nature of aerosols, and this can result in significant biases in the retrieved size distribution. We describe here a new versatile optical particle/sizer counter (OPC) named LOAC (Light Optical Aerosol Counter), which is light and compact enough to perform measurements not only at the surface but under all kinds of balloons in the troposphere
- and in the stratosphere. LOAC is an original OPC performing observations at two scattering angles. The first one is around 12°, and is almost insensitive to the nature of the particles; the second one is around 60° and is strongly sensitive to the refractive index of the particles. By combining measurement at the two angles, it is possible to retrieve accurately the size distribution and to estimate the nature of the dominant par-
- ticles (droplets, carbonaceous, salts and mineral particles) in several size classes. This topology is based on calibration charts obtained in the laboratory. Several campaigns of cross-comparison of LOAC with other particle counting instruments and remote sensing photometers have been conducted to validate both the size distribution derived by LOAC and the retrieved particle number density. The topology of the aerosols has been
- validated in well-defined conditions including urban pollution, desert dust episodes, fog, and cloud. Comparison with reference aerosol mass monitoring instruments also shows that the LOAC measurements can be successfully converted to mass concentrations. All these tests indicate that no bias is present in the LOAC measurements and in the corresponding data processing.



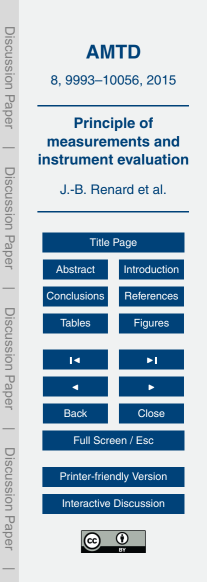
1 Introduction

The importance of measuring the concentration and size distribution of aerosols in the lower atmosphere has been highlighted by various studies. For instance, their presence in ambient air can have direct effects on human health (e.g. Zemp et al., 1999;

- ⁵ Brunekreef and Holgate, 2002), and their interaction with solar radiation and clouds are affecting regional and global climate (Ramanathan et al., 2001; Diner et al., 2004; Kanakidou et al., 2005; Quaas et al., 2008). When very high concentrations of ashes after volcanic eruptions are present at cruise altitude, they can affect air traffic (e.g. Chazette et al., 2012). In the middle atmosphere, aerosols play a significant role in
 ¹⁰ stratospheric chemistry through heterogeneous reactions with nitrogen and halogen
- species (e.g. Hanson et al., 1994, 1996), and they can affect climate through their role in the global radiative balance of the Earth (e.g. Hansen et al., 1992; Ammann et al., 2003). The concentration and size of the particles are highly variable due to the large variety of aerosol sources and properties, both of natural and man-made origin, and
- because of their relatively short residence time in the atmosphere. To understand and predict aerosol impacts, it is important to develop observation and monitoring systems allowing for their full characterization.

Instruments have been developed for routine measurements or for dedicated campaigns. Observations can be conducted from the ground, from unmanned aerial vehicles (UAV), from aircrafts, from balloons, and from satellites. To retrieve the physical properties of the aerosols, it is necessary to combine the information obtained with different instruments. In situ mass-spectrometers (Murphy et al., 2007) and aerosol collecting instruments (Brownlee, 1985; Blake and Kato, 1995; Allan et al., 2003; Bahreini et al., 2003; Ciucci et al., 2011) provide their composition. Optical instruments perform-

ing remote sensing measurements from the ground or from space with photometric, lidar, and extinction techniques (Shaw et al., 1973; Dubovik and King, 2000; Bitar et al., 2010; Winker et al., 2010; Salazar et al., 2013) provide indications on the size distribution and on the nature of the particles (liquid, carbon, minerals, ice, ...), generally



assuming a priori hypotheses in the retrieval process. Complementarily, in situ optical measurements with optical particle counters can provide more accurate information on the size distributions of the particles.

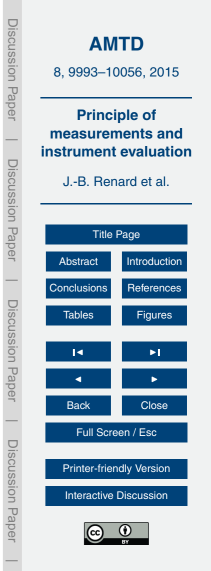
The present study deals with optical aerosol particles counters (OPCs). The corresponding measurement principle relies on the properties of light scattered by particles injected in an optical chamber and crossing a light beam (e.g. Grimm and Eatough, 2009). The measurements are usually conducted at "large" scattering angles, typically around 90° with collecting angle of a few tens of degrees. At such angles, the light scattered is depending both on the size of the particles and on their refractive index. Conventional counters are calibrated using latex and glass beads and are postcalibrated using Mie calculations (Mie 1908) for liquid aerosols (the refractive index of latex beads and liquid aerosols is well known, assuming no imaginary part of the index

i.e. non-absorbing aerosols). Some instruments can be also be post-calibrated for the observation of specific particles, as desert dust or urban pollutants, assuming a given value of their refractive index.

The refractive index dependence can be partially determined by performing measurements at different scattering angles, since the variation of the scattered intensity with scattering angles is strongly dependent on the refractive index of the particles (Volten et al., 2006; Francis et al., 2011). Thus, performing simultaneous measurements at different angles can provide an indication of the nature of the particles. Such an approach was used by Eidhammer et al. (2008) at angles of 40 and 74° mainly for

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- the identification of mineral particles, and by Gayet et al. (1997) with a ring of detectors covering the whole scattering angle range for the identification of cloud droplets and icy particles.
- ²⁵ Another approach was proposed by Renard et al. (2010a); in this case, measurements are conducted at small scattering angles, below 20°, where the light scattered is less sensitive to the refractive index of the particles. In this angular region, the scattered light is dominated by diffraction (which is not sensitive to the refractive index), at least for irregular grains as those found in the atmosphere. Such non-dependence of



the refractive index was confirmed by measurements conducted at a scattering angle around 15° for different types of irregular grains (Lurton et al., 2014). In this case, the light scattered is dependent only on the size of the particles, allowing a better determination of the corresponding size distribution. However, the main problem of measurement at small angles is stread light contamination. Thus a real time correction of the

5 ment at small angles is stray-light contamination. Thus a real-time correction of this signal offset due to the stray-light, which can vary with time, must be developed.

Aerosol particles counters are often used on the ground; some of them are used in the free atmosphere on-board aircraft or large balloons during dedicated campaigns, for example for the studies of desert dust events or volcanic aerosols (Bukowiecki et al.,

- ¹⁰ 2011; Jégou et al., 2013; Ryder et al., 2013) or for stratospheric studies (Rosen, 1964; Ovarlez and Ovarlez, 1995; Deshler et al., 2003; Renard et al., 2008, 2010b). We propose here a new optical particle counter concept, called LOAC (Light Optical Aerosol Counter) that is light and compact enough to perform measurements on the ground and under all kinds of balloons in the troposphere and in the stratosphere, including
- ¹⁵ meteorological balloons. LOAC uses a new approach combining measurements at two scattering angles. The first one is around 12°, an angle for which scattering is weakly sensitive to the imaginary part of the refractive index nature of the aerosols, allowing the retrieval of the particle size distribution. The second one is around 60° where the light scattered is strongly sensitive to the refractive index of the particles, and thus can
- ²⁰ be used to evaluate their topology (liquid droplet are transparent, minerals are semitransparent, and carbonaceous particles are strongly absorbing).

In this first paper, we will present the principle of measurements and calibration, and cross-comparison exercises with different instruments that detect atmospheric aerosols. In the companion paper, we illustrate first scientific results from airborne ob-

²⁵ servations on-board balloons and unmanned aircraft (Renard et al., 2015).

| Discussion Paper | | AMTD 8, 9993–10056, 2015 Principle of measurements and instrument evaluation JB. Renard et al. | | |
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2 Principle of measurements

2.1 Instrument concept

LOAC is a modular instrument, for which some parts can be changed depending on the measurements conditions. For measurements under balloon or on the ground in low

- wind conditions, the aerosols are collected by a metal profiled inlet designed to optimize the sampling conditions when oriented in the wind direction. The particles are drawn up to the optical chamber, through an isostatic tube and to the injector that focusses the flux inside the laser beam. LOAC uses a small vane-type pump (having a life-time of 3 weeks in continuous operation) working at ~ 2 L min⁻¹. The pump is connected to
- the exit of the optical chamber by a flexible plastic tube. The optical chamber is open, thus the pressure is the same as outside. In-flight tests under sounding balloons have shown that the rotation speed of the pump is not affected by pressure variations.

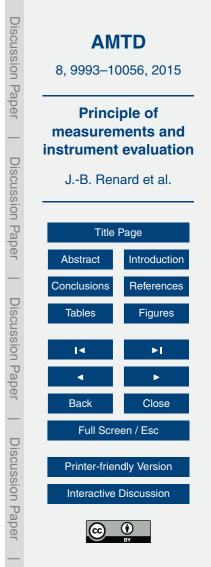
For measurements in windy and rainy conditions, the inlet can be replaced by a total suspended particulate or TSP inlet rejecting rain droplets and particles greater than

¹⁵ 100 μ m. For long-duration measurements, the small pump can be replaced by a robust pump; to maintain the aerosol detection efficiency, the pump flux must be in the range 1.3–2.7 L min⁻¹.

To minimize its weight, the optical chamber is in plastic Delrin[®]. The weight, including the pump, is of 300 g. The electric consumption is of 340 mA under 8 V (which corresponds to a power of 3 W). The optical chamber and the pump can fit in a rectangle box of about 20 cm × 10 cm × 5 cm.

LOAC is mainly designed for the detection of irregular grains, as those present in the ambient air (Fig. 1, from Institute of Physics of the University of Sao Paulo, Brazil). It uses a statistical approach for the size and concentration retrievals, as done for

the laboratory PROGRA2 instruments dedicated to the study of optical properties of irregular levitating grains (Renard et al., 2002). Because of their shape, their orientation and their rotation in the air flow, the scattering properties of an individual grain vary with time at a given scattering angle (this variation could be more than a factor 2, as shown



during laboratory tests by photodiodes and imagery measurements with PROGRA2). This must be taken into account for the calibration and data analysis. Thus, we propose here a calibration approach that can differ from the one used for other optical counters.

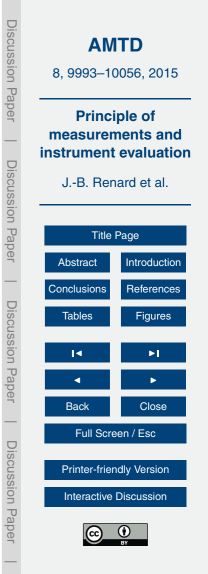
The sampled air crosses a laser beam of 25 mW working at the wavelength 650 nm.

- The homogeneity of the beam is of ±20%. The scattered light is recorded by two photodiodes at scattering angles respectively in the 12–16° (hereafter called the 12° channel) and 55–65° (hereafter called the 60° channel), as shown on Fig. 2. Instead of using lenses to collect the light, the photons travel directly to the photodiodes through pipes, providing fields of view with a few degrees. The collecting area of the photodiodes is
 larger than the diameter of the pipes. This system prevents optical misalignment prob-
- In a ger than the diameter of the pipes. This system prevents optical misalignment problems in case of vibrations and strong temperature variations like those encountered during atmospheric balloon flights. Such a concept of scattering measurements without collecting lenses was tested and validated by Daugeron et al. (2007).

The electronic sampling is at 40 kHz and the transit time of particles inside the laser beam is equal or lower than 500 µs. As said before, stray light contamination is high at small scattering angles and needs real-time correction. The stray light correction method presented in Renard et al. (2010a) was applied to the LOAC measurements. The stray light acts as a flux continuum, which can slightly vary over time due to changes in the temperature and pressure conditions and possible dust contamination in the optical chamber. The light scattered by the particles is superimposed on this

²⁰ In the optical chamber. The light scattered by the particles is superimposed on this continuum, which can be assumed as a continuous base-line over a short time interval. This baseline is determined before and after the intensity pulse produced by the particles that cross the laser beam.

The maximum of the intensity pulse is obtained after subtracting the stray-light contamination. Figure 3 presents an example of real ambient air measurements of the time evolution of the flux scattered by a 5 μm particle and by few submicronic particles. The pulse is slightly asymmetric, because the particles decelerate when crossing the optical chamber. This deceleration occurs because the diameter of the optical chamber is larger than the diameter of the inlet; thus the particles encounter pressure relaxation.

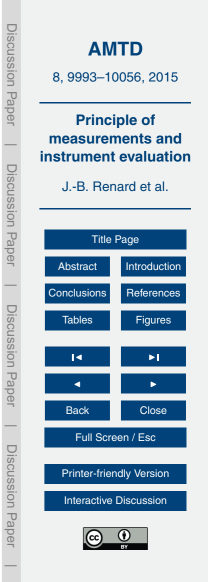


Some secondary flux maxima may be present in the pulse and can be attributed to the rotation of irregular shaped particles in the air flow. The search for a new intensity peak is inhibited until the flux decreases to a given threshold, represented in Fig. 3 by the red line. This procedure prevents multiple counting of the same particle (of irregular shape) that exhibits secondary flux maxima.

This pulse analysis is performed by the on the on-board electronics, which provides every 10s the concentrations detected by the 2 channels. Until now, an additional computer has been necessary to record and analyse the data.

