Facility for production of ambient-like model aerosols (PALMA) in the laboratory: application in the intercomparison of automated PM monitors with the reference gravimetric method

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Abstract. A new facility has been developed which allows for a stable and reproducible production of ambient-like model aerosols (PALMA) in the laboratory. The set-up consists of multiple aerosol generators, a custom-made flow tube homogeniser, isokinetic sampling probes, and a system to control aerosol temperature and humidity. Model aerosols containing elemental carbon, secondary organic matter from the ozonolysis of α-pinene, inorganic salts such as ammonium sulfate and ammonium nitrate, mineral dust particles, and water were generated under different environmental conditions and at different number and mass concentrations. The aerosol physical and chemical properties were characterised with an array of experimental methods, including scanning mobility particle sizing, ion chromatography, total reflection X-ray fluorescence spectroscopy and thermo-optical analysis. The facility is very versatile and can find applications in the calibration and performance characterisation of aerosol instruments monitoring ambient air. In this study, we performed, as proof of concept, an intercomparison of three different commercial PM (particulate matter) monitors (TEOM 1405, DustTrak DRX 8533 and Fidas Frog) with the gravimetric reference method under three simulated environmental scenarios. The results are presented and compared to previous field studies. We believe that the laboratory-based method for simulating ambient aerosols presented here could provide in the future a useful alternative to time-consuming and expensive field campaigns, which are often required for instrument certification and calibration.

1 Introduction

Atmospheric pollution by airborne particles significantly contributes to climate change and has been linked to respiratory and cardiovascular diseases and lung cancer (Fuzzi et al., 2015; Kim et al., 2015; WHO, 2013). It has been estimated that in Europe alone more than 500 000 deaths per year can be attributed to PM exposure, and that pollution hot spots of PM are responsible for a loss in life expectancy of up to 36 months (Fuzzi et al., 2015). For EU member states, air quality monitoring – as laid down in the Air Quality Directive 2008/50/EC (European Parliament, 2008, 2015) – is mandatory and comprises quantification of airborne particulate matter (PM) and some of its constituents. The most important regulated metric to monitor particulate air pollution is the mass concentration, or more specifically the total mass per unit volume of air of particulate matter which is small enough to pass through a size-selective inlet with a
50 % efficiency cut-off at 2.5 and 10 µm aerodynamic diameter, commonly referred to as PM$_{2.5}$ and PM$_{10}$ respectively. Ambient limit values for PM$_{2.5}$ and PM$_{10}$ have been established in Europe (European Parliament, 2008, 2015; FOEN, 2018), the USA (US-EPA, 2016) and other countries worldwide.

Regulatory bodies, air quality networks and atmospheric instrument manufacturers all strive to improve air quality monitoring, yet there is still a lack of metrological traceability in airborne PM measurements. PM mass concentration was established as the default metric of PM based on the assumption that mass measurements are straightforward; they can be performed with a conventional balance. The gravimetric filter-based reference methods for PM$_{10}$ and PM$_{2.5}$ are set out in the standards EN 12341:2014 (CEN/TC 264/WG-15, 2014) and EN 14907:2005; however, they fall short in areas such as time resolution and ongoing quality assurance and quality control to control the effects of semi-volatile particles and water absorption by particles, for example (CEN/TC 264/WG-15, 2014; Eisner and Wiener, 2002; Hauck et al., 2004; Zhu et al., 2007). The measurement uncertainties for PM mass concentration in the directive (European Parliament, 2008, 2015) are 25 %, and thus much higher than those for gaseous pollutants (typically 15 %).

Automatic PM monitoring systems were developed in order to avoid these drawbacks and enable time resolutions below 24 h (Schwab et al., 2006; Weingartner et al., 2011; Zhu et al., 2007); however, demonstrating their equivalence to the reference manual gravimetric method is time consuming and expensive (Hauck et al., 2004; Zhu et al., 2007). There are also inconsistencies in the automatic instruments based on different working principles (e.g. light scattering, beta absorption, oscillating microbalance) and the variations of the aerosols used for comparison. Ambient PM is not uniform with respect to chemical composition, particle size and shape. In most cases, PM does not refer to a single pollutant with a distinct chemical signature, but rather to a highly variable mixture of combustion particles, salts, mineral dust, organic substances and other materials (Hueglin et al., 2005; Putaud et al., 2010; McNamara et al., 2011). Such model aerosols, however, are only partially representative of ambient air since they fail to account for organic particles, such as sucrose and adipic acid (Zhang et al., 2018). Such model aerosols, however, are only partially representative of ambient air since they fail to account for carbonaceous particles and the complex organic matter, which constitute a considerable mass fraction of airborne particulates (Hueglin et al., 2005; Putaud et al., 2010). Light-scattering PM monitors are very sensitive to the aerosol size distribution, refractive index (i.e. chemistry) and humidity, and research findings suggest that a rigorous calibration with “tailored” aerosols, i.e. aerosols representative of the environment of their intended use, is needed (Jayaratne et al., 2020; McNamara et al., 2011).

