



Laboratory Validation and Field Deployment of a Compact Single-Scattering Albedo (SSA) Monitor

Julia Perim de Faria¹, Ulrich Bundke¹, *, Andrew Freedman², Timothy B. Onasch², and Andreas Petzold¹ ¹Forschungszentrum Jülich GmbH, IEK-8, 52425 Jülich, Germany

⁵ ²Aerodyne Research, Inc., Billerica, MA 01821-3976, USA *Correspondence to*: Ulrich Bundke (u.bundke@fz-juelich.de)

Abstract. An evaluation of the performance and accuracy of a Cavity Attenuated Phase-Shift Single Scattering Albedo Monitor (CAPS PM_{ssa}, Aerodyne Res. Inc.) was conducted in an optical closure study with proven technologies: Cavity Attenuated Phase-Shift Particle Extinction Monitor (CAPS PM_{ex}, Aerodyne Res. Inc.); 3-wavelengh Integrating

- 10 Nephelometer (TSI Model 3563); and 3-wavelength filter-based Particle Soot Absorption Photometer (PSAP, Radiance Research). The evaluation was conducted by connecting the instruments to a controlled aerosol generation system and comparing the measured scattering, extinction, and absorption coefficients measured by the CAPS PM_{ssa} with the independent measurements. Three different particle types were used to generate aerosol samples with single-scattering albedos (SSA) ranging from 0.4 to 1.0 at 630 nm wavelength. The CAPS PM_{ssa} measurements compared well with the
- 15 proven technologies. Extinction measurement comparisons exhibited a slope of the linear regression line for the full data set of 0.96 (-0.02/+0.06), an intercept near zero, and a regression coefficient $R^2>0.99$; whereas, scattering measurements had a slope of 1.01 (-0.07/+0.06), an intercept of less than +/-2×10⁻⁶ m⁻¹ (Mm⁻¹), and a coefficient $R^2\sim1.0$. The derived CAPS PM_{ssa} absorption compared well to the PSAP measurements at low levels (< 70 Mm⁻¹) for the small particle sizes and modest (0.4 to 0.6) SSA values tested, with a linear regression slope of 1.0, an intercept of -4 Mm⁻¹, and a coefficient
- 20 R^2 =0.97. Comparisons at higher particle loadings were compromised by loading effects on the PSAP filters. For the SSA measurements, agreement was highest (regression slopes within 1%) for SSA = 1.0 particles, though the difference between the measured values increased to 9% for extinction coefficients lower than 55 Mm⁻¹. SSA measurements for absorbing particles exhibited absolute differences up to 18%, though it is not clear which measurement had the lowest accuracy. For a given particle type, the CAPS PM_{ssa} instrument exhibited the lowest scatter around the average. This study demonstrates
- that the CAPS PM_{ssa} is a robust and reliable instrument for the direct measurement of the scattering and extinction coefficients and thus SSA. This conclusion also holds as well for the indirect measurement of the absorption coefficient with the constraint that the accuracy of this particular measurement degrades as the SSA and particle size increases.

Keywords: CAPS PM_{ssa}, optical closure, single scattering albedo.

1 Introduction

- 30 Airborne aerosols impact climate directly though the interaction with incident solar light by scattering, generating a cooling effect, or by absorbing it and reemitting infrared radiation, having a heating effect. According to Haywood and Shine (1995), the effect of aerosols on the atmospheric radiation budget in the visible spectral range depends on the aerosols optical depth (AOD), the single-scattering albedo (SSA), and the backscattered fraction (BF). The radiative forcing efficiency (RFE) describes the resulting aerosol direct forcing per unit AOD (Andrews et al., 2011; Haywood and Shine,
- 35 1995; Sheridan et al., 2012) and is widely used to describing the radiative impact of a given aerosol type. As an aerosol





intensive parameter the RFE value depends only on SSA and BF. As is stated in the latest IPCC report (Boucher et al., 2013), uncertainties in SSA and the vertical distribution of aerosol contribute significantly to the overall uncertainties in the direct aerosol radiative forcing, while AOD and aerosol size distribution are relatively well constrained.