2.2 Calibration

- The calibration of an optical counter is not an easy task, especially for the detection of irregular particles (Whitby and Vomela, 1967; Gebhart, 1991; Hering and McMurry, 1991; Belosi et al., 2013). A first presentation of the calibration procedure for measurements at small scattering angles using a LOAC optical chamber can be found in Lurton et al. (2014).
- ¹⁵ Only the 12° channel, which is insensitive to the refractive index of the particles, must be calibrated. The 60° channel will be used as a comparison to the 12° channel measurements to determine the typology of the aerosol, as explained in the Sect. 2.4. Monodisperse latex beads, which are perfect transparent spheres, have been used for diameter calibration in the 0.2–4.8 µm range; glass beads have been used at 5 µm
- (see Figs. 2 and 3 of the Lurton et al., 2014 paper for the LOAC response to monodisperse beads). In fact, Mie calculations show that the scattered flux encounters strong oscillations linked to small changes both in diameter and in scattering angle. Conventional aerosol counters use large field of view, typically a few tens of degrees, to average these oscillations. On the opposite, the LOAC 12 and 60° measurement chan-
- nels have a field of view only of few degrees and use no lens. The detected flux at the 12° channel is then very sensitive to the position of the individual bead inside the laser beam, and thus to its scattering angle. Taking into account this constraint, we

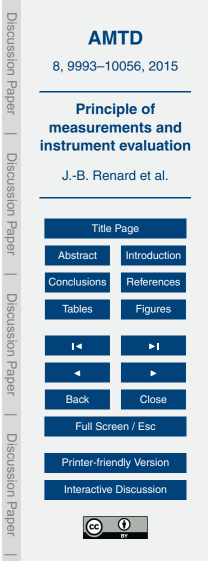


considered here only the highest flux scattered by each size class of monodisperse beads.

The electronic noise is lower than 20 mV at ambient temperature and lower than 10 mV at low air temperatures. Statistically speaking, the noise is divided by the root mean square of the number of identical measurements (here the number of events detected in a given size class). To reach a 1 mV accuracy in case of 20 mV noise, which is necessary to be able to discriminate the smaller size classes and to establish

- which is necessary to be able to discriminate the smaller size classes and to establish accurately the size distribution, at least 20×20 (= 400) particles must be detected for each size class.
- ¹⁰ During laboratory calibration, it is easy to reach such concentration levels using monodisperse beads. During real measurements inside the atmosphere, we must ensure that such particle concentrations are indeed present for all size classes below 1 μ m. The LOAC has an integration time of 10 s, with a pumping flow of about 2 L min⁻¹. Even in low polluted ambient air at ground ("background conditions"), all counting in-
- ¹⁵ struments have shown in the past that concentrations are greater than 1 particle cm⁻³ for size classes smaller than 0.5 µm (having a 0.1 µm width), which corresponds to 2000/6 = more than 300 particles during the 10 s LOAC integration time. For particles in the 0.5–1 µm size classes (having a width of 0.1 or 0.2 µm), concentrations are greater than 0.1 particle cm⁻³, giving more than 30 particles per size class.
- For all the cross-comparison exercise presented below, the measurements were integrated form 2 to 15 min. For the 2 min integration time, the number of particles given above must be multiplied by 12, giving are at least 3000 for the 3 first size classes and 300 for the other ones. For a 15 min integration time, these numbers must be multiply again by 7.5. Thus, the LOAC class identification can be conducted with the expected accuracy in the ambient air. Obviously, in case of polluted air, all these values could be also 2 to 3 orders of magnitude higher (1000 particles cm⁻³ between 0.2 and 0.3 um is
 - also 2 to 3 orders of magnitude higher (1000 particles cm^{-3} between 0.2 and 0.3 μ m is often encountered).

In case of very low particle concentrations, as those that can be encountered during flights in the stratosphere with typically less than 1 particle cm⁻³ greater than 0.2 μ m,



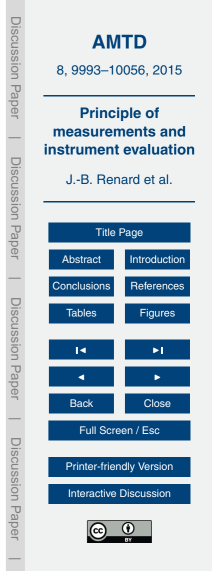
the size attribution will be less accurate. Thus, the retrieved size distributions and the time-evolution of the concentration will be more scattered and need to be averaged in altitude.

- For the calibration in the 7–45 μ m size range, different natures of irregular grains have been used: carbon particles, dust sand of various types, ashes and salts (see for example Fig. 4 of the Lurton et al., 2014 paper). The size selection was obtained using sieves. For diameters at ~ 90 μ m, calibrated silicon carbide grains were used, the size being characterised by the provider. The diameter presented here corresponds to an equivalent (or optical) diameter, which can differ significantly from the aerodynamic diameter or from the electric mobility diameter used by non-optical instruments
- ¹⁰ namic diameter or from the electric mobility diameter used by non-optical instruments for ambient air measurements. Several tens of grains are necessary to ensure a mean random orientation, to be able to derive a mean equivalent diameter. The relation between flux and size was derived by considering the diameter where the concentration distribution is at its maximum The measurements with different nature of grains con-¹⁵ firms that no significant dependence with the particle type exists for the variation of the
- scattered flux with their diameter, as presented in Fig. 8 of Renard et al. (2010a) and Fig. 5 of Lurton et al. (2014).

Taking into account the laser departure from homogeneity, the electronic noise, and the statistical approach, the uncertainties in size calibration is of $\pm 0.025 \,\mu$ m for particles

²⁰ smaller than 0.6 µm, 5% for particles in the 0.7–2 µm range, and of 10% for particles greater than 2 µm. Figure 4 presents the calibration curve for the 12° channel, where the scattered flux is given in mV, which corresponds to the photodiode output voltage (updated from Lurton et al., 2014). The offset due to the electronic dark current and high frequency noise of the detector were added to the calculations; thus the curve asymptotically decreases with decreasing size to this offset value.

The calibration with the latex beads captures well the large-amplitude Mie oscillations up to $5\,\mu\text{m}$ in diameter, calculated by integrating the scattered fluxes over the whole LOAC field of view (12–16°). In particular, the amplitude of the oscillations at 1, 2 and $5\,\mu\text{m}$ are well reproduced. For the larger sizes, calibrated with irregular grains, the



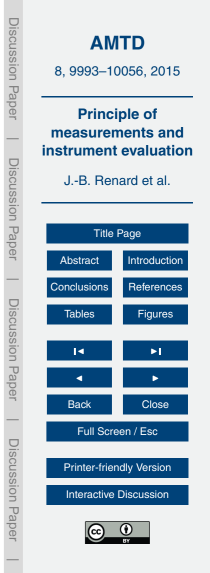
evolution of the scattered flux with size is lower than the one expected from the Mie calculation. This difference was attributed to the natural roughness of the particulates and the small LOAC field of view, as shown by theoretical calculations (Lurton et al., 2014). It was found that a relatively small roughness parameter caused the scattered intensities to collapse on a lower saturation limit corresponding solely to the diffraction part of the scattered light.

Solid particles found in ambient air are not perfectly spherical and have some irregularity on their surface, even for the sub-micrometre (sub-µm) sizes (e.g. Xiong and Friedlander, 2001; McDonald and Biswas, 2004). Thus, the Mie oscillations will disappear and the scattered flux will roughly cross the middle of the oscillations amplitude (except perhaps for some iced crystals encountered in some clouds). A good illustration of the light scattering properties by irregular grains can be found in Weiss-Wrana (1983).

The flux evolution with diameter for the particles larger than a few μ m can be fitted ¹⁵ using a power law. As the lower saturation limit of Lurton et al. (2014) is likely to exist even for very slightly rough particulates, this power law can be reasonably used to estimate the sizes of any sort of real particulates found in the atmosphere. The best fit is obtained using a power law in $D^{-1.0}$ (adding an offset of 17 mV) where *D* is the particle diameter, as shown in Fig. 4.

²⁰ Such an approach is validated by performing measurements with particles in urban air, as shown in Fig. 5. Measurements were integrated over 15 min (90 acquisitions of 10 s). LOAC captures well the continuous decrease of concentration with increasing size, as previously established by various kinds of instruments, such as electrostatic low pressure impactors (e.g. Shi et al., 1999). A calibration error would produce oscil-

²⁵ lations in the size distribution, correlated to the Mie oscillations. On the other hand, we would expect that the LOAC calibration could be inappropriate for droplets, which are supposed to be perfectly spherical. In fact, such a phenomenon was not observed during laboratory tests conducted for droplets only, nor during measurements inside fog and clouds. We can expect that droplets are slightly distorted when entering the optical



chamber due to changes in the air flow speed between the aerosols injector and the optical chamber.

Thus, the LOAC detection size range is between 0.2 and ~ 100 μm. LOAC, with its present calibration procedure, is operated to the detection of irregular grains and droplets, but not to perfect spherical solid grains, such as latex or metal beads for which uncertainties arise from the smoothing of Mie oscillations by the calibration curve (in this case, the total concentration is correct but the size attribution can be erroneous).

Overall, a total of 19 size classes are defined for diameters between 0.2 and 100 μ m (Table 1). The upper limit can be lower, however, depending on the sampling collection cut-off of the inlet. The size classes are chosen as a good compromise between the instrument sensitivity and the expected size distribution of ambient air aerosols.

2.3 Concentration measurements

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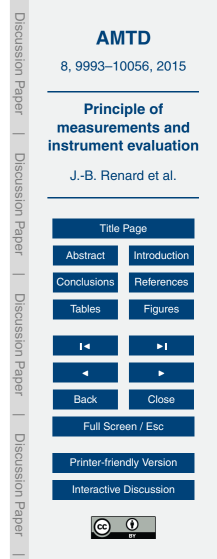
Counting is conducted while the particles cross the laser beam one by one, and are classified in size classes corresponding to the scattered flux. The measurements are integrated during 10s and are converted to number densities or particles cm^{-3} . The

detectors of the two channels (12 and 60°) work asynchronously.

This discrete detection works well for large particles greater than $2 \mu m$, with uncertainty in size attribution of 10%. For smaller particles, the size determination is within the calibration errors bars ($\pm 0.025 \mu m$ for particles smaller than 0.6 μm , 5% in the 0.7– 1 μm range) if more than 400 particles are detected for each size classes.

The optical and electronic response of the system has been modelled by a numerical Monte-Carlo method, taking into account the shape of the laser beam, the speed of the particles inside the laser beam and the instrument noise. To ensure a good statistical approach, 10⁴ particles were randomly injected for each size class. The ratio

 $_{25}$ of the number of detected particles over the number of injected particles provides the detection efficiency. For the smallest particles, only the brightest part of the peak of the pulse of the scattered flux is observable and the apparent transit time in the laser beam is a few tens of μs (in Figs. 3 and 4 small peaks are present). Thus, some par-

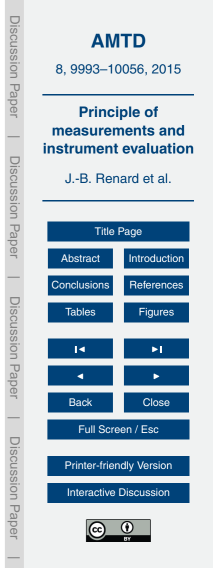


ticles cannot be detected. The detection efficiency increases as the diameter of the particles increases, and is reaching 100% for particles larger than 1 μm . The concentrations of submicron size particles are then corrected using these detection efficiency coefficients.

- ⁵ For particles larger than 1 μ m, the observed transit time in the laser beam is at its maximum (~ 500 μ s) and the expected maximum concentration is of about 20 particles cm⁻³. In fact, even higher concentrations can be determined using a statistical approach when several particles cross the laser beam almost simultaneously. The higher are the concentration, the lower is the probability that the scattered intensity peaks decreases below the threshold to start a new counting. In this extreme case, the
- real concentrations are inversely proportional to the detected concentrations. Another Monte-Carlo numerical modelling was conducted to establish the relationship between the number of particles > 1 μ m detected and the number of particles injected in the laser beam (Fig. 6). In the simulations, particles were randomly injected (in time), with
- ¹⁵ concentrations increasing from 0 to 500 particles cm⁻³ by step of 1 particle cm⁻³. The response is almost linear up to 10 particles cm³, reaching a kind of saturation value at around 15 particles cm⁻³, and decreases for larger concentrations. It is obvious that such a corrective procedure must be used only in dense aerosol media (more than 10 particles greater than 1 μ m cm⁻³), as fog or clouds, i.e. in conditions which must be confirmed by independent measurements. At present, this procedure is applied only
- when large droplets are detected by LOAC using the topology procedure presented below. In this case, up to 200 large particles cm⁻³ can be detected.

For the smaller particles, the signal of the scattered flux is close or inside the noise. The photodiodes cannot detect the whole transit of the particles inside the laser beam,

²⁵ but just the brighter part of the peak. Thus, the effective acquisition time can be reduced down to 35 μ s instead of around 500 μ s for the largest particles. Then, a greater number of particles can be detected. Taking into account also the detection efficiency for the smaller particles, up to 3000 particles cm⁻³ can be (statistically) detected.



For the LOAC integration time of 10 s, the counting uncertainty can be derived from the Poisson counting statistics. This uncertainty, defined as the relative standard deviation, is 60 % for aerosol concentrations of 10^{-2} cm⁻³, 20 % for 10^{-1} cm⁻³, and 6 % for concentrations higher than 1 cm⁻³. Such uncertainties can be reduced by averaging the concentration measurements for each size classes or by increasing the integration time. Nevertheless, such calculation does not take into account the real instrumental

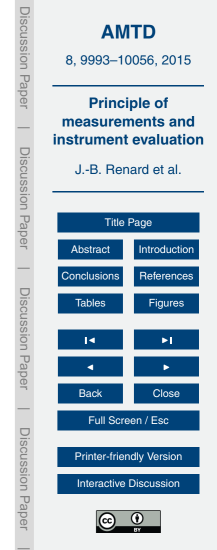
uncertainties dominated by the electronic noise and the inlet sampling efficiency. LOAC is designed to be used in various atmospheric conditions. The temperature can dramatically change, in particular during balloon flights up to the middle strato-

- sphere. The electronics offset can change with time because of the sensitivity of the electronic components to atmospheric temperature variations. The instrument performs a check of its noise level after 10 min of measurements. If the noise differ by more than 50 % from the previous check, an electronic re-calibration is automatically performed to estimate the offset variation and to adjust the calibration. A processing software is applied after the experiment to check the offset time-evolution during the
- 10 min periods and to then correct the raw measurements.