The goal of this study was to develop a standardised laboratory-based calibration procedure for automatic PM-measuring instruments under well-controlled and reproducible experimental conditions. Multi-component model aerosols were generated in order to reproduce the main properties of real ambient air in terms of particle size distribution, chemical composition and number/mass concentration, including semi-volatility and hygroscopicity. The properties of ambient air, of course, may differ dramatically from place to place. Here, the main focus was on simulating aerosols encountered in Europe (Putaud et al., 2010), which are dominated by organic matter, inorganic ions (predominantly sulfate and nitrate, and to a lesser extent ammonium), carbonaceous particles (mostly from fossil fuel combustion rather than biomass burning), mineral dust and water.

Apart from the aerosol generation system (detailed below), the new set-up comprises a flow tube homogeniser and a system for reference gravimetric measurements. The facility is very versatile: the total PM mass concentration of the model aerosols can be adjusted in a range from a few micrograms per cubic metre up to about 500 µg m$^{-3}$; the percentage fraction of each PM constituent can be tuned to simulate different urban, suburban or rural aerosols; and the aerosol temperature and relative humidity can be adjusted to simulate winter or summer-like environmental conditions. As a proof of concept, three different automated PM monitors, the TEOM 1405 (Thermo Scientific, USA), the DustTrak DRX 8533 (TSI Inc., USA) and the Fidas Frog (Palas, Germany), were compared with the reference gravimetric method under three different environmental scenarios. To our knowledge, this is the very first intercomparison involving the Fidas Frog.

Here, we focused on the calibration of the PM monitors’ particle quantification, rather than the particle inlet size selection; i.e. the TEOM 1405 unit was calibrated without...
its PM sampling inlet. The Fidas Frog and DustTrak DRX 8533, which are optical instruments, do not possess any size-selective inlet. The facility could be, however, extended in the future to calibrate PM monitors together with their sampling inlets, if needed. Finally, the facility for generating ambient-like model aerosols presented in this study is not only relevant for the calibration of PM monitors but can find applications in the performance evaluation and quality assurance of other aerosol instruments meant for monitoring ambient, indoor and workplace air as well as in controlled health studies and in vitro toxicology.

2 Design and validation of the experimental set-up

The experimental set-up consists of three distinct parts: (i) the generators of the primary aerosols (dust, salts, soot and aged soot); (ii) a flow tube homogeniser for aerosol mixing, including isokinetic sampling probes; and (iii) a system for reference gravimetric measurements. Each part is described in more detail in the following subsections.

2.1 Aerosol generation

Four primary aerosols, fresh soot, aged (i.e. organically coated) soot, inorganic salt and mineral dust particles, were generated as depicted in Fig. 1. Fresh soot particles were generated with a miniCAST 6204 burner (Jing Ltd., Switzerland). The operation point was optimised to produce combustion particles with a geometric mean mobility diameter (GMD) of 90 nm and EC / TC (elemental carbon to total carbon) mass fraction of > 90%. The combustion aerosol was split in two portions; one portion was led to the exhaust and the other through a metallic agglomeration tube (1.2 m long, 5 mm internal diameter), where the soot particles grew to about 120 nm. The mobility diameter was measured by a scanning mobility particle sizer (SMPS). The combustion aerosol was subsequently diluted by a factor of 10 with a VKL10 dilution unit (Palas, Germany). The outlet flow was divided into an oxidation flow reactor known as the Micro Smog Chamber (MSC prototype developed by Keller and Burtscher, 2012, and used by Bruns et al., 2015; Corbin et al., 2015b, 2015a; Keller and Burtscher, 2012), where fresh soot particles (120 nm mobility diameter) were sampled from the exhaust of the VKL10 dilution unit with the use of a second Micronel blower at flows between 1 and 2 L min\(^{-1}\). Inorganic salt particles were generated by nebulising aqueous mixtures of ammonium sulfate and ammonium nitrate at various ratios with the use of a TSI 3076 atomiser (TSI Inc., USA). The particles were then mixed with a controlled amount of \(\alpha\)-pinene vapours (≥ 97 % purity, Sigma Aldrich, Switzerland) under dry conditions (RH < 5 %). The concentration of \(\alpha\)-pinene at the inlet of the MSC was determined with a photoionisation detector (PID PhoCheck TIGER, Ion Science Ltd, UK) after filtering out the particles. The concentration could be varied by adjusting the flow of air through the \(\alpha\)-pinene container (gas bubbler) and typically ranged between 60 and 70 ppm. RH was measured with a digital humidity sensor (FHAD 46 series/Almemo D6, Ahlborn, Germany). The aerosol flow through the MSC was set to 1.2 L min\(^{-1}\) with the use of a miniature radial air blower (model H015X-525A9 with controller, Micronel AG, Switzerland). Higher aerosol flows through the MSC would lead to the residence time in the reactor being too short and should be avoided. \(\alpha\)-Pinene underwent ozonolysis in the MSC, forming secondary organic aerosol (SOA), part of which condensed on the surface of the soot particles, simulating atmospheric ageing procedures (Ess et al., 2020).