The measurement of SSA requires the simultaneous but independent observation of two parameters since, by 40 definition, the SSA is the ratio of the scattering to the extinction coefficient (where extinction is the sum of the scattering and absorption – see Equation (1) and (2); the index p refers to the contribution of aerosol particles to overall light extinction, which has also a contribution by gas molecules, identified by the index g not shown in the equation).

$$\sigma_{ep} = \sigma_{ap} + \sigma_{sp} \tag{1}$$

$$SSA = \frac{\sigma_{sp}}{\sigma_{ep}}$$
(2)

45 Measuring all three aerosol optical parameters independently allows for the closure of optical properties and thus the determination of uncertainties of the involved instruments.

The aerosol optical parameters are typically measured *in-situ* by instruments such as Integrating Nephelometers (NEPH) for the scattering coefficient (Heintzenberg and Charlson, 1996); photoacoustic (see e.g., Lack et al. (2006); Arnott et al. (2006)) and filter-based methods such as the Particle-Soot Absorption Photometer (PSAP; Bond et al. (1999)), the Multi Angle Absorption Photometer (MAAP; Petzold and Schönlinner (2004)) and more recently the Tricolor Absorption

- Photometer (TAP; Ogren et al. (2017)) for the absorption coefficient; and for the extinction coefficient, the Cavity Ring Down (CRD) technology (Moosmüller et al., 2005) or, since 2007, the Cavity Attenuated Phase Shift Particle Extinction Monitor (CAPS PM_{ex}) (Massoli et al., 2010). To measure the SSA using the optical closure approach involves separate instruments with different principles and uncertainties, leading to potential sources of significant errors and biases.
- 55 A novel instrument based on cavity attenuated phase-shift technology and incorporating an integrating sphere was recently developed by Aerodyne Research, Inc. This novel instrument represents a major step forward in the observation of aerosol optical properties since it simultaneously measures two of the three aerosol optical parameters from the same air sample, reducing the potential sources of sampling biases (Onasch et al., 2015). The two main applications of the CAPS PM_{sss} instrument, apart from the direct measurement of scattering and extinction coefficients, are the indirect measurement
- 60 of the aerosol absorption coefficient and the measurement of the single-scattering albedo. A few recent *in-situ* application studies of the CAPS PM_{ssa} instrument are already available (Corbin et al., 2018; Han et al., 2017). The present optical closure study intends to quantify uncertainties in the measurement of the primary aerosol optical properties and the resulting SSA by the CAPS PM_{ssa} for several types of laboratory aerosol by applying a full set of established instrumentation for measuring the extinction (CAPS PM_{ex}), absorption (PSAP), and scattering (Integrating Nephelometer TELMs del 25(2) estificients at multiple properties.
- 65 TSI Model 3563) coefficients at multiple wavelengths

2 Instruments and Methods

2.1 Instrumental Set-up

The laboratory study was conceived to evaluate the operational principle of the CAPS PM_{ssa} and its performance and accuracy when compared to proven technologies. The instrumental set-up used is shown in Figure 1.

70

50

In this study, similar to previous work (Massoli et al., 2010; Petzold et al., 2013); two collision-type aerosol generators (TSI Model 3076) were used; one containing a solution of deionized water and purely scattering aerosol, Ammonium Sulphate (AS), and a second containing absorbing aerosol, water-soluble colloidal graphite (Aquadag – AD – from Agar Scientific) or Black Carbon (REGAL 400R Pigment Black – BC – from Cabot Corporation). The SSA of the





dispersed aerosol ranged from approximately 0.4 (pure AD or BC) to 1.0 (pure AS), with the modal value of the particle size distribution being below 100 nm in all cases. A drying tube filled with silica gel was positioned after each particle generator in order to reduce the relative humidity below 30%. Once the samples were passed through the dryer, they entered a mixing chamber where effective ensemble particle SSA values of 0.4 < SSA < 1.0 could be produced by mixing aerosol flows containing both absorbing and scattering aerosols. The aerosol generation set-up specifications are shown in Table 1, whereas Table 2 compiles the information about the applied instrument and correction schemes.</p>