2.4 Aerosols typology

The scattered flux recorded at 60° is very sensitive to the refractive index of the particles and thus to their nature (as said before this phenomenon appears at scattering angles greater than ~ 20°). The more absorbing they are, the lower the recorded fluxes. Thus, we use the "60° channel" as a diagnostic for the effect of the refractive index on the scattered fluxes. This channel uses the same flux threshold (in mV) as the 12° channel, in order to perform a direct comparison of the counting detected by two channels. For a given size class and for a given particle concentration recorded in the 12° channel, the concentration detected by the 60° channel decreases when the imaginary

²⁵ channel, the concentration detected by the 60° channel decreases when the imaginary part of the refractive index increases. This increase of the imaginary part leads to an underestimation of the real size of the particles, and thus produces a diameter bias in the size distribution (diameter vs. concentration) for the 60° channel with respect to the



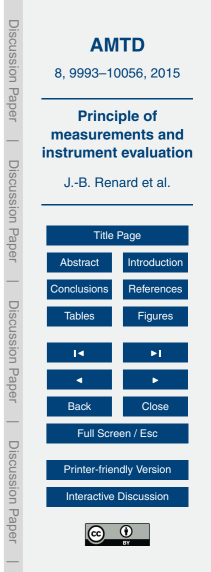
12° channel. An example of the procedure used to determine this effect is presented in Fig. 7, where the size distributions of the two channels are presented. For a given particle size of the 12° channel (noted D1), we consider the concentration value of the 60° channel. Then we search for the same concentration value on the 12° channel (a linear interpolation is used if needed). The corresponding diameter is then determined (D2). Finally, we define a so-called "speciation index" as the ratio D2/D1. The more absorbing the particles are, the higher this ratio. This procedure is conducted for each size class.

This procedure works well for irregular particles, but not for solid symmetrical particles; in this later case, the Mie oscillation will produce strong fluctuations in the evolution of the speciation index with size (we have indeed observed this effect inside some cirrus clouds). Also, this procedure must be used only for a large enough number of detected particles per size class, because of the irregular shape of the particles. Laboratory tests have shown that about 20 particles in a size class are sufficient to be able

to indicate the aerosol topology. In its nominal operating mode, LOAC provides the speciation index every 1 min. For the analysis of continuous ground-based measurements presented below, we have conducted the topology detection with an integration time of 15 min (assuming that the aerosols are stationary).

Different types of particles have been tested in the laboratory to assess the amplitude of the speciation index throughout the measurement size range: organic carbon, black carbon, desert dust or sand from different origins (excluding black sand), volcanic ashes, plaster, salt (NaCl), water droplets, droplets of mixture of water and sulphuric acid. They can be classified in 4 families: carbonaceous particles, minerals, salts and liquid droplets. Figure 8 presents the curves obtained in laboratory for the various sam-

²⁵ ples. Then, "speciation zones" charts (speciation index vs. real diameter) are defined by the minimum and maximum speciation index values reached by each family, taking into account the measurement uncertainties. Among solid particles, carbonaceous particles produce the higher speciation index and salt the lower, mineral particles being in between. Detailed analysis has shown that most of the carbon particles are in the



lower part of the carbon speciation zone while some strongly absorbing particles, perhaps black carbon having fractal shape, are in the middle and upper part of the carbon speciation zone. For all solid particles, the global trend is a decrease of the speciation index with increasing size. On the contrary, the liquid droplets speciation index exhibits an increase with increasing diameter.

The case presented in Fig. 7 has $D1 = 0.35 \,\mu\text{m}$ and $D2 = 0.51 \,\mu\text{m}$, leading to a ratio of 1.46, which is in the carbon speciation zone.

The speciation indices obtained from LOAC observations in the atmosphere are compared to these reference charts obtained in the laboratory. The position of the data ¹⁰ points in the various speciation zones provides the main topology of the particles. In principle, this procedure can be conducted for each size class. In fact, due to the statistical dispersion of the results, it is better to consider several consecutive size classes to better conduct the identification. This is in particular necessary for the identification of droplets, whose speciation zone crosses all the speciation zones of the solid particles.

It is obvious that the identification of the typology of the particles works well in case of a homogenous medium, when the speciation indices are not scattered through the various speciation zones.

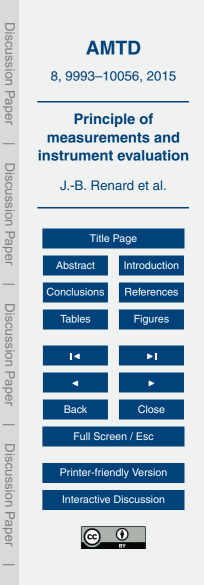
At present, the speciation zones are established for particles expected to be found in the troposphere and stratosphere, but it is an evolving data base. Additional laboratory measurements can be conducted to retrieve the speciation zones for specific particles in case of measurements in new specific environments.

2.5 Reproducibility

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The instrument is industrially produced by Environnement-SA (http://www. environnement-sa.com); more than 100 copies were produced up to mid-2015. Tests have been conducted for the different parts of the instrument: diode, pump, photodiode and electronics, to assess their reproducibility and thus to establish the measurements uncertainty.



The variation of the laser flux from one copy to another is less than 5%, which has no significant effect on the flux scattered by the particles. The variability of the pump flow was less than $\pm 0.2 L$ from one pump to another. The flow can be controlled manually by a flow-metre before a balloon flight or during ground based measurements;

- ⁵ it can used in the post-processing for the concentration retrieval. The stability of the pump flow over one hour is of about ±5%, which induces a ±5% concentration uncertainty. The pump was tested at low temperature and low pressure in balloon flights in the stratosphere and no obvious instability nor loss of performance has been detected. Finally, optical tests have been conducted with a rotating wheel inserted between the laser and the photodiodes. The wheel had pinholes of different diameters to produce
- ¹⁰ laser and the photodiodes. The wheel had pinholes of different diameters to produce different flux levels. Overall, the photodiodes and the electronic contribute to a reproducibility uncertainty of less than 5 %.

Taking into account all these uncertainties, we can expect a total reproducibility of about ± 15 %, assuming no systematic bias. To confirm this estimate, tests have been conducted with 8 LOAC in a "pollution test room" at LPC2E laboratory (Orléans, France). Various types of solid particles, following mainly a power law size distribution, have been injected in the chamber. A standard deviation of ± 15 % (1 σ) has been obtained between the different instruments for particles smaller than 10 µm from mea-

surements of these 8 LOACs using the two channels. The standard deviation increases ²⁰ up to ± 30 % for particles larger than 10 µm. This increase is due to the low concentrations of large particles, as resulting from the Poisson counting statistics.

2.6 Inlet sampling efficiency

LOAC will be used in different conditions, mainly on the ground and under balloons. Depending on the chosen inlet and the relative speed between the inlet and the wind,

the isokinetic sampling is respected or not, and the efficiency of collecting the largest particles can change.

On the ground, a total suspended particulate (TSP) inlet can be used, ensuring an efficiency close to 100 % for collecting all the particles up to a few tens of $\mu m.$ For some



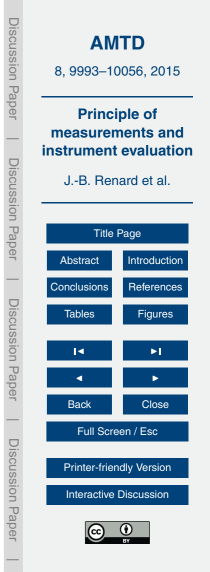
specific studies where very large particles dominate, as measurements inside fog or clouds, or because of mechanical constraint if a TSP inlet cannot be mounted, the particles can be collected by a tube having a bevelled metal inlet and oriented downwards. In this case, the largest particles are generally under-sampled and a corrective coefficient must be applied, taking into account the direction and the speed of the wind.

For measurements under balloons floating at constant altitude, the relative speed between ambient air and the inlet is close to zero. The sampling efficiency assessed using the Agarwal and Liu (1980) criterion for an upward-facing inlet shows that the sampling is unbiased for particle with diameter below $20 \,\mu m$.

¹⁰ The sampling line used during the flights is composed of a thin wall metallic probe and antistatic tubing. The thin wall aerosol probe has an inlet diameter equal to 5.4 mm and is connected to a tube of about 20 cm long and 6.7 mm internal diameter. The sampling line is connected vertically to the LOAC. Nevertheless, due to the tube stiffness, the line can be inclined with a maximum sampling angle of 30° from vertical.

- The sampling efficiency of the line was assessed using modelling calculations in order to account for changes in atmospheric pressure, temperature and possible changes of the probe orientation during these flights. For that purpose, the values of pressure and temperature as a function of altitude are taken from the international standard atmosphere. Sampling efficiency calculations have been made by considering a balloon
- ²⁰ ascending velocity of 5 m s⁻¹, a LOAC sampling flow rate equal to 1.7 L min⁻¹ and two angles of the sampling line from the vertical (0 and 30°). According to these parameters, the inlet aspiration velocity of the probe is equal to 1.24 m s⁻¹ (sub-isokinetic) and the flow is laminar in the tubing for all altitudes.

The mechanisms considered to calculate the sampling efficiency are the inlet efficiency of the probe in isoaxial and isokinetic sampling conditions (Belyaev and Levin, 1974; Hangal and Willeke, 1990) and particle losses in the tubing due to gravitational settling when the line is not perfectly vertical (Heyder and Gehbart, 1977). Calculations have been conducted for particles with diameter ranging from 0 to 20 µm, and from the ground to an altitude of 30 km. Figure 9 presents the sampling efficiency for a 0°



deviation (isoaxial) and for a 30° deviation of the sampling line with respect to the vertical. Data are plotted according to the particle aerodynamic diameter which describes particle settling and inertia phenomenon.

In isoaxial conditions, results show for all altitudes an increase of sampling efficiency with the particle diameter, up to a factor > 3 for the largest particles. In this case, there is no particle deposition in the sampling line and the sampling is dominated by subisokinetic conditions (apparent wind velocity higher than inlet probe velocity). A sampling efficiency higher than unity is explained by the particle inertial effect resulting from the divergence of the flow field at the inlet of the probe. The increase in sampling efficiency with altitude is due to changes in air viscosity and gas mean free path with temperature and pressure.

When the tube is inclined by 30° from the vertical, the sampling efficiency is between 1 and 2. The sampling efficiency is lower than for the 0° isoaxial conditions. Firstly, the sub-isokinetic effect is reduced by the orientation of the tube, and secondly, deposition can occur in the tubing due to particle settling.

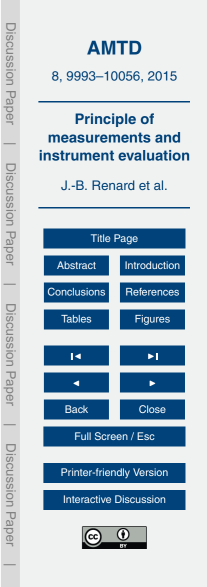
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Since the tube always has a deviation of about 30° during the balloon flights, we consider only the results at 30° from the vertical. The over-sampling effect is negligible for particles smaller than 5 μ m up to the lower stratosphere and for particles smaller than 2 μ m in the middle stratosphere. Thus, this effect will just affect the retrieved concentrations of the largest particles.

The results of these theoretical calculations are not yet fully validated by an experimental approach with LOAC itself. Thus, all balloon measurements in the stratosphere will not be corrected from this aerodynamic effect. It could be taken into account in future work involving large particles, for example when converting concentrations to

²⁵ extinction by comparison with remote sensing instruments, or to estimate the real concentration of the interplanetary dust in the middle atmosphere.



3 Cross-comparison with other instruments

Various cross-comparisons have been conducted in ambient air at ground and during balloon flights for concentrations and topology identification. LOAC concentrations are compared to other commercial particle counter instruments and photometer mea-

⁵ surements, although there is no absolute reference, many of them are using different technical approaches and calibration procedures. The LOAC topologies are validated during well-identified atmospheric events of liquid and solid particles. Finally, the LOAC particle concentrations are converted to mass concentrations to be compared to commercial microbalance mass instruments used as reference instruments in air quality monitoring. Table 2 summarizes the conditions of measurements.

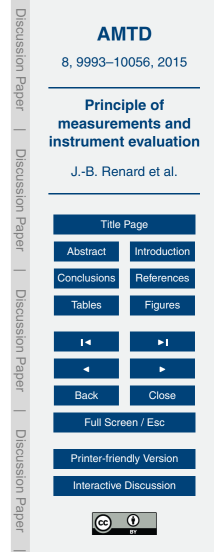
The LOAC was used under different conditions. When deployed underneath meteorological and tropospheric balloons, the data are transmitted in real time by telemetry. For deployments under large stratospheric balloons, the data are stored on-board using a specific module. On the OAG tethered balloon, the data are sent to the ground

¹⁵ using a Wi-Fi link and are stored on a computer. An autonomous version for automatic ground-based applications is now also available, including an on-board computer to record the data.

3.1 Concentrations and size distribution

Continuous measurements have been conducted in ambient air at the SIRTA observatory (Site Instrumental de Recherche par Télédétection Atmosphérique, http: //sirta.ipsl.fr/) at Palaiseau, South of Paris, France (48.713° N, 2.208° E), during the ParisFog campaign, http://parisfog.sirta.fr/, from November 2012 to April 2013. During this period, the total concentrations of aerosols have been monitored by a WELAS aerosols counter, a Fog Monitor (counter for large droplets) and a SMPS (Scanning Mo-

bility Particle Sizer) in common size range domains with LOAC. SMPS measurements are based on the electric mobility diameter of the particles, while the other instruments determine optical diameters. These two diameters could differ depending on the nature



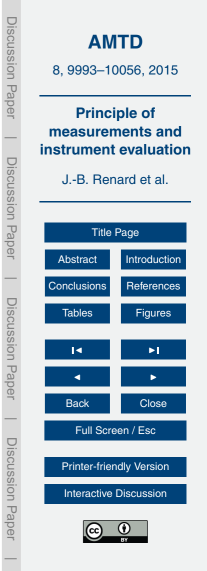
of the particles; thus the direct comparison of the measurements (even in the same air mass) could be sometimes biased.