The GMD of the soot mobility size distribution was shifted to 160 nm upon coating with SOA, and the EC / TC mass fraction dropped to about 20 %. In parallel, fresh soot particles (120 nm mobility diameter) were sampled from the exhaust of the VKL10 dilution unit with the use of a second Micronel blower at flows between 1 and 2 L min\(^{-1}\). Mineral dust particles (ISO 12103-1 A2 fine dust test, Powder Technology Inc., USA) were generated with a rotating brush generator (RBG 1000, Palas, Germany) and were injected horizontally into an empty vessel, which acted as a swirl separator, filtering out the largest size fraction above PM\(_{10}\). Alternatively, whenever calibration with respect to the PM\(_{2.5}\) fraction is desired, a PM\(_{2.5}\) impactor can be installed right before injecting the dust particles into the homogeniser.

The primary aerosols were introduced into a flow tube homogeniser (see Sect. 2.2) through separate injection ports. The flow of each primary aerosol entering the homogeniser could be regulated with separate mass flow controllers (Redy MFC, Vögtlin, Switzerland) by splitting and directing part of the main primary aerosol flow to the exhaust. A filter (HEPA capsule, Pall Corporation, USA) was placed upstream of each MFC to remove the particles from the air flow. All four MFCs were connected to the same aerosol pump (VTE8, Thomas, Germany) as shown in Fig. 1.

The mobility diameter and number concentration of the soot and salt particles were determined with a scanning mobility particle sizer (SMPS 4.500, Grimm Aerosol Technik GmbH & Co. KG, Germany, L-DMA, Am-241 neutraliser,
scan time 695 s). The mass concentration of each primary aerosol was measured with a tapered element oscillating microbalance (TEOM 1405, Thermo Scientific, USA), operated at a flow rate of 3 L min\(^{-1}\) and a temperature of 30 °C. The TEOM data were recorded via a custom-made LabVIEW routine every 6 s without averaging. The size distribution of the dust particles was measured with a Fidas Frog fine-dust monitor (Palas, Germany) and a high-resolution optical particle counter LAS-X II (Particle Measuring Systems, USA).

### 2.2 Aerosol homogenisation and sampling

The homogeniser is a 2.1 m long custom-made stainless steel tube with an inner diameter of 16.4 cm, placed vertically. The design is based on a previous study but has been significantly improved, and the facility has been shortened (Horender et al., 2019). The tube is equipped with five identical inlets, placed at the very top as shown in Figs. 1 and 2a. Dilution air (filtered, humidity and temperature controlled) is delivered to each one of the inlets at a flow rate of 24 L min\(^{-1}\). The air is conditioned in two steps (Niedermeier et al., 2020) in such a way that the humidified air is particle-free: first, the dew point is adjusted by passing the air through a Nafion humidifier (Series FC125-240-10MP, PermaPure, USA) filled with water (ultra-analytic grade, Purelab ultra, ELGA, Switzerland) at a preselected water temperature, adjusted between 3 and 30 °C with a cryostat–thermostat (LAUDA Ecoline Staredition RE 306, Lauda DR. R. Wobser GmbH & Co. KG, Germany). After being put through the Nafion humidifier, the air is fully saturated with water. Subsequently, the air is guided through a heated hose (Series T-7000, Thermocoax Isopad GmbH, Germany), where the temperature can be adjusted up to 100 °C. The temperature and RH of the aerosol were monitored in the homogeniser at the height of the sampling probes with digital sensors (FHAD 46 series/Almemo D6, Ahlborn, Germany).

The primary aerosols are injected in the middle of the tube through separate ports located 50 cm downstream as shown in Fig. 2b. The dilution air sweeps the particles down the tube, where they are further mixed by three turbulent jets of air. The three air-jet injection tubes (flow rate 20 L min\(^{-1}\) each) are placed symmetrically around the homogeniser tube pointing 60° downwards (Fig. 2b). The total flow rate of...
the homogenised aerosol is hence equal to 180 L min\(^{-1}\) plus the flows of the four primary aerosols (in total less than 10 L min\(^{-1}\)). The temperature and relative humidity of the air jets are adjusted as described above for the dilution air. Finally, the homogeniser is surrounded by copper tubes with flowing water in order to maintain the stainless-steel tube at the same temperature as the aerosol. The temperature of water is adjusted by a flow-type cooler (AS-160 Green Line, Lindr, Czech Republic) or a thermostat (LAUDA EcoGold E4, Lauda DR. R. Wobser GmbH & Co. KG, Germany). The water flows in a closed loop, i.e. circulates back to the cryostat–thermostat as shown in Fig. 1. Currently, the homogeniser can only be cooled down to about 10\(^\circ\)C, and this poses limitations to the environmental conditions which can be simulated in the laboratory; even though the aerosol entering the homogeniser can be preconditioned at a temperature down to about 5\(^\circ\)C, the aerosol temperature at the outlet of the homogeniser will always be \(\geq 10\,^\circ\)C.