80

Three mass flow controllers (MFC), one at each generator's head and a third after the mixing chamber, supplied particle-free compressed air to the sample to both reach the desired humidity and particle number concentration and to make-up the flow required by the instruments. The particle number concentration was measured by a condensation particle counter (CPC).

85 Table 1. Type of generated aerosol, targeted SSA (630 nm), and targeted max. aerosol extinction values

Aerosol type	SSA	Run 1 200 Mm ⁻¹	Run 2 150 Mm ⁻¹	Run 3 100 Mm ⁻¹	Run 4 50 Mm ⁻¹	Run 5 25 Mm ⁻¹
Aquadag (AD)	0.4	Х	Х	х	х	Х
Black Carbon (BC)	0.4		х	Х	х	х
Mixture (AS+AD)	0.6			Х	х	Х
Ammonium Sulphate (AS)	1.0		х	х	х	х



90 Figure 1. Instrumental set-up applied in the optical closure study





Table 2. List and specifications of optical instrumentation	n and applied correction algorithms
---	-------------------------------------

Instrument	Manufacturer	Property	λ (nm)	Aerosol	Correction Algorithm
CAPS PM _{ssa}	Aerodyne	σ_{sp}, σ_{ep}	630	AS, AD,	Mie Amigo (Aerodyne) for σ_{sp} truncation
	Research Inc.			BC, MIX	correction (Onasch et al., 2015)
CAPS PM _{ex}	Aerodyne	σ_{ep}	630	AS, AD,	No correction required
	Research Inc.	-		BC, MIX	
NEPH	TSI Inc.	σ_{sp}	450, 550,	AS	Müller et al. (2009), Anderson and
		-	700		Ogren (1998)
				AD, BC,	Massoli et al. (2009)
				MIX	
PSAP	Radiance	σ_{ap}	467, 530,	AS, AD,	Ogren (2010) and Virkkula (2010)
	Research Inc.	·	660	BC, MIX	

The samples were produced at up to five nominal concentration levels, as shown in Table 1, defined by the aerosol extinction. This was achieved by holding the aerosol generation system constant (MFC#1 and MFC#2) and regulating the make-up air MFCs (MFC#3, MFC#4 and MFC#5). Extinction coefficient levels were varied from ~10 up to 200 Mm⁻¹. For each level, a sampling time of at least 5 minutes was sustained.

To ensure an isoaxial, isokinetic sampling by all instruments, special sampling tips made of stainless steel were designed such that the sample air extraction tips were each concentrically placed along the centre line of the sample tube of 1 inch inner diameter. The inlet nozzles diameters are dimensioned such that the flow velocities in the sample tube and inside extraction tip nozzles match. Distances between the extraction points for the different instruments were 20 cm.

All scattering instruments were calibrated using CO_2 (high span gas) and particle-free air (low span gas), before starting the experiments. This procedure includes also, as recommended by the manufacturers, the calibration of scattering channel of the CAPS PM_{ssa} , against the extinction channel of the instrument. For the filter-based absorption instruments, no calibration is necessary since they both operate with a blank filter in parallel as reference (see description in the subsections below).

105

100

The optical instruments were placed downstream of the generation system, as shown, and will be described in more detail in the following subsections.

2.1.1 Integrating Nephelometer

110

In this optical closure study, an integrating nephelometer (NEPH) of the type TSI Model 3563 was used. The NEPH collects scattering measurements both in the forward and backscatter directions at three wavelengths 450, 550, and 700 nm (Heintzenberg et al., 2006). The NEPH data was corrected for truncation angle effects using the approach proposed by Massoli et al. (2009) for strongly light-absorbing aerosol and the approaches proposed by Anderson et al. (1996) and Müller et al. (2009) for predominantly light-scattering aerosols.