Figure 10 presents the cross-comparison of the instruments in January 2013. Most of the measurements were conducted in background aerosol conditions, although some
small fog events were detected and can be identified by concentration peaks in the Fog Monitor measurements. Roughly speaking, the order of magnitude of the concentrations is similar, although some significant discrepancies are present. To investigate their possible origin, the size distribution obtained in different conditions of aerosol content can be compared. Figure 11 (upper panel) presents an example where the agreement in total concentration during background aerosol conditions is very good between LOAC and SMPS. On the other hand, the shape of the size distribution of the WELAS instrument is unusual with a decrease of the sub-µm aerosol concentrations

with decreasing size (the opposite trend is expected for background aerosol conditions). This could be due to calibration problem of the latter instrument; Indeed, Heim

- et al. (2008) report a strong decrease in counting efficiency with decreasing particle diameter for submicron particles for the WELAS-2100 OPC. Finally, as expected, the Fog Monitor measurements are not useable (no fog at this time and only noise is present). Figure 11 (lower panel) presents the same cross-comparison in case of significant disagreements between all the instruments. LOAC seems to underestimate the con-
- ²⁰ centrations of the smallest particles, but this can be due to the difficulty of reconciling the retrieved diameters for the different measurement techniques in case of different types of particles.

Strong fog events were observed in November 2012. LOAC, WELAS and Fog Monitor are in very good agreement during these events (Fig. 12). This result validates the ²⁵ correction procedure applied to the LOAC measurements in case of dense medium of liquid particles. Between fog events, LOAC and WELAS were sometimes in disagreement. This was due to the difference in the concentration values obtained by the two instruments for the particles smaller than ~ $0.5 \,\mu$ m, which may be attributed to the WELAS undercounting. Figure 13 presents the size distribution at the beginning



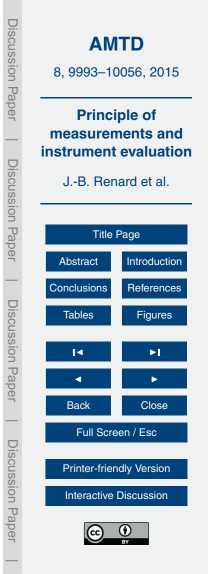
of a fog event, with the typical enhancement around a diameter of $10 \,\mu\text{m}$ (e.g. Singh et al., 2011), and at the end of the event. Both LOAC and WELAS found a bimodal size distribution although the WELAS shows a doubtful sharp decrease in concentration for the smallest sizes., but disagree for the size of the second mode at ~ $10 \,\mu\text{m}$. On the

opposite, LOAC and Fog Monitor were in good agreement for the position of the second mode, although the population of the first size class of Fog Monitor was obviously underestimated. Finally, for the largest sizes, LOAC concentrations are in-between those of the WELAS and Fog Monitor.

A ground-based measurement session was conducted from Minorca (Spain) during the ChArMEx campaign (Chemistry Aerosol Mediterranean Experiment, https: //charmex.lsce.ipsl.fr/) in parallel with measurements of an HHPC-6 aerosol counter in the period 12 June–2 July 2013. The orders of magnitude for the different size classes were in good agreement. In particular, both instruments captured an aerosol enhancement of large solid particles between 18 and 21 June 2013, as shown in Fig. 14 for the size distribution.

The last cross-comparison exercise was conducted during an ambient air campaign at SIRTA observatory, site 5 near Gif-sur-Yvette, South of Paris, France (48.709° N, 2.149° E), in the beginning of 2014. LOAC performed measurements from 3 to 13 February 2014 in parallel with a SMPS, a Grimm aerosol counter and a HHPC-6
²⁰ aerosol counter. Due to the sampling conditions that vary from one instrument to another (direct sampling, TSP inlet, dryer, direct or curved tubes), the analysis is limited to the smallest particles (diameter < 1 μm) which are almost insensitive to the sampling techniques. Figure 15 presents the temporal cross-comparison for 4 size-classes: 0.2–0.3, 0.3–0.5, 0.5–0.7 and 0.7–1 μm. In fact, the size classes of the 4 instruments are
²⁵ not always the same, thus the closest ones have been considered for the comparison.

Globally, all the instruments give similar concentrations for all size classes, the better agreement being for the 0.5–0.7 μ m diameter range. Some discrepancies appear for some time periods, but several reasons can explain them. First, the SMPS instrument determines the electric mobility diameter that can depend on the nature of the aerosols,



whereas the other instruments determine optical diameters. SMPS measurements could lead to some uncertainties in size determination, and thus in concentrations, when compared to other kinds of instruments for irregular particles (e.g. Gulijk et al., 2003). This could explain why LOAC has missed some concentration peaks detected

- ⁵ by SMPS. Secondly, the particles size distribution of sub-μm particles exhibits a strong decreasing when the diameter increases. Thus the uncertainty in the size calibration of a few hundredth of μm could induce concentration differences of at least a factor 2. This is presented in Fig. 15 for the 0.7–1 μm comparison with the Grimm instrument for which both 0.65–1 and 0.8–1 μm concentrations are plotted. Finally, the Grimm and HHPC 6 instruments are constitue to the nature of the particles, and changes in the
- ¹⁰ HHPC-6 instruments are sensitive to the nature of the particles, and changes in the type of aerosol (for example mineral or carbon particles) could partially affect their size determination.

An indirect evaluation of the LOAC size calibration has been conducted during the ChArMEx campaign on the Balearic island of Minorca, Spain. A total of 9 flights of

- LOAC have been performed under a meteorological sounding balloon launched from Sant Lluís airfield (39.865° N, 4.254° E) in the 15–19 June 2013 period during a desert dust transport event. The aerosol concentration has been integrated for all size classes from the ground to the highest altitude reached by the balloon, i.e. an altitude of about 30 km, to be compared to ground-based remote sensing measurements provided by
- ²⁰ the AERONET photometer network (http://aeronet.gsfc.nasa.gov/) station of Cap d'En Font (39.826° N, 4.208° E), which performed measurements close the trajectory of the LOAC balloon measurements. AERONET provides the vertically integrated volume concentration of aerosols (in dV/dln(r), where r is the radius of the particles) in the 0.13–30 µm radius range (Dubovik et al., 2000).
- The LOAC integrated concentrations are converted to volume concentrations by using the mean volumetric diameter D_v calculated for each size class by the formula:

$$D_{\rm v} = 0.5 \times \left[\left(D_{\rm min}^3 + D_{\rm max}^3 \right) \right]^{(1/3)},$$

where D_{min} and D_{max} are the lower and upper diameter of a given size class, respectively. With such a formula, the mean volumetric diameter is at about 60% of the size class width instead of 50% for the mean geometric diameter. For each size class, the volume of the particles is calculated assuming sphericity. To be consistent with the AERONET data, the LOAC results are presented in radius instead of diameter.

Figure 16 presents two examples of comparison between LOAC and AERONET size distributions for two different amounts of sand particles in the troposphere (the contribution of the stratospheric particles is negligible). The bi-modal distribution is typical for a desert dust or sand plume event. The two instruments are in excellent agreement, both in size distribution and volume concentrations It is worth noting that the volume concentrations are proportional to the cube of the size of the particles, an error in the LOAC calibration would lead to strong discrepancies, which is not the case.

The cross-comparison measurements presented above have been conducted for different air temperature, including day-night cycles and seasonal temperature variations.

No effect of the temperature on the accuracy of the retrieved concentrations has been pointed out. These results confirm that the LOAC real-time noise-checking process works well.

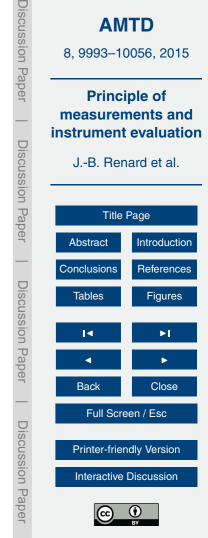
All these cross-comparison exercises have shown that the LOAC measurements are consistent with those of the other instruments considered here, accounting for the errors and the limitation of the various techniques. This confirms that no systematic bias are present in the LOAC calibration and in the data analysis when LOAC is used at ground.

3.2 Tropospheric vertical distribution

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Cross-comparison exercises have been also conducted for balloon-borne LOAC mea-²⁵ surements.

LOAC has performed tropospheric flights during the ChArMEx campaign from Minorca Island in time-coincidence with the WALI aerosols lidar measurements (Chazette et al., 2014) at a few tens of km apart. One LOAC flight was conducted under meteo-



rological balloon on 16 June 2014; two LOAC flights were conducted on 19 June 2013 at the same time, the first one being under a meteorological balloon and the second being under a pressurized tropospheric balloon (see the companion paper for more information of the balloons and the gondolas, Renard et al., 2015). To be compared to lidar extinction data at 350 nm, the LOAC data were converted to extinction using Mie scattering theory, assuming spherical sand particles having an refractive index of n = 1.6 + 0.05i (such index is an average of values available in the literature). Uncertainties of the refractive index values are included in the errors bars calculations of the retrieved LOAC extinctions. Figure 17 presents the tropospheric vertical profile 10 of LOAC and WALI lidar extinctions. Taking into account the instrumental errors bars, LOAC and WALI have captured the same main vertical structures and the extinction

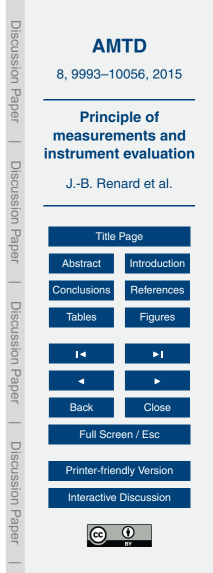
values are in average in good agreement in the lower troposphere.

3.3 Topology of the particles

The speciation zones, obtained from laboratory measurements, must be validated in real atmospheric conditions.

Urban ambient air measurements are proper for the detection of carbon particles (black and organic carbon), especially during well-identified pollution events. Permanent LOAC measurements have been conducted at "Observatoire Atmosphérique Generali" (OAG) in the South-West of Paris since May 2013 (48.841° N, 2.274° E). This

- ²⁰ observatory is a recreational tethered balloon operated in a public park; the LOAC measurements nominal maximum altitude is 120 m but some flights could be conducted up to an altitude of 270 m. The measurements can be sorted out between measurements with the balloon at ground level and measurements during flight. Figure 18 presents an example of light absorbing particles (probably carbonaceous ones) detected at the
- $_{25}$ OAG on 29 December 2013 around 07:30 UT. In this example, the speciation index curve is well inside the carbon speciation zone in the whole size range up to $\sim 10\,\mu m$. In addition to sounding balloons mentioned above, measurements under drifting balloons launched from Sant Lluís on Minorca Island were also conducted during sev-



eral well-identified desert dust events above the Mediterranean sea during the summer ChArMEx campaign. Figure 19 presents an example on 17 June 2013, around 14:30 UT (approximate balloon position: 41.9° N, 4.1° E) at an altitude of 2050 m under a low altitude pressurized drifting balloon. The speciation curve is well inside the mineral dust zone, showing that LOAC has indeed detected the desert dust event.

Measurements in the marine atmospheric boundary layer were also conducted with a low altitude balloon on 22 July 2013 drifting in an altitude range of 250–400 m, launched from the French Levant Island off Hyères on the Mediterranean French coast (43.021° N, 6.461° E). Figure 20 presents the measurements at 21:25 UT (approx. balloon position: 43.0° N, 6.55° E, alt. ~ 275 m), and the topology is mainly in the "salt zone", as expected for a measurement close to the sea surface.

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Droplet topology was validated in fog events during the ParisFog campaign; but also during cloud measurements conducted in May 2013 at the Puy de Dôme observatory (45.772° N, 2.964° E, alt. 1465 m). Figure 21 presents an example of measurements in-

- ¹⁵ side a cloud on 15 May 2013 at 10:30 UT. Globally, the topology identification is inside the droplets zone, which indicates that all of the particles were indeed liquid. In addition, measurements were conducted inside haze or thin cloud at an altitude of 1.2 km during a flight under a meteorological balloon launched from Reykjavik, Iceland(64.127° N, 21.904° W), on 7 November 2013 at 12:30 UT in the frame of the VOLTAIRE-LOAC
- ²⁰ campaign for the study of the stratospheric aerosol trend. The presence of the droplets was confirmed by the on-board humidity sensor, with a hygrometry of 90 %. The topology in Fig. 22 is well inside the droplets zone.

Finally, most of the measurements under meteorological balloons in the middle atmosphere show that (pure) liquid water and sulphuric acid droplets largely present

²⁵ in the stratosphere are close to the lower part of the droplets zone, and sometimes slightly below. Vertical profiles of LOAC concentration and topology measurements are presented in the second part (Renard et al., 2015).

These examples show that the topology determination works well in case of homogeneous aerosol media. Nevertheless, there are two limitations of this process. First,



the analysis of measurements conducted in heterogeneous media could be difficult or even inaccurate, in particular when different size modes are present. In this case, the speciation curve exhibits unusual oscillations that match none of the speciation zones. Secondly, some high porosity aerosols can exhibits high values for the speciation index, even if they are not black (as fluffy silica). Then, the topology determination is providing most of the time the nature of the particles, but one has to be cautious in the analysis when the speciation curves are non-typical.

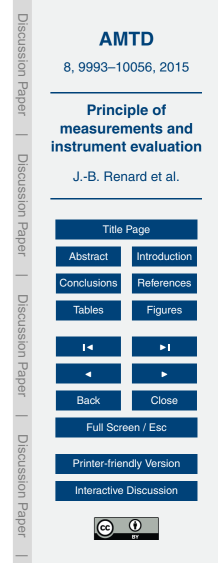
3.4 Mass concentrations

Our final test to evaluate both the calibration of LOAC and the retrieval of concentrations in all size classes (but especially large particles) is to convert the number size distribution measurements to mass concentrations and to compare the results to reference mass measurements. This is the most sensitive test to evaluate LOAC, since mass concentrations are proportional to the cubic diameter of the particles (and to their density). The topology helps to determine the type of aerosols, from which a density can be deduced. The density determination is necessary for the conversion of number concentrations (in cm⁻³) to mass concentrations (in µg m⁻³).