The sampling zone is located 1.25 m downstream of the injection position and accommodates isokinetic sampling probes (funnels) placed at the bottom end of the homogeniser as illustrated in Fig. 2c. Isokinetic conditions are necessary when sampling with instruments operating at different flow rates to ensure representative sampling, e.g. by minimising sampling artefacts of larger particles. Several custom-made sampling probes with different cross sections have been therefore designed to match the flow rate of the various automated PM monitors, which typically ranges between 0.2 and 20 L min\(^{-1}\). It is worth noting that the sampling system is highly adaptable; the lower end (outlet) of each sampling probe has custom-made threads so that it can be screwed in and out of the bottom metallic plate of the homogeniser. This ensures that the sampling probes can be readily exchanged before each experiment depending on the specifications of the PM monitors under testing. Finally, the excess aerosol flow exits the homogeniser through an exhaust outlet connected to a vacuum line as illustrated schematically in Fig. 1.

Figure 2. (a) Computer-aided design (CAD, Inventor Professional 2019, Autodesk, USA) of the homogeniser. Panels (b) and (c) show enlarged views of the primary aerosol inlets and isokinetic sampling probes, respectively.

To characterise the aerosol homogeneity in the flow tube as a function of particle size, sodium chloride (NaCl) particles with a geometric mean mobility diameter of 50 nm and mineral dust particles with an aerodynamic diameter in the lower micrometre range (ISO A2 dust) were generated with a nebuliser and a rotating-brush generator, respectively, as described in Sect. 2.1. Two parallel sampling lines were inserted into the flow tube at the height where the sampling probes would be normally located; the position of the first sampling line was kept fixed at the centre of the flow tube (radial position 0), whereas the second one was placed consecutively at a distance \(i = -70, -50, -30, -10, +10, +30, +50\) and +70 mm with respect to the centre. The outlet of each sampling line was connected to a calibrated condensation particle counter (CPC; Models 3775 and 3776, respectively, TSI inc., USA). In total, concentration measurements at eight different positions along the diameter of the flow tube were performed. The particle number concentration measured at the centre was used as reference \((C_{\text{ref}} = C_{0})\), and the aerosol homogeneity was calculated as \(C_i / C_{\text{ref}}\). The flow rate of each CPC was 0.3 L min\(^{-1}\), and the inner diameter of the sampling line was 6 mm. This configuration ensured nearly isokinetic sampling.

The tests were performed with NaCl and mineral dust particles separately. In both cases the aerosol spatial homogeneity was found to be well within 3 % in number concentration as shown in Fig. 3a and b, respectively, indicating that the particle mixing characteristics do not depend on particle size in the tested range (i.e. from lower nanometre to lower micrometre range). A final test was performed by mixing NaCl and dust particles to investigate whether the particle mixing properties are affected when two primary aerosols are introduced into the homogeniser simultaneously. It was confirmed that the aerosol homogeneity remains well within \(\pm 3\,\text{\%} (\text{measurements not shown})\), indicating that the simultaneous injection of primary aerosols into the homogeniser through separate ports (see Fig. 2b) does not compromise particle mixing in any way.

By calculating the standard deviation of all 28 measured data points, the spatial inhomogeneity of the aerosol in terms of number concentration was found to be 1.3 % for coverage factor \(k = 1\) (i.e. 68 % confidence level) or 2.6 % for \(k = 2\).
Table 1. Example of the uncertainty budget for a PM$_{10}$ mass concentration of 40 µg m$^{-3}$ and a sampling time of 240 min.

| Quantity | Value (example) | Standard uncertainty $(k = 1)$ | Relative uncertainty $(95 \%$ confidence level) |
|----------|----------------|--------------------------------|-----------------------------------------------|
| $t$      | 240 min        | negligible                     | negligible                                    |
| $P_{rel}$ | 1.00           | 0.01                           | 2 %                                          |
| $\eta_{hom}$ | 1.000         | 0.013                           | 2.6 %                                        |
| $Q$      | 38.333 L min$^{-1}$ | 0.058 L min$^{-1}$ | 0.30 % $^a$                                   |
| $m$      | 368.0$^b$ µg m$^{-3}$ | 8.4 µg m$^{-3}$ | 4.6 %                                        |
| $C_{m, ref}$ | 40.00 µg m$^{-3}$ | 1.13 µg m$^{-3}$ | 5.7 %                                        |

$^a$ The mass flow meter (Natec Sensors GmbH, Germany) was calibrated at METAS in a traceable manner. The expanded relative uncertainty on the calibration certificate amounts to 0.15 %. Here, a conservative estimation of 0.30 % was made to account for possible drifts since the time of calibration. $^b$ Assuming no loss of particulate mass during filter conditioning.