2.1.2 Particle-Soot Absorption Photometer

115 The PSAP is a filter-based three wavelength (467, 530, 660 nm) instrument, manufactured by Radiance Research, that provides continuous measurement of the light absorption coefficient. The instrument uses two spots on a quartz fibre filter; one receives the particle containing sample, and the second clean air. The instrument measures then the difference in the transmission of light between a loaded and a blank filter spot (Bond et al., 1999). Absorption coefficient data were determined using the approach proposed by Ogren (2010).



125

150



120 2.1.3 The CAPS PM_{ex}

The CAPS PM_{ex} system, described in detail and assessed in several studies, such as Massoli et al. (2010), Petzold et al. (2013) and Perim de Faria et al. (2017) measures light extinction by determining the change in signal phase shift caused by the introduction of particles into an optical cavity. The use of high reflectivity mirrors (reflectivity approx. 99.99%) in the optical cavity creates the long measurement path of approx. 2 km required to measure very low values of light extinction (LOD of 1-2 Mm⁻¹ in 1 second sample period).

2.1.4 The CAPS PM_{ssa}

The CAPS PM_{ssa} (Onasch et al., 2015), uses the same principle to measure light extinction as the CAPS PM_{ex}, but it also contains, located at the centre of the measurement cell, a 10 cm diameter integrating sphere capable of measuring light scattering on the same aerosol sample, as shown in Figure 2. The integrating sphere acts as an integrating nephelometer, which measures the scattering of light by particles at all angles, only excluding the near 0 and near 180° angles since at these directions the opening of the extinction chamber is located, allowing the sample and light beam to pass through. The sphere shows 98-99% Lambertian reflectance efficiency due to its high reflectivity coating (Avian D from Avian Technologies). The usage of an integrating sphere increases the collection of scattered light at the photomultiplier compared to a traditional cosine corrected detector arrangement.

- 135 The scattering channel is calibrated against the extinction channel using small particles (<250 nm) that have SSA=1.0, in this case ammonium sulphate, and set equal to the extinction measurement. Thus, the monitor should be thought of as providing separate extinction and SSA values with the scattering channel a derived measurement. This calibration procedure also allows the user to prove monitor linearity over a wide range of optical extinctions without the limitation of using individual gases with sometimes not particularly well-known Rayleigh scattering coefficients.
- 140 As in the CAPS PM_{ex} , the sample flow is set to 0.85 lpm and is controlled by a critical orifice. The measurement sample enters the chamber in one end and exits through an opening located in the other end flowing through a glass tube inside the integrating sphere (Figure 2). The mirrors are kept particle-free by a continuously flowing purge flow (25 cm³ min⁻¹).



145 Figure 2. CAPS PMssa components and set-up (Onasch et al., 2015).

The baseline determination system is identical to the one used in the CAPS PM_{ex} , in which filtered and thus particle-free sample air fills the measurement chamber and is used to quantify contributions of gas molecules to the instrument response by Rayleigh scattering and potential absorption of light, and to determine interferences of system components. Both the CAPS PM_{ex} and CAPS PM_{ssa} used in this study operate at a wavelength of 630 nm and thus show minimal interference from absorption by ambient gaseous species like NO₂ and H₂O.





2.2 Data Treatment

All multi-wavelength instruments were adjusted to match the other instruments' wavelengths for the intercomparison by using the Ångström exponent approach; see Equation (3) and (4),

155

$$\mathring{a} = -\frac{\log \frac{\sigma_x}{\sigma_y}}{\left/\log \frac{x}{v}\right|}$$
(3)

$$\sigma_w = \sigma_y \times (^W/_y)^{-a} \tag{4}$$

where å is the Ångström exponent, σ is the optical property measured (extinction, scattering or absorption coefficient), x and y are the operating wavelengths of the instrument, and w refers to the wavelength, to which the property should be adjusted. For a better understanding of the wavelength adjustment, the complete description is given in Figure 3 from Petzold et al. (2013).