Measurements were conducted first in indoor air (in the "pollution room" at the LPC2E laboratory) in autumn 2013, by injecting in the air of the room different kinds of carbonaceous and mineral particles (smaller than $20\,\mu m$) in various concentrations

to produce a large range of mass concentrations. The reference mass measurements were achieved with a calibrated TEOM microbalance. An air flow system was used (when needed) to prevent sedimentation of the particles in the room. Also, some measurements have been conducted without injecting particles, to detect only the smallest particles present in the ambient air, in particular during the night without convection in the room.

The volume concentration is calculated for each size class, using the mean volumetric diameter, assuming spherical particles, and is multiplied by the corresponding concentrations. The mass concentration is obtained by multiplying these results by the



particle density. The mass densities were determined for each size class by identifying the typology of the particles though their speciation index. The mass densities chosen here are:

 $-2.2 \,\mathrm{g\,cm^{-3}}$ for salt; a value corresponding to NaCl particles;

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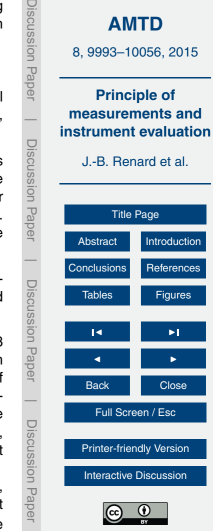
- 2.2 g cm⁻³ for mineral particles; this value is a compromise for common mineral particles present in ambient air: compact sand (2.1 g cm⁻³), quartz (2.7 g cm⁻³), limestone (2.5 g cm⁻³) and silicon (2.3 g cm⁻³);

1.4 g cm⁻³ for carbonaceous particles. This value was derived after detailed tests during the comparison between LOAC and microbalance measurements in the laboratory. It lies well inside the range of values proposed in the literature for such particles (e.g. Chen et al., 2010; Virtanen et al., 2006; Spencer et al., 2007). Sensitivity tests have shown that a 10% variation of this value will not induce strong changes in the results presented below.

A value of 0.0 g cm⁻³ was used for water droplets, for comparing LOAC measurements to those of the TEOM instrument, which tends to evaporate condensed water.

The duration of the sessions was from several hours to several days. Figure 23 presents the mass measurements for particles smaller than $20 \,\mu$ m, averaged on 24 h for the two instruments. The variability of the concentrations is related to the amount of particles injected into the room. The lowest values correspond to measurements without injection. In this case, LOAC indicates that only particles smaller than $2 \,\mu$ m were present in the air. The LOAC and TEOM measurements are in very good agreements, with a correlation of 0.97. The correlation curve has the slope of 0.98, with an offset at the origin of 2.2 μ g m⁻³, and a mean error of 4.8 μ g m⁻³.

²⁵ Sessions of ambient air measurements were conducted in Paris and in its suburbs, to test the retrieval of $PM_{2.5}$ and PM_{10} mass concentrations, with pumps working at $2.7 Lmin^{-1}$. The first location of measurements is at the "Observatoire Atmosphérique



Generali (OAG)" in Paris (latitude 48.8417° N, longitude 2.2736° E). The LOAC measurements were conducted using a TSP inlet. The second location is at SIRTA observatory at Palaiseau (48.7180° N, 2.2075° E) during the ParisFog campaign. The LOAC measurements were conducted with the metal inlet directed towards the ground.

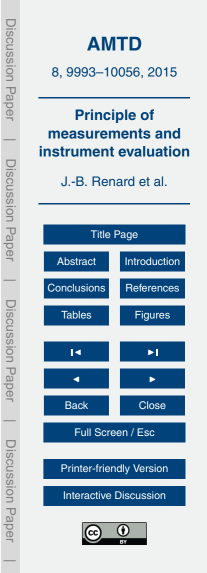
⁵ The OAG and SIRTA measurements considered here were conducted in the periods September 2013–April 2014 and September 2013–December 2013, respectively. The PM_{2.5} and PM₁₀ LOAC mass concentrations were retrieved by combining the results for particles smaller than 3 μm an smaller than 10 μm, respectively, taking into account the sampling efficiency of the PM_{2.5} and PM₁₀ inlets currently used by the air quality
 ¹⁰ networks (cut-off at 2.5 μm for PM_{2.5} inlet and cut-off at 10 μm for PM₁₀ inlet).

Reference mass concentrations data of urban ambient air in the Paris region are provided by the Airparif network (http://www.airparif.asso.fr/), operating TEOM microbalance instruments. Unfortunately, there is no Airparif station very close to the OAG site nor to the SIRTA site at the time of the measurements. Therefore, we decided to use

¹⁵ data recorded at 3 stations that have environmental conditions close to those at OAG and SIRTA: Paris Centre (latitude 48. 8528° N, longitude 2.3600° E), Vitry-Sur-Seine (latitude 48.7820° N, longitude 2.3992° E) in the south-eastern suburb area of Paris, and "Rural South" at Bois-Herpin (latitude 48.3725° N, longitude 2.2258° E) in the South of the Paris region; the last station provides background conditions measurements.

Figures 24 and 25 present the comparison of PM_{2.5} and PM₁₀ concentrations, for the 2013 and 2014 period, respectively. The LOAC measurements are most of the time between the background and the urban conditions, the small discrepancies with the reference mass concentrations are probably due to a difference in the wind direction and to the regional-scale transport of the particles. It is worth noting that LOAC did capture well the 10–15 December 2013 and the 11–14 March 2014 pollution peaks.

These measurement sessions have been conducted with different kinds of pumps and of inlet systems. The agreement with reference mass concentration measurements is very good. This confirms that no obvious bias is present in LOAC observations for the sizes of particles considered here ($\sim 0.2-20\,\mu m$), and that the topology procedure is



providing useful information to convert the LOAC concentrations for the 19 size classes to mass concentrations.

4 Conclusions

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- LOAC is a modular optical particle counter/sizer, whose pump and air inlet can be changed, depending on the conditions of measurements. Extensive tests performed in different atmospheric conditions have shown that LOAC can be used to retrieve the size distribution of irregular-shaped or liquid aerosols with a satisfactory accuracy at ground level and from all kinds of balloons, if the total concentrations of particles greater than 0.2 μm is more than 1 cm⁻³.
- ¹⁰ LOAC can also provide an estimate of the typology of the particles in case of a relatively homogeneous media. Finally, LOAC can be used for monitoring the mass concentration of PM_{2.5} and PM₁₀ (and of course of larger particles) in ambient air with reasonable accuracy. The companion paper will present and discuss the first scientific results from balloons and an unmanned aerial vehicle.
- Acknowledgement. The LOAC project was funded by the French National Research Agency's ANR ECOTECH. The instrument and the gondola are built by Environnement-SA and MeteoModem companies. The balloons flights of the ChArMEx campaigns were funded and performed by the French Space Agency CNES. The Icelandic flights were conducted by the Iceland Meteorological Office. The various copies of LOAC used in the campaigns were funded with the support of CNES. ADEME, and INSUL CNES in the framework of the MISTRAL S Programme.
- ²⁰ support of CNES, ADEME, and INSU-CNRS in the framework of the MISTRALS Programme, and of the French VOLTAIRE Labex (Laboratoire d'Excellence ANR-10-LABX-100-01). The QAIDOMUS laboratory tests were funded by the French Ministry of Industry. Some calibration tests were conducted at the Aerolab Company.

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This work is in memory of Jean-Luc Mineau.



References

Agarwal, J. and Liu, B. Y. H.: A criterion for accurate sampling in calm air, Am. Ind. Hyg. Assoc. J., 41, 191–197, 1980.

Allan, J. D., Jimenez, J. L., Williams, P. I., Alfarra, M. R., Bower, K. N., Jayne, J. T., Coe, H.,

- and Worsnop, D. R.: Quantitative sampling using an Aerodyne aerosol mass spectrometer 1. Techniques of data interpretation and error analysis, J. Geophys. Res., 108, 4090, doi:10.1029/2002JD002358, 2003.
 - Ammann, C. M., Meehl, G. A., Washington, W. M., and Zender, C. S.: A monthly and latitudinally varying volcanic forcing dataset in simulations of 20th century climate, Geophys. Res. Lett., 30, 1657, doi:10.1029/2003GL016875, 2003.
- Bahreini, R., Jimenez, J. L., Wang, J., Flagan, R. C., Seinfeld, J. H., Jayne, J. T., and Worsnop, D. R.: Aircraft-based aerosol size and composition measurements during ACE-Asia using an Aerodyne aerosol mass spectrometer, J. Geophys. Res., 108, 8645, doi:10.1029/2002JD003226, 2003.
- ¹⁵ Belosi, F., Santachiara, G., and Prodi, F.: Performance evaluation of four commercial optical particle counters, Atmos. Clim. Sci., 3, 41–46, doi:10.4236/acs.2013.31006, 2013.
 - Belyaev, S. P. and Levin, L. M.: Techniques for collection of representative aerosol samples, J. Aerosol Sci., 5, 325–338, 1974.

Bitar, L., Duck, T. J., Kristiansen, N. I., Stohl, A., and Beauchamp, S.: Lidar observations of

20 Kasatochi volcano aerosols in the troposphere and stratosphere, J. Geophys. Res., 115, D00L13, doi:10.1029/2009JD013650, 2010.

Blake, D. F. and Kato, K.: Latitudinal distribution of black carbon soot in the upper troposphere and the lower stratosphere, J. Geophys. Res., 100, 7195–7202, 1995.

Brownlee, D.: Cosmic dust: collection and research, Annu. Rev. Earth Pl. Sc., 13, 147–173, 1985.

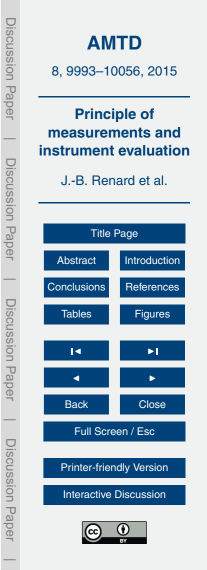
25

10

Brunekreef, B. and Holgate, S. T.: Air pollution and health, Lancet, 360, 1233–1242, 2002.

Bukowiecki, N., Zieger, P., Weingartner, E., Jurányi, Z., Gysel, M., Neininger, B., Schneider, B., Hueglin, C., Ulrich, A., Wichser, A., Henne, S., Brunner, D., Kaegi, R., Schwikowski, M., Tobler, L., Wienhold, F. G., Engel, I., Buchmann, B., Peter, T., and Baltensperger, U.: Ground-

³⁰ based and airborne in-situ measurements of the Eyjafjallajökull volcanic aerosol plume in Switzerland in spring 2010, Atmos. Chem. Phys., 11, 10011–10030, doi:10.5194/acp-11-10011-2011, 2011.



Chazette, P., Bocquet, M., Royer, P., Winiarek, V., Raut, J.-C., Labazuy, P., Lardier, M., and Cariou, J.-P.: Eyjafjallajökull ash concentrations derived from both lidar and modelling, J. Geophys. Res., 117 D00U14, doi:10.1029/2011JD015755, 2012.

Chazette, P., Marnas, F., and Totems, J.: The mobile Water vapor Aerosol Raman Lldar and

- its implication in the framework of the HyMeX and ChArMEx programs: application to a dust 5 transport process, Atmos. Meas. Tech., 7, 1629–1647, doi:10.5194/amt-7-1629-2014, 2014. Chen, S.-C., Tsai, C.-J., Chou, C. C.-K., Roam, G.-D., Cheng, S.-S., and Wang, Y.-N.: Ultrafine particles at three different sampling locations in Taiwan, Atmos. Environ., 44, 533–540, 2010. Ciucci, A., Palumbo, P., Brunetto, R., Della Corte, V., De Angelis, S., Rotundi, A., Rietmei
 - jer, F. J. M., Zona, E., Colangeli, L., Esposito, F., Mazzotta Epifani, E., Mennella, V., Inarta, S., Peterzen, S., Masi, S., and Ibba, R.: DUSTER (Dust in the Upper Stratosphere Tracking Ex-

periment and Retrieval) - preliminary results, Mem. S. A.It Suppl., 6, 119-124, 2011.

Daugeron, D., Renard, J.-B., Personne, P., Brun, G., and André, J. M.: Laboratory polarization nephelometer for measurements of optical properties of aerosols, Meas. Sci. Technol., 18, 632-638, 2007.

15

10

- Deshler, T., Hervig, M. E., Hofmann, D. J., Rosen, J. M., and Liley, J. B.: Thirty years of in situ stratospheric aerosol size distribution measurements from Laramie, Wyoming (41°N) using balloon-borne instruments, J. Geophys. Res., 108, 4167, doi:10.1029/2002JD002514, 2003. Diner, D. J., Ackerman, T. P., Anderson, T. L., Bösenberg, J., Braverman, A. J., Charlson, R. J.,
- Collins, W. D., Davies R, Holben, B. N., Hostetler, C. A., Kahn, R. A., Martonchik, J. V., 20 Menzies, R. T., Miller, M. A., Ogren, J. A., Penner, J. E., Rasch, P. J., Schwartz, S. E., Seinfeld, J. H., Stephens, G. L., Torres, O, Travis, L. D., Wielicki, B. A., and Yu, B.: PARAGON: An integrated approach for characterizing aerosol climate impacts and environmental interactions, B. Am. Meteorol. Soc., 85, 1491–1501, 2004.
- Dubovik, O. and King, M. D.: A flexible inversion algorithm for retrieval of aerosol optical prop-25 erties from Sun and sky radiance measurements, J. Geophys. Res., 105, 20673–20696, 2000.

Eidhammer, T., Montague, D. C., and Deshler, T.: Determination of index of refraction and size of supermicrometer particles from light scattering measurements at two angles, J. Geophys.