Briefly, model aerosols were drawn through 47 mm PTFE-coated glass fibre filters (Measurement Technology Laboratories, USA) placed in a metallic filter holder (C806 standard aerosol filter holder, Merck Millipore, Germany). The aerosol flow was controlled with a needle valve and measured with a calibrated mass flow meter (Natec Sensors GmbH, Germany) connected to an aerosol pump (VTE8, Thomas, Germany) in such a way that the volumetric flow corresponded to 2.3 m$^3$ h$^{-1}$ at ambient conditions. Here, ambient conditions refer to the aerosol temperature and pressure in the homogeniser at the height of the sampling probes. In the EN 12341 standard, the requirement that the aerosol flow be set to 2.3 m$^3$ h$^{-1}$ ($=38.33$ L min$^{-1}$) at ambient conditions arises from the need to accurately define the size cut-off of the PM inlets, a property that depends on the inlet flow. Since the custom-made facility developed in this study aims at calibrating the PM monitors without their respective PM inlet, this flow requirement is here largely superfluous, apart from effects on sampling from the velocity of air through the filter. Nevertheless, during the experiments the aerosol flow was set to $2.3$ m$^3$ h$^{-1}$ at ambient conditions to facilitate comparison between the conventional field-based and the new laboratory-based procedures. The connecting tube between the isokinetic sampling probe (i.e. central sampling funnel in Fig. 2c) and the filter holder was made of inert, electrically conducting rubber material and was kept as short as possible ($\approx 5$ cm) without bends to minimise deposition losses of particulate matter by kinetic processes as well as losses due to thermal, chemical or electrostatic processes. Finally, the laboratory temperature and pressure were kept constant at $(21 \pm 1)$ °C and $(950 \pm 20)$ hPa, respectively.

Before sampling, the filters were conditioned and weighed at the National Physical Laboratory (NPL) and shipped in individual plastic containers to the Federal Institute of Metrology METAS. After sampling, the filter samples were placed in Petri dishes, wrapped tightly in plastic covers and stored at 4 °C for about a week. They were then shipped to NPL for conditioning and weighing. NPL use a Measurement Technology Laboratories robotic filter weighing system that com-
prises an environmental chamber (20°C±1°C and 47.5%±2.5% relative humidity), an autohandler system and a Mettler Toledo XP2U balance. The filters are conditioned in the chamber for 48 h before weighing. The filters are weighed, and then the system pauses for 24 h before reweighing the filters to identify any time variation in filter mass. Numerous quality assurance and quality control checks are made before each set of weighings.

2.4 Uncertainty budget for the laboratory-based calibration of PM monitors

The reference mass concentration, \( C_{m,\text{ref}} \), is given by the equation

\[ C_{m,\text{ref}} = \eta_{\text{hom}} \frac{m}{V} P_{\text{rel}}, \]

where \( \eta_{\text{hom}} \) is the aerosol homogeneity in the flow tube, \( m \) is the particulate mass collected on the filter and \( V \) is the sampled volume. \( V \) is given by the aerosol flow through the filter, \( Q \), multiplied by the time duration of the measurement \( t \). \( P_{\text{rel}} \) is defined as the relative particle penetration, \( P_{\text{rel}} = P_{\text{DUT}} / P_{\text{ref}} \), where \( P_{\text{DUT}} \) and \( P_{\text{ref}} \) are the penetration through the sampling probe and connecting tube of the device under testing (DUT) and the reference method, respectively. The associated uncertainties are listed in Table 1.

Since sampling is carried out with isokinetic sampling probes and the tubes leading to the filter holder and the DUT are kept straight and as short as possible, particle losses are minimised. Penetration \( P_{\text{rel}} \) was set to 1; however, an uncertainty of 2% was assigned to account for the higher impaction losses of supermicrometre particles in the sampling funnel of the reference method due to the higher sampling flow (von der Weiden et al., 2009). These losses are to some extent counteracted by the lower diffusion losses of submicrometre particles, which decrease with increasing sampling flow. Here, we followed a rather conservative approach and kept the uncertainty of \( P_{\text{rel}} \) at 2%.

3 Chemical characterisation of model aerosols

Ion chromatography was performed with a Thermo Scientific Dionex™ICS-1500 Ion Chromatography System for analysis of anions and the ICS-2100 model for cations. The systems consist of a liquid eluent, a high-pressure pump, an automatic sample injector, a guard and separator column, an electrolytic suppressor, and a conductivity cell. Before running a sample, the systems were calibrated using a traceable set of calibration standard solutions, which were prepared in-house. The data produced by the range of calibration standard solutions were used to calculate calibration coefficients, which were used to quantitate the sample ions.

Thermo-optical analysis of carbonaceous particles was performed with an OC–EC Analyzer (Lab OC–EC Aerosol Analyzer, Sunset Laboratory Inc., USA), which classified the carbonaceous material as elemental carbon (EC) and organic carbon (OC). The particles were sampled on quartz filters (Advantec, Tokyo, Japan, QR-100, 47 mm). For the analysis, the EUSAAR2 protocol (Cavalli et al., 2010) was modified by extending the last temperature step (850°C) from 80 s in the original protocol to 120 s in order to ensure the complete evolution of carbon (Ess and Vasilatou, 2019). The charring correction for pyrolysed OC was performed by transmittance. OC, EC and TC (total carbon = sum of OC and EC) masses were calculated by the software based on instrument calibration with sucrose solutions.