Res., 113, D16206, doi:10.1029/2007JD009607 10.1088/0957-0233/21/8/085901, 2008, 30 Francis, M., Renard, J.-B., Hadamcik, E., Couté, B., Gaubicher, B., and Jeannot, M.: New studies on scattering properties of different kinds of soot, J. Quant. Spectrosc. Ra., 112, 1766-1775, 2011.

| Discussion Paper | | AMTD 8, 9993–10056, 2015 | |
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10026

Gayet, J. F., Crépel, O., Fournol, J. F., and Oshchepkov, S.: A new airborne polar Nephelometer for the measurements of optical and microphysical cloud properties. Part I: Theoretical design, Ann. Geophys., 15, 451–459, doi:10.1007/s00585-997-0451-1, 1997.

Grimm, H. and Eatough, D. J.: Aerosol measurement: the use of optical light scattering for the

determination of particulate size distribution, and particulate mass, including the semi-volatile fraction, Japca. J. Air Waste Ma., 59, 101–107, 2009.

Gulijk, C., Marijnissen, J., Makkee, M., and Moulijn, J.: The Choice of Instrument (ELPI and/or SMPS) for Diesel Soot Particulate Measurements, SAE Technical Paper 2003-01-0784, doi:10.4271/2003-01-0784, 2003.

¹⁰ Hangal, S. and Willeke, K.: Aspiration Efficiency: Unified model for all forward sampling angles, Environ. Sci. Technol., 24, 688–691, 1990.

Hansen, J., Lacis, A., and Sato, M.: Potential climate impact of Mount Pinatubo eruption, Geophys. Res. Lett., 19, 215–218, 2003.

Hanson, D. R., Ravishankara, A. R., and S. Solomon: Heterogeneous reactions in sulphuric acid aerosols: a framework for model calculation, J. Geophys. Res., 99, 3615–3629, 1994.

acid aerosols: a framework for model calculation, J. Geophys. Res., 99, 3615–3629, 1994.
 Hanson, D. R., Ravishankara, A. R., and E. R. Lovejoy: Reactions of BrONO₂ with H₂O on submicron sulphuric acid aerosol and implication for the lower stratosphere, J. Geophys. Res., 101, 9063–9069, 1996.

Heim, M., Mullins, B. J., Umhauer, H., and Kasper, G.: Performance evaluation of three optical

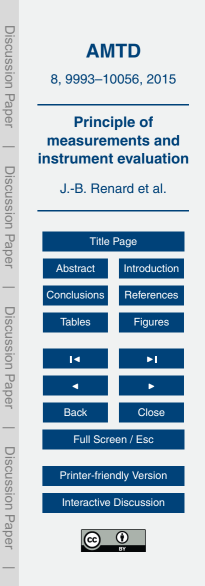
20 particle counters with an efficient "multimodal" calibration method, J. Aerosol Sci., 39, 1019– 1031, doi:10.1016/j.jaerosci.2008.07.006, 2008.

Hering, S. V. and McMurry, P. H.: Optical counter response to monodisperse atmospheric aerosols, Atmos. Environ., 25, 463–468, 1991.

Heyder, J. and Gehbart, J.: Gravitational deposition of particles from laminar aerosol flow through inclined circular tubes, J. Aerosol Sci., 8, 289–295, 1977.

25

- Jégou, F., Berthet, G., Brogniez, C., Renard, J.-B., François, P., Haywood, J. M., Jones, A., Bourgeois, Q., Lurton, T., Auriol, F., Godin-Beekmann, S., Guimbaud, C., Krysztofiak, G., Gaubicher, B., Chartier, M., Clarisse, L., Clerbaux, C., Balois, J. Y., Verwaerde, C., and Daugeron, D.: Stratospheric aerosols from the Sarychev volcano eruption in the 2009 Arctic
- ³⁰ summer, Atmos. Chem. Phys., 13, 6533–6552, doi:10.5194/acp-13-6533-2013, 2013. Kanakidou, M., Seinfeld, J. H., Pandis, S. N., Barnes, I., Dentener, F. J., Facchini, M. C., Van Dingenen, R., Ervens, B., Nenes, A., Nielsen, C. J., Swietlicki, E., Putaud, J. P., Balkanski, Y., Fuzzi, S., Horth, J., Moortgat, G. K., Winterhalter, R., Myhre, C. E. L., Tsigaridis, K.,



Vignati, E., Stephanou, E. G., and Wilson, J.: Organic aerosol and global climate modelling: a review, Atmos. Chem. Phys., 5, 1053–1123, doi:10.5194/acp-5-1053-2005, 2005.

- Lurton, T., Renard, J.-B., Vignelles, D., Jeannot, M., Akiki, R., Mineau, J.-L., and Tonnelier, T.: Light scattering at small angles by atmospheric irregular particles: modelling and laboratory
- measurements, Atmos. Meas. Tech., 7, 931–939, doi:10.5194/amt-7-931-2014, 2014.
 McDonald, R. and Biswas, P.: A methodology to establish the morphology of ambient aerosols, J. Air Waste Manage., 54, 1069–1078, 2004.

Mie, G.: Beiträge zur Optik trüber Medien, speziell kolloidaler Metallösungen, Annal. Phys., 3, 377–445, 1908.

Murphy, D. M., Cziczo, D. J., Hudson, P. K., and Thomson, D. S.: Carbonaceous material in aerosol particles in the lower stratosphere and tropopause region, J. Geophys. Res., 112, D04203, doi:10.1029/2006JD007297, 2007.

Ovarlez, J. and Ovarlez, H.: Water vapour and aerosol measurements during SESAME, and the observation of low water vapour content layers, Proc. Third European Workshop Polar

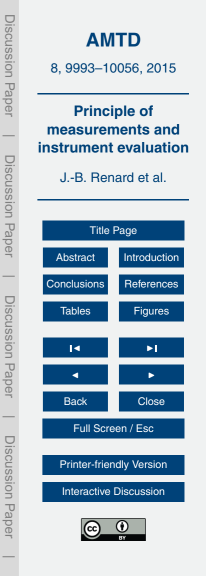
Stratospheric Ozone, Schliersee, Germany, 18–22 September 1995, Air Pollution Rep. 56, edited by: Pyle, J. A., Harris, N. R. P., and Amanatidis, G. T., European Commission, 205–208, 1995.

Quaas, J., Boucher, O., Bellouin, N., and Kinne, S.: Satellite-based estimate of the direct and indirect aerosol climate forcing, J. Geophys. Res., 113, D5, doi:10.1029/2007JD008962, 2008.

- Ramanathan, V., Crutzen, P. J., Lelieveld, J., et al.: Indian Ocean experiment: An integrated analysis of the climate forcing and effects of the great Indo-Asian haze, J. Geophys. Res., 106, 28371–28398, 2001.
 - Renard, J.-B., Worms, J.-C., Lemaire, T., Hadamcik, E., and Huret, N.: Light scattering by dust particles in microgravity: polarization and brightness imaging with the new version of the PROGRA2 instrument, Appl. Optics, 41, 609–618, 2002.

25

- Renard, J.-B., Brogniez, C., Berthet, G., Bourgeois, Q., Gaubicher, B., Chartier, M., Balois, J.-Y., Verwaerde, C., Auriol, F. Francois, P., Daugeron, D., and Engrand, C.: Vertical distribution of the different types of aerosols in the stratosphere: Detection of solid particles and analysis of their spatial variability, J. Geophys. Res., 113, D21303, doi:10.1029/2008JD010150, 2008.
- Renard, J.-B., Thaury, C., Mineau, J.-L., and Gaubicher, B.: Small-angle light scattering by airborne particulates: Environnement- S. A. continuous particulate monitor, Meas. Sci. Technol., 21, 931–939, doi:10.1088/0957-0233/21/8/085901, 2010a.



Renard, J.-B., Berthet, G., Salazar, V., Catoire, V., Tagger, M., Gaubicher, B., and Robert, C.: In situ detection of aerosol layers in the middle stratosphere, Geophys. Res. Lett., 37, L20803, doi:10.1029/2010GL044307, 2010b.

Renard, J.-B., Dulac, F., Berthet, G., Lurton, T., Vignelles, D., Jégou, F., Tonnelier, T., Thaury,

- ⁵ C., Jeannot, M., Couté, B., Akiki, R., Verdier, N., Mallet, M., Gensdarmes, F., Charpentier, P., Mesmin, S., Duverger, V., Dupont, J. C., Elias, T., Crenn, V., Sciare, J., Giacomoni, J., Gobbi, M., Hamonou, E., Olafsson, H., Dagsson-Waldhauserova, P., Camy-Peyret, C., Mazel, C., Décamps, T., Piringer, M., Surcin, J., and Daugeron, D.: LOAC: a small aerosol optical counter/sizer for ground-based and balloon measurements of the size distribution and nature of atmospheric particles. Part 2: First results from heliaen and unmanned particles.
- ture of atmospheric particles Part 2: First results from balloon and unmanned aerial vehicle flights, Atmos. Meas. Tech. Discuss., 8, 10057–10096, doi:10.5194/amtd-8-10057-2015, 2015

Rosen, J. M.: The vertical distribution of dust to 30 kilometers, J. Geophys. Res., 69, 4673–4676, 1964.

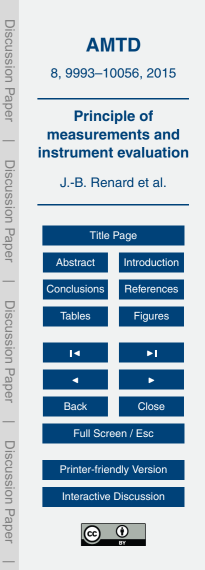
- ¹⁵ Ryder, C. L., Highwood, E. J., Rosenberg, P. D., Trembath, J., Brooke, J. K., Bart, M., Dean, A., Crosier, J., Dorsey, J., Brindley, H., Banks, J., Marsham, J. H., McQuaid, J. B., Sodemann, H., and Washington, R.: Optical properties of Saharan dust aerosol and contribution from the coarse mode as measured during the Fennec 2011 aircraft campaign, Atmos. Chem. Phys., 13, 303–325, doi:10.5194/acp-13-303-2013, 2013.
- ²⁰ Salazar, V., Renard, J.-B., Hauchecorne, A., Bekki, S., and Berthet, G.: A new climatology of aerosols in the middle and upper stratosphere by alternative analysis of GOMOS observations during 2002–2006, Int. J. Remote Sens., 34, 4986–5029, doi:10.1080/01431161.2013.786196, 2013.

Shaw, G. E., Reagan, J. A., and Herman, B. M.: Investigations of atmospheric extinction using

- direct solar radiation measurements made with a multiple wavelength radiometer, J. Appl. Meteorol., 12, 374–380, 1973.
 - Shi, J. P., Khan, A. A., and Harrison, R. M.: Measurements of ultrafine particle concentration and size distribution in the urban atmosphere, Sci. Total Environ., 235, 51–64, 1999.
 - Singh, V. P., Gupta, T., Tripathi, S. N., Jariwala, C., and Das, U.: Experimental study of the effects of environmental and fog condensation nuclei parameters on the rate of fog formation

30

and dissipation using a new laboratory scale fog generation facility, Aerosol Air Qual. Res., 11, 140–154, 2011.



Spencer, M. T., Shields, L. G., and Prather, K. A.: Simultaneous measurement of the effective density and chemical composition of ambient aerosol particles, Environ. Sci. Technol., 41, 1303–1309, 2007.

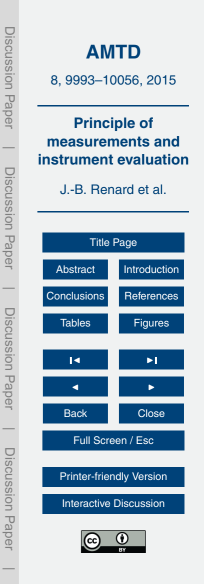
Virtanen, A., Rönkkö, T., Kannosto, J., Ristimäki, J., Mäkelä, J. M., Keskinen, J., Pakkanen, T.,

- Hillamo, R., Pirjola, L., and Hämeri, K.: Winter and summer time size distributions and densities of traffic-related aerosol particles at a busy highway in Helsinki, Atmos. Chem. Phys., 6, 2411–2421, doi:10.5194/acp-6-2411-2006, 2006.
 - Volten, H., Muñoz, O., Hovenier, J. W., and Waters, L. B.: An update of the Amsterdam Light Scattering Database, J. Quant. Spectrosc. Ra., 100, 437–443, 2006.
- ¹⁰ Winker, D. M., Pelon, J., Coakley Jr., J. A., Ackerman, S., A., Charlson, R. J., Colarco, P. R., Flamant, P., Fu, Q., Hoff, R. M., Kittaka, C., Kubar, T. L., le Treut, H., McCormick, M. P., Mégie, G., Poole, L., Powell, K., Trepte, C., Vaughan, M. A., and Wielicki, B. A.: The CALIPSO Mission, a global 3D view of aerosols and clouds, B. Am. Meteorol. Soc., 91, 1211–1229, 2010.
- ¹⁵ Whitby, K. T. and Vomela, R. A.: Response of single particle optical counters to nonideal particles, Environ. Sci. Technol., 1, 810–814, 1967.
 - Weiss-Wrana, K.: Optical properties of interplanetary dust comparison with light scattering by larger meteoritic and terrestrial grains, Astron. Astrophys., 126, 240–250, 1983.

Xiong, C. and Friedlander, S. K., Morphological properties of atmospheric aerosol aggregates, P. Natl. Acad. Sci. USA, 98, 11851–11856, 2001.

- Zemp, E., Elsasser, S., Schindler, C., Kunzli, N., Perruchoud, A. P., Domenighetti, G., Medici, T., Ackermann-Liebrich, U., Leuenberger, P., Monn, C., Bolognini, G., Bongard, J.-P., Brandli, O., Karrer, W., Keller, R., Schoni, M.-H., Tschopp, J.-M., Villiger, N., and Zellweger, J.-P.: Longterm ambient air pollution and respiratory symptoms in adults (SAPALDIA study), Am. J. Deep Orth Core, 150, 1057, 1000.
- ²⁵ Resp. Crit. Care, 159, 1257–1266, 1999.