The elemental composition of the model aerosols was characterised by combining a cascade impactor for PM sampling with Total Reflection X-ray Fluorescence Spectroscopy (TXRF, Bruker TStar S4™, Germany) (Osán et al., 2020). A 13-stage low-pressure cascade impactor (Dekati DLPI 10™, Finland) with particle size range from 30 nm to 10 µm was modified to sample at a rate of 10 L min⁻¹ on smooth and clean commercial-grade acrylic discs with 30 mm diameter, suitable for TXRF. In TXRF, the incident X-ray beam hits the disc’s surface at the total reflection angle. The fluorescence spectrum is detected perpendicular to the surface and is dominated by the contributions from the deposit, i.e. the sampled particles. This allows for the detection of element masses as low as ≈10 to 100 pg and thus short sampling periods. The measured element quantities, combined with the sampled air volume, provide the particle size-selected element mass concentrations in the aerosol. The discs were prepared with a 50 ng yttrium standard for TXRF calibration.

As an example, the TXRF analysis of model aerosol 1 is shown in Fig. 4. The analysis revealed that the mineral dust particles contain primarily the elements Si and Al, and it was assumed that these are present as oxides SiO₂ and Al₂O₃. The mass-based aerodynamic distribution of the SiO₂ particles exhibits a maximum in the range 1–2 µm while the Al₂O₃ particles are larger (≈7 µm). Sulfur (i.e. in the form of sulfate ions) appears predominantly in the submicrometre range (aerodynamic diameter of 30 nm–1 µm), but a second weaker mode is visible at ≈4–7 µm, thus simulating the aerodynamic size distribution of sulfates in ambient air (Wall et al., 1988; Zhuang et al., 1999) reasonably well. The coarse-mode arises most probably from internal mixing of sulfate ions with mineral dust particles. Since nitrates and sulfates were generated with the same method, nitrates are expected to exhibit a similar bimodal size distribution, but this could not be experimentally confirmed since nitrogen is difficult to detect with TXRF spectroscopy. Finally, K⁺ and Cl⁻ ions appear in the micrometre range (>2 µm). It is reasonable to expect that Na⁺ ions also appear in this size range; however, this could not be investigated by TXRF. By comparing the results of ion chromatography with those of TXRF spectroscopy, there is no evidence of insoluble potassium.

The results of the chemical analysis of the model aerosols with ion chromatography, EC/OC analysis and TXRF spectroscopy are summarised in Table 2 and presented graphically in Fig. 5.
ambient-like mixtures; i.e. they contained inorganic salts, facility described in Sect. 2. All three model aerosols were aerosols, which were generated in the laboratory with the algorithms.

and size distribution using optical techniques, from which directly but record instead the particle number concentration and more cost efficient. These do not measure particle mass

air quality monitoring stations. The DustTrak DRX 8533 and mated instruments for monitoring PM mass concentration at

is considered to be one of the most well established auto-

icates using a tapered element oscillating microbalance and TEOM takes continuous direct mass measurements of partic-

USA), a DustTrak DRX 8533 (TSI Inc., USA) and a Fidas Frog (Palas, Germany), were used in this study. The 1405 seems to agree well with the reference method in the...
beginning but indicates a decrease of about 15% in mass concentration at the end of the 4 h measurement. Particle number concentration measurements of the primary aerosols before and after the experiment revealed that the number concentration of the fresh soot particles decreased by about 60% during the measurement period, whereas the number concentration of the dust, salt and aged soot particles remained largely constant. The reason was a defect in the valve regulating the flow of the fresh soot particles into the homogeniser. The decrease in the aerosol mass concentration recorded by the TEOM is therefore real and can be attributed predominantly to the decreasing number and mass concentration of the uncoated soot particles. Since the concentration of the model aerosol decreased during measurement, the best way to assess the performance of the TEOM 1405 with respect to the reference method is to calculate the 4 h average mass concentration. This amounts to 41.6 µg m\(^{-3}\) (see Table 3), only 3.7% lower than the reference measurement (43.2 µg m\(^{-3}\)).

The fresh soot particles consist mainly of EC and have a geometric mean mobility diameter of about 120 nm, i.e. below the cut-off limit of the Fidas Frog. Indeed, experiments with miniCAST soot showed that the Fidas Frog and DustTrak 8533 failed to detect soot particles of this size. This explains why the TEOM reported a constant mass concentration over the whole measurement period. In Table 3, it can be seen that the Fidas Frog reported an average PM\(_{10}\) mass concentration of 38.8 µg m\(^{-3}\), i.e. \(-4.4\) µg m\(^{-3}\) with respect to the reference method. This deviation agrees well with the EC mass concentration of 5.0 µg m\(^{-3}\) (Table 2), as determined with EC/OC analysis. Note that the cut-off curve of optical instruments depends on the refractive index of the particles: the Fidas Frog fails to detect fresh soot particles below \(\approx 200\) nm but detects a considerable mass fraction of the coated soot and salt particles despite their small size.