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| Discussion Paper | AMTD 8, 9993–10056, 2015 | | | | |
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Table 1. The 19 size classes of LOAC for concentration measurements.

| | Diameter range (µm) |
|----|---------------------|
| 1 | 0.2–0.3 |
| 2 | 0.3-0.4 |
| 3 | 0.4-0.5 |
| 4 | 0.5–0.6 |
| 5 | 0.6-0.7 |
| 6 | 0.7–0.9 |
| 7 | 0.9–1.1 |
| 8 | 1.1–3.0 |
| 9 | 3.0-5.0 |
| 10 | 5.0-7.5 |
| 11 | 7.5–10.0 |
| 12 | 10.0–12.5 |
| 13 | 12.5–15.0 |
| 14 | 15.0–17.5 |
| 15 | 17.5–20.0 |
| 16 | 20.0-22.0 |
| 17 | 20.0-30.0 |
| 18 | 30.0-40.0 |
| 19 | 40.0–100.0 |
| | |

| Table 2. Conditions of measurements for evaluation exercises | | | | | | | |
|--|----------|-------------------|-------------------|----------------------------|--|--|--|
| Campaign | Location | Date | Installation | Instruments for validation | | | |
| ParisFog | SIBTA | Nov 2012–Apr 2013 | Continuous ground | – WELAS counter | | | |

| ParisFog | SIRTA Observatory, Palaiseau (France) | Nov 2012–Apr 2013 Sep 2013–Jan 2014 | Continuous ground measurements | WELAS counter Fog monitor counter Scanning Mobility Particle Sizer (SMPS) |
|--|---|--|---|---|
| Cloud measurement | Puy de Dôme s (France) | May 2013 | Continuous ground measurements | Well-known atmospheric conditions for the topology identification |
| ChArMEx | Minorca (Spain) | 17 Jun 2013 | Tropospheric pressurized balloon | Well-known atmospheric conditions for the topology identification |
| ChArMEx | lle du Levant (France) | 22 Jul 2013 | Tropospheric pressurized balloon | Well-known atmospheric conditions for the topology identification |
| ChArMEx | Minorca (Spain) | 15 Jun 2013–2 Jul 2013 | Continuous ground measurements | HHPC-6 counter |
| ChArMEx | Minorca (Spain) | 16 and 17 Jun 2013 | Meteorological sounding balloon flights | Well-known atmospheric conditions for the topology identification |
| ChArMEx | Minorca (Spain) | 16 and 19 Jun 2013 | Meteorological and pressurized tropospheric balloon flights | WALI lidar |
| QAIDOMUS | Orléans (France) | Sep-Nov 2013 | Indoor air | TEOM microbalance |
| VOLTAIRE- LOAC | Reykjavik (Iceland) | 7 Nov 2013 | Meteorological balloon flight | Well-known atmospheric conditions for the topology identification |
| Observatoire Atmo- sphérique Generali | Paris (France) | Jan–Apr 2014 | Permanent measurements on tethered balloon (at ground and up to an altitude of 270 m) | TEOM microbalances (Airparif air quality network) Well-known atmospheric conditions for the topology identification |
| SIRTA5 campaign | Gif-sur-Yvette (France) | 3–13 Feb 2014 | Continuous ground measurements at SIRTA | – Grimm counter – HHPC-6 counter – SMPS |



Discussion Paper

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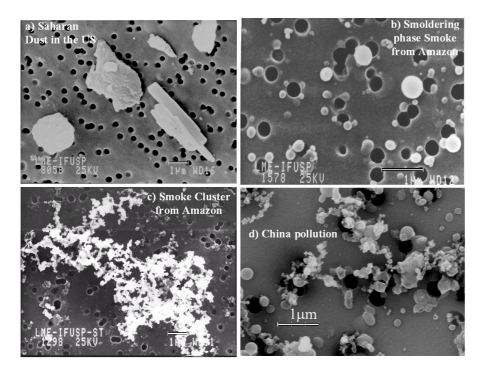


Figure 1. Scanning electron microscope image of ambient air aerosols (courtesy Jose Vanderlei Martins, Institute of Physics of the University of Sao Paulo, Brazil).



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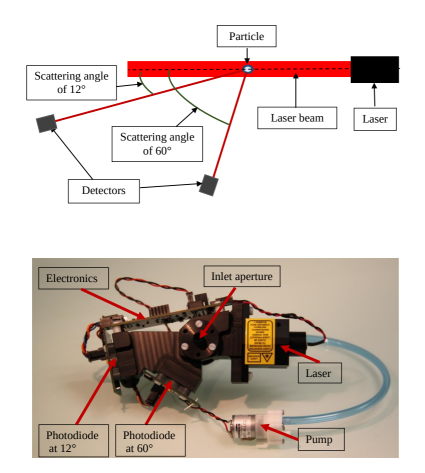
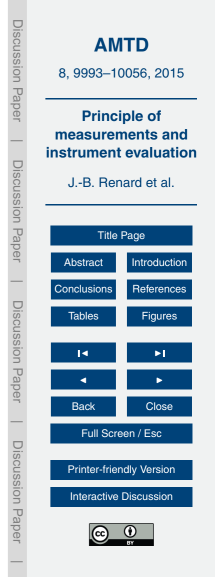
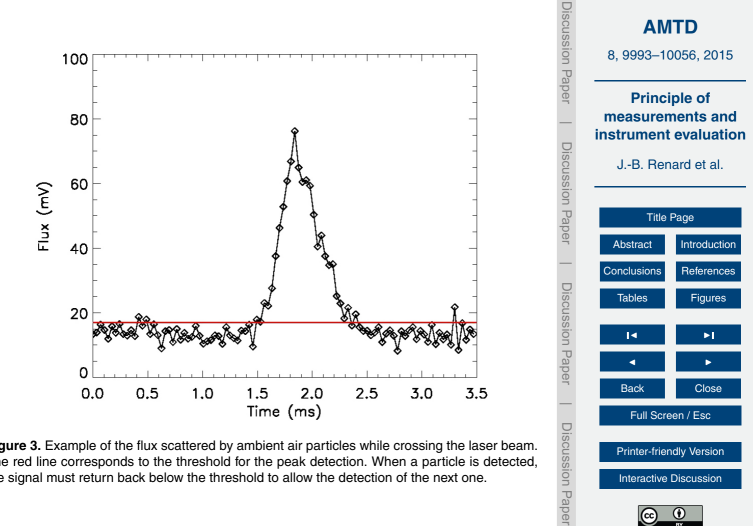


Figure 2. The LOAC instrument; upper panel: principle of measurement; lower panel: picture of the instrument (the inlet tube is not presented here).





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Figure 3. Example of the flux scattered by ambient air particles while crossing the laser beam. The red line corresponds to the threshold for the peak detection. When a particle is detected, the signal must return back below the threshold to allow the detection of the next one.

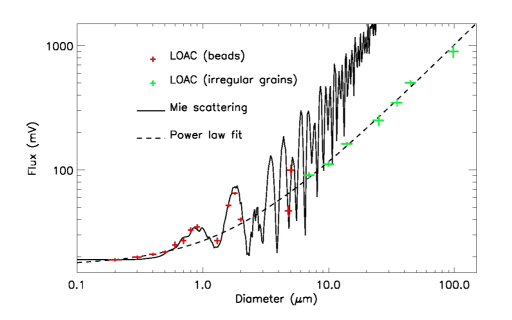
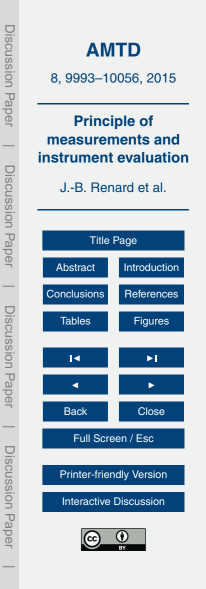


Figure 4. Calibration curve of the scattered flux at the 12° channel as a function of particle diameter. Beads were used in the 0.2–5.0 μ m range; irregular grains selected by sifters were used for the largest size. The offset due to the electronic dark current and high frequency noise of the detector were added to the calculations; thus the curve asymptotically decreases with decreasing size to this offset value. The Mie calculations were conducted for the LOAC field of view. The difference between the Mie scattering calculations and LOAC measurements for diameters greater than 5 μ m is due to the small aperture of the field of view coupled with the roughness of the particle shapes; the measurement curve is fitted by a power law (adding an offset of 17 mV).



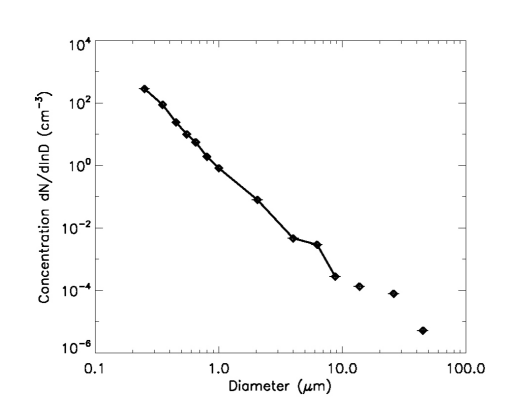
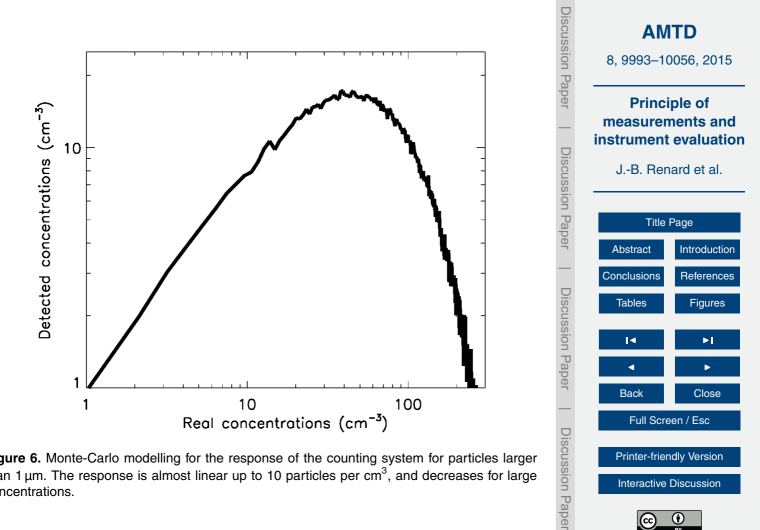
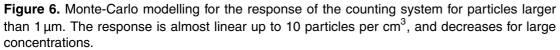


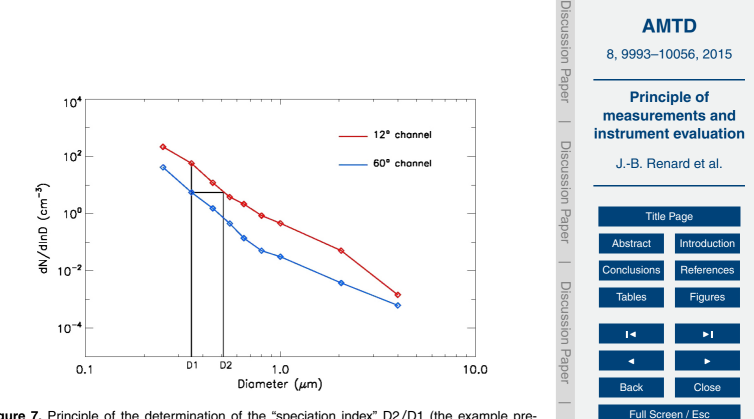
Figure 5. Typical size distribution in a suburban ambient air with carbon particles (Palaiseau, South of Paris) on 14 October 2013 during ParisFog campaign; the data are integrated during 15 min; the last points are not related because of zero concentration measured between them.





Interactive Discussion





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Figure 7. Principle of the determination of the "speciation index" D2/D1 (the example presented here uses real measurements).

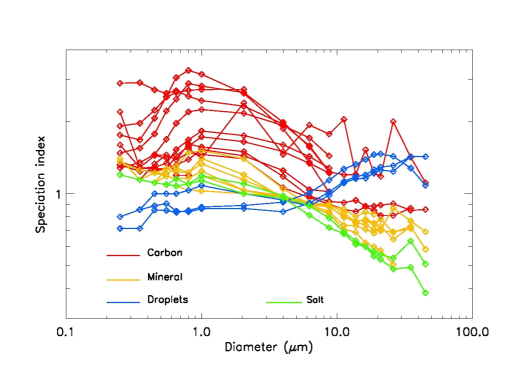
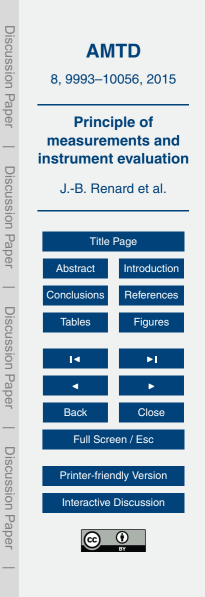
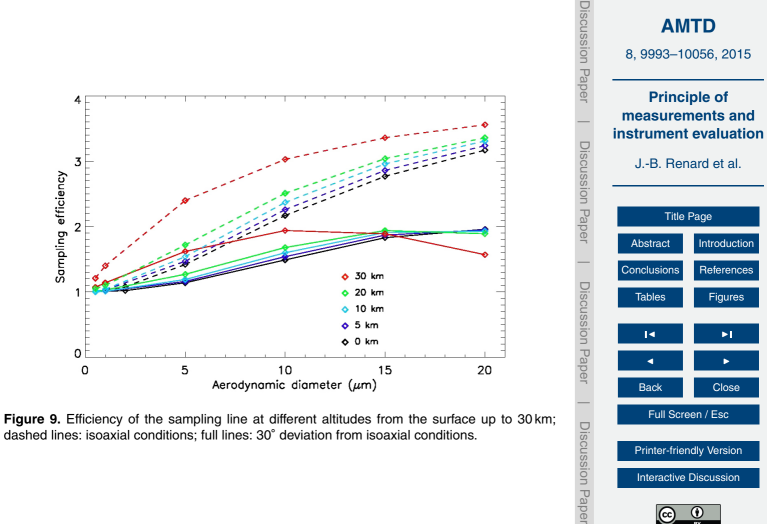


Figure 8. Evolution of the speciation index with diameter for various families of samples; measurements were conducted in laboratory with LOAC using pure samples.