The results obtained with model aerosol 2 are displayed in Fig. 6b. Here, the concentration of the aerosol remained constant throughout the measurement period. The Fidas Frog and TEOM 1405 monitors underestimate the mass concentration by 29% and 14%, respectively, compared to the reference method, while the DustTrak 8533 overestimates the mass concentration by 50%. The larger deviation between the TEOM 1405 and the reference method compared to model aerosol 1 results from the winter-like environmental conditions; the temperature of model aerosol 2 was set to 12\(^{\circ}\)C, the relative humidity was set to 70% and the nitrate content was relatively high (about 15%) as shown in Table 2. Since the aerosol stream sampled by the TEOM 1405 is heated to 30\(^{\circ}\)C, a fraction of the (semi)volatile components (e.g. nitrate, secondary organic aerosol and water) evolves into the gas phase and is therefore not collected on the filter. These results are in agreement with previous studies reporting that TEOM monitors set at a lower temperature than the standard configuration (50\(^{\circ}\)C) still could lose semivolatile materials (Lee et al., 2005), especially in cooler months (Sofowote et al., 2014; Su et al., 2018).

The large positive deviation of the DustTrak 8533 by a factor of about 1.5 is not surprising. Previous studies have found that different DustTrak models over-recorded PM values by a factor of 1.2–3 (Chung et al., 2001; Grzyb and Lenart-Boron, 2019; Heal et al., 2000; Kingham et al., 2006; Liu et al.,

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**Figure 5.** PM composition (%) of the three model aerosols and environmental conditions during each experiment.

**Table 3.** Average PM\(_{10}\) mass concentration (µg m\(^{-3}\)) reported by the TEOM 1405, Fidas Frog and DustTrak 8533 automated PM monitors and the reference gravimetric method.

| Model aerosol | TEOM 1405 | Fidas Frog | DustTrak 8533 | Reference gravimetric method |
|---------------|-----------|------------|---------------|-----------------------------|
| 1             | 41.6      | 38.8       | –*            | 43.2 ± 2.7                 |
| 2             | 25.3      | 21.0       | 44.0          | 29.4 ± 2.8                 |
| 3             | 19.2      | 15.0       | 25.6          | 19.3 ± 2.2                 |

*The result was discarded because of a technical issue during measurement.*
Figure 6. PM$_{10}$ mass concentrations reported by the TEOM 1405, DustTrak DRX 8533 and Fidas Frog monitors compared to the results of the reference gravimetric method in the case of (a) model aerosol 1, (b) model aerosol 2 and (c) model aerosol 3. In Fig. 6a, the results of the DustTrak 8533 are not plotted because of technical issues during measurement (see text for more details). The dashed lines designate the expanded uncertainties (95% confidence level) of the reference PM$_{10}$ value.

In the case of Fidas Frog, if the reading of the monitor (21.0 µg m$^{-3}$; Table 3) is corrected for the undetected mass of fresh soot (3.8 µg m$^{-3}$; Table 2), then the Fidas Frog still underestimates the mass concentration by $\approx$ 15% with respect to the reference method.

The results obtained in the case of model aerosol 3 are illustrated in Fig. 6c. With an average PM$_{10}$ mass concentration of 19.2 µg m$^{-3}$, the TEOM 1405 exhibits an excellent agreement with the reference method (19.3 µg m$^{-3}$; see Table 2). The DustTrak 8533 overestimates the mass concentration by approx. 33% and thus performs slightly better than in the case of model aerosol 2. Fidas Frog underestimates the mass concentration by about 23%, or $\approx$ 15% after correction for the undetected mass of fresh soot, in agreement with the findings of the experiment with model aerosol 2. As mentioned above, PM monitors based on light scattering, such as the Fidas Frog and the DustTrak, measure particle number concentration and convert this into mass concentration by using a size-dependent particle density function. This function is integrated into the software of the instrument. Deviations may occur if the built-in functions differ substantially from the real density function of the aerosol. Hygroscopic growth of aerosol particles can also lead to considerable measurement artefacts especially when low-cost PM sensors are used (Di Antonio et al., 2018; Crilley et al., 2018). More experiments with ambient-like model aerosols under low and high relative humidity would be needed to define a comprehensive set of calibration factors for these instruments.

5 Conclusions

In this study, we present the first steps towards the generation of ambient-like model aerosols in the laboratory. A custom-made facility (PALMA) for the stable and reproducible generation of such model aerosols was developed, which presents the following advantages:

- The model aerosols are complex, consisting of elemental carbon (fresh soot), soot coated with SOA (aged soot), inorganic ions (such as ammonium, sulfate and nitrate) and mineral dust particles.
- The aerosol mixture can therefore have a controlled amount of semi-volatile and hygroscopic material.
- The total PM mass concentration of the model aerosols can be adjusted in a range from a few micrograms per cubic metre up to about 500 µg m$^{-3}$ and remains stable over several hours.
- The percentage fraction of each PM constituent can be tuned to simulate different urban, suburban or rural aerosols.
- The size distribution (geometric mean and width of accumulation and coarse mode) can be adjusted by tuning the size distribution of the primary aerosols.
- The aerosol temperature and relative humidity can be adjusted to simulate winter-like or summer-like environmental conditions (10–40 °C, 5%–90% RH).

2017; McNamara et al., 2011; Wallace et al., 2011; Yanosky et al., 2002) depending on the aerosol properties. It has been suggested that the “over-estimation is a simple calibration issue in which differences between the optical properties of the manufacturer’s factory calibration PM (Arizona Road Dust) and the PM under study explained the uniform relative errors recorded” (Kingham et al., 2006). The results are nevertheless puzzling. Considering that the device fails completely to detect fresh soot and underestimates the amount of aged soot, we would have rather expected to observe a negative deviation with respect to the reference method. In any case, the large range of the positive systematic bias (factor of 1.2–3) highlights the need for source-specific calibration procedures against a reference method.

In the case of Fidas Frog, if the reading of the monitor (21.0 µg m$^{-3}$; Table 3) is corrected for the undetected mass of fresh soot (3.8 µg m$^{-3}$; Table 2), then the Fidas Frog still underestimates the mass concentration by $\approx$ 15% with respect to the reference method.

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- A spatial aerosol homogeneity of 2.6 % ($k = 2$) in number concentration can be attained in the mixing chamber, a parameter not evaluated so rigorously, if at all, in previous chamber studies (Hogrefe et al., 2004; Liu et al., 2017; Papapostolou et al., 2017; Schwab et al., 2004; Zhu et al., 2007).

- The isokinetic sampling system is highly adaptable and can accommodate instruments with flows up to at least 40 L min$^{-1}$.

- The design is much more compact compared to other mixing chambers described in the literature (Hogrefe et al., 2004; Horender et al., 2019; Papapostolou et al., 2017; Schwab et al., 2004; Zhu et al., 2007) and can therefore easily fit into a typical laboratory.

As a proof of concept, three different automated PM monitors, the TEOM 1405 (Thermo Scientific, USA), the DustTrak DRX 8533 (TSI Inc., USA) and the Fidas Frog (Palas, Germany), were compared with the reference gravimetric method under three different environmental scenarios. The TEOM 1405, operated at 30 °C, agreed very well with the reference gravimetric method in the case of summertime aerosols (21 °C) but showed a negative deviation in PM$\text{10}$ mass concentration of $\approx 15\%$ when the model aerosol was conditioned at 12 °C due to losses of semi-volatile material. The Fidas Frog underestimated the PM$\text{10}$ mass concentration by $\approx 10\%$–$30\%$, whereas the DustTrak 8533 overestimated the PM$\text{10}$ mass concentration by $\approx 30\%$–$50\%$ depending on the aerosol chemical composition and environmental conditions.

Currently, one limitation of the facility is that the model aerosols cannot be conditioned to temperatures lower than 10 °C, but this could be improved by thermally insulating the homogeniser (e.g. with black nitrile foam insulation). Moreover, the composition of the model aerosols could be further refined by adding more components, such as metallic particles with the use of a spark-discharge generator, bioaerosols (e.g. with a Sparging Liquid Aerosol Generator – SLAG, CH Technologies, USA) and particles from biomass burning. This last step could pose challenges since the mass output is usually not very stable over time and the physicochemical properties of the aerosol depend heavily on the combustion material, as well as the stove design.

To conclude, the facility presented in this study can be used to generate ambient-like model aerosols for quality assurance testing, intercomparisons of different instruments, and performance evaluation and calibration with respect to PM mass concentration. The same facility could also be used for other PM measurements such as number concentration and absorption properties (e.g. those related to black carbon). The aerosol facility also provides excellent opportunities for basic aerosol research and aerosol health-related studies.

Data availability. All data presented in the paper are available for research purposes on request to the authors of the paper.

Author contributions. SH and KV designed, validated and operated the experimental facility; coordinated the intercomparison; and prepared the paper with contributions from all other authors. KA designed the isokinetic sampling probes. CCA assisted during the preparation of the intercomparison and DMK performed EC/OC analysis. SIs performed TXRF analysis. PQ helped design the study, TOMS weighed the filter samples and KW performed ion chromatography analysis. FGL advised on aerosol generation. KD performed high-resolution measurements with a reference optical particle counter. SNS operated the DustTrak DRX during the intercomparison.

Competing interests. The authors declare that they have no conflict of interest.

Disclaimer. The opinions expressed and arguments employed herein do not necessarily reflect the official views of the Swiss Government.

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