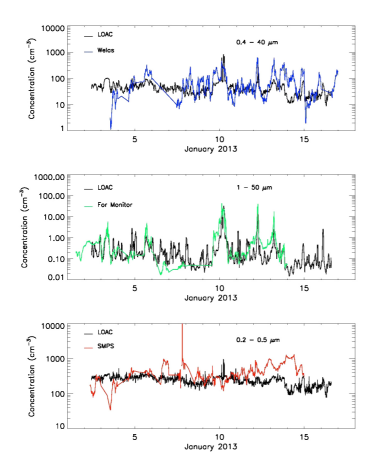


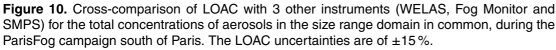


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dashed lines: isoaxial conditions; full lines: 30° deviation from isoaxial conditions.







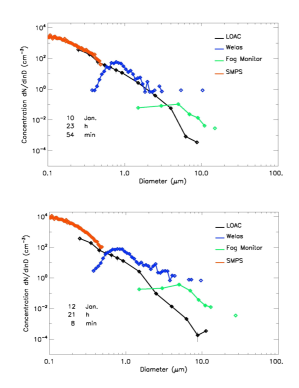
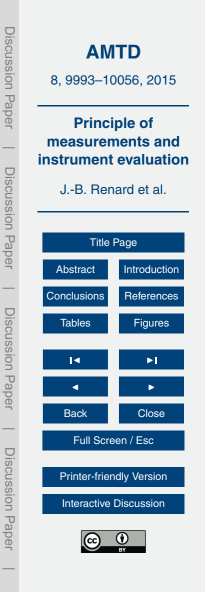
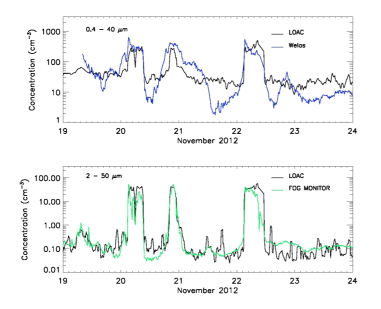
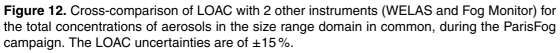


Figure 11. Cross-comparison of the 4 instruments during background conditions, in case of good agreement for the total concentrations measurements, during the ParisFog campaign. Upper panel: 10 January 2013, good agreement between the instruments; lower panel: 12 January 2013, poor agreement. The LOAC uncertainties are of ± 15 %. The WELAS probably underestimates sub-µm particles (Heim et al., 2008).









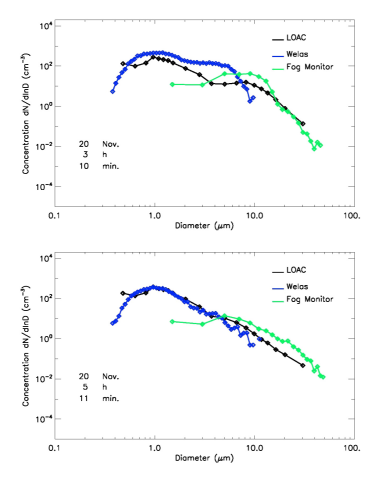
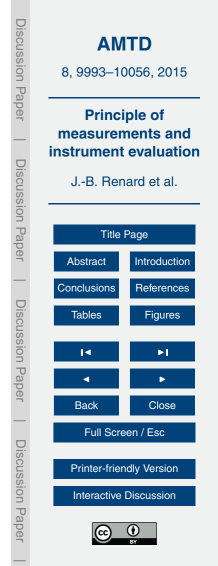


Figure 13. Cross-comparison of the 3 instruments at the beginning of the fog event (top) and at the end (bottom), during the ParisFog campaign on 20 November 2012 during a fog event. The LOAC uncertainties are of ± 15 %. The WELAS probably underestimates sub-µm particles.



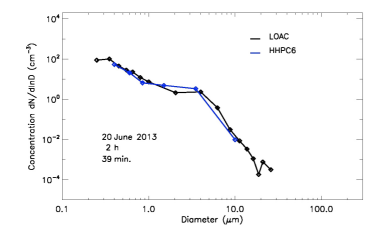
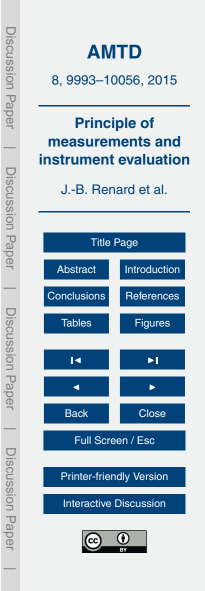
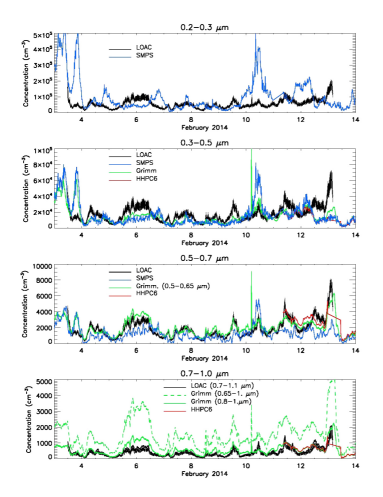
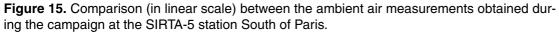
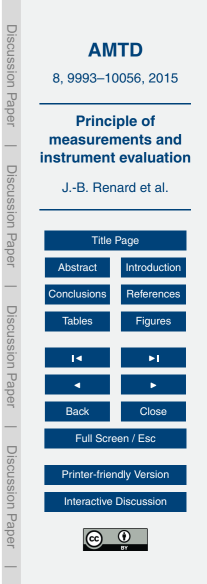


Figure 14. Example of size distribution for LOAC and HHPC-6 during an event of solid particles during the ChArMEx campaign at Minorca on 20 June 2013.









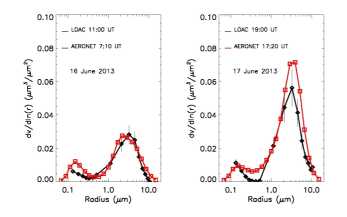


Figure 16. Comparison between integrated LOAC measurements from vertical profiles obtained under meteorological balloons and AERONET measurements during an African dust transport event during the ChArMEx 2013 campaign (note that the LOAC data are given in radius to match the AERONET format).



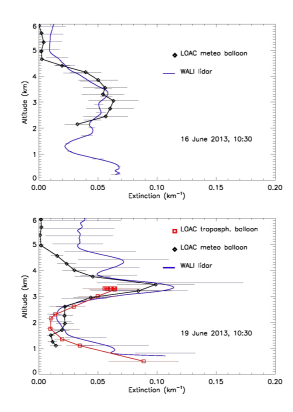




Figure 17. Extinction profiles of the WALI lidar and extinction profiles calculated from LOAC measurements under meteorological and pressurized tropospheric balloons, from Minorca Island during the ChArMEx campaign.

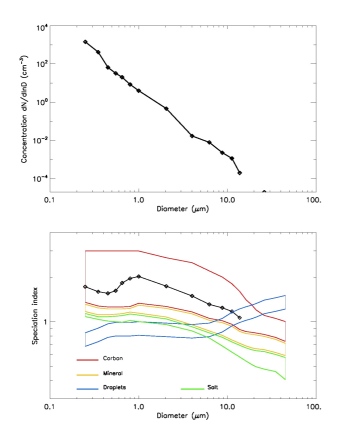
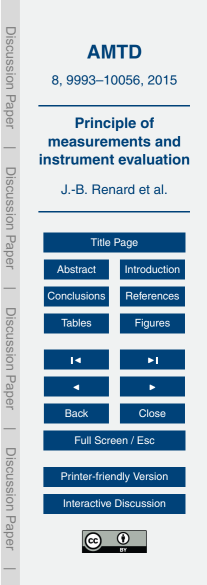


Figure 18. Example of the detection of carbon particles in urban air, in South-West of Paris on 29 December 2013 around 07:30 UT, at the "Observatoire Atmosphérique Generali"; upper panel: size distribution; lower panel: topology.



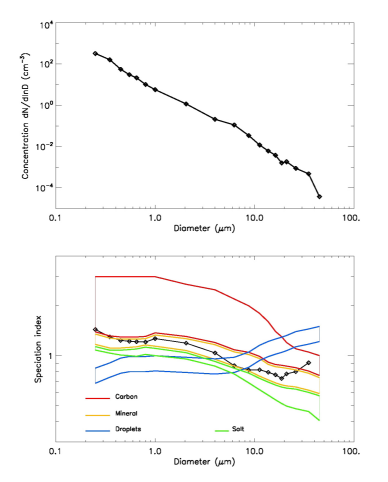
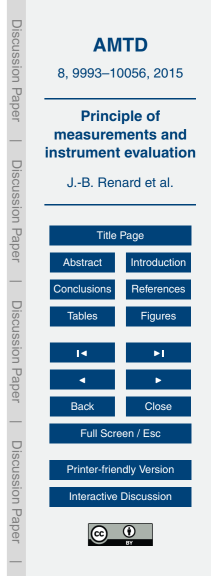


Figure 19. Example of the detection of sand particles above Mediterranean Sea (close to Minorca) from a drifting pressurized tropospheric balloon on 17 June 2013 around 14:30 UT at an altitude of 2050 m, during the ChArMEx campaign; upper panel: size distribution; lower panel: topology.



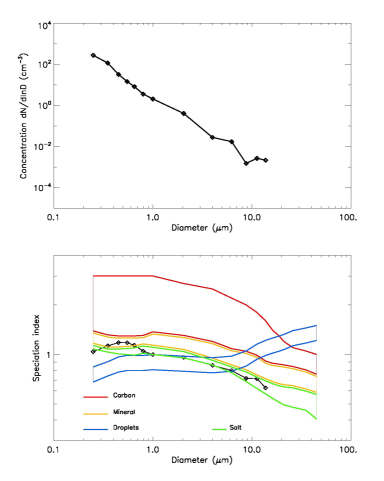
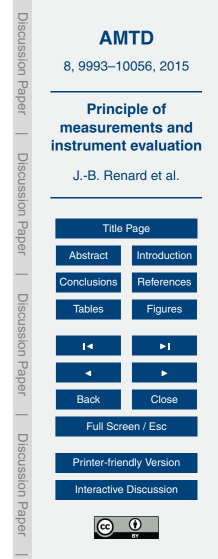


Figure 20. Example of the detection of salt particles above Mediterranean Sea (close to Minorca, Spain) from balloon on 22 July 2013 at 21:25 UT at an altitude of 275 m during the ChArMEx campaign; upper panel: size distribution; lower panel: topology.



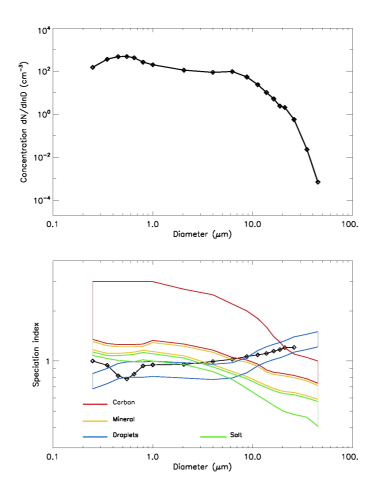
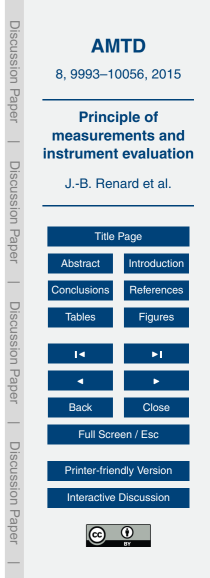


Figure 21. Example of measurements inside a cloud at Puy de Dôme observatory (France) on 15 May 2013 at 10:30 UT; upper panel: size distribution; lower panel: topology.



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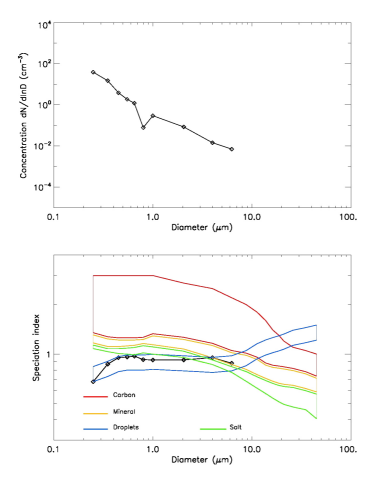
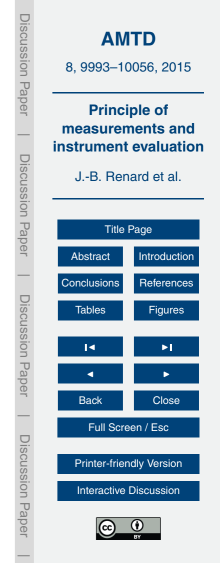


Figure 22. Example of measurements inside a haze or thin cloud at an altitude of 1.2 km during a flight under meteorological balloon from Reykjavik (Iceland) on 7 November 2013 at 12:30 UT; upper panel: size distribution; lower panel: topology.



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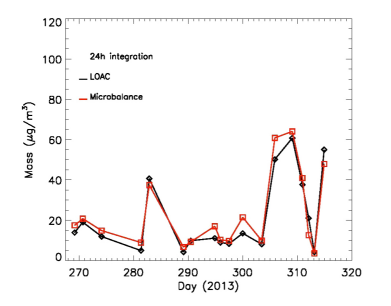


Figure 23. Comparison of coincident LOAC and TEOM microbalance measurement in indoor air (averaged over 24 h); particles have been injected with various concentrations to document a large range of mass concentration.