All instruments provide 1 second resolution data. Data was collected over 5 minutes for each experimental point to remove any effect of differences in response times and fluctuations in the aerosol generation system. The data was averaged for each extinction/scattering/absorption level, and the standard deviation was calculated from the mean.

165

Standard linear regression analysis was performed for the mean values of each level. For the cases with the standard deviation of the intercept value being higher than the value itself, the regression model interception was forced to zero intercept, since the intercept value shows no significant difference to zero.

3 Results and Discussion

In this section, we present the results and relevant discussion of findings for the optical closure study. All the measurements presented here were corrected to the CAPS PM_{ssa} operational wavelength of 630 nm.

3.1 Extinction Coefficient

The extinction coefficient measured by the CAPS PM_{ssa} was analysed in comparison with proven technologies. On the direct measurement of σ_{ep}, we compared the two CAPS systems for AS and AD (Petzold et al., 2013). The direct measurement of σ_{ep} from the CAPS PM_{ssa} was also compared with the indirect measurement given by the sum of the absorption coefficient measured by the PSAP with the scattering coefficient measured by the NEPH for BC, AD, and MIX (as defined in Table 1). For AS with the measured SSA value of 1.0, extinction coefficients provided by the CAPS extinction channels and scattering coefficients provided by the CAPS scattering channel and the NEPH instrument are used for the evaluation of the light scattering measurements in the next subsection. The time series for the extinction channels are shown in Figure 3 and the averages and standard deviations for each test point are shown in Table A1 in the

supplemental information. The higher variability observed in the last plot of the figure is due to particle load fluctuations

180

from_generation system when operating at very high loads.







185 Figure 3. Time series of the measurements by the extinction channel.

Figure 4 shows the scatter plot of the measured extinction coefficient for the two CAPS systems for AD and AS and the comparison with the sum of the NEPH and PSAP for AD and BC. The best results for the AD and BC were found when applying the Massoli et al. (2009) correction with the assumption, that no particle size cut has been used for the inlet system (no-cut approach) to the NEPH data, and Virkkula (2010) for strongly light-absorbing aerosols AD and BC to the PSAP data. For the mixture, the applied corrections were Anderson et al. (1996) for the NEPH data and Ogren (2010) for the PSAP data. The extinction channels from the two CAPS and the sum of the NEPH and PSAP (PSAP-NEPH) signals show a good agreement for all aerosol types, with linear regression slopes (m) between 0.94 and 1.02 and correlation coefficients above 0.99 (all regression analysis data for the averaged values of each level is presented in Table 3 together
with their standard deviation). For the linear regression analysis of the full data set including all types of aerosols, the slope found was 0.96 (R²=0.99) for the comparison of the CAPS PM_{ssa} extinction data. The slopes of the regression analysis and their standard deviation are shown in Figure 5 as a function of the sampled aerosol single-scattering albedo. As it can be seen there is no systematic difference in the slope with increase or decrease of the aerosol SSA.





Table 3 Linear regression parameters including standard deviation of the mean, intercept, standard intercept, and R^2 for the comparison of the CAPS PM_{ssa} extinction channel with proven technologies

Aerosol	Reference Instrument	SSA	М	Std m	В	Std b	\mathbb{R}^2
AD	PSAP-NEPH	0.4	0.94	0.01	0.00	< 0.01	1.00
BC	PSAP-NEPH	0.4	1.00	0.01	0.00	< 0.01	1.00
MIX	PSAP-NEPH	0.6	1.02	0.00	0.00	< 0.01	1.00
ALL	PSAP-NEPH	NA	0.96	0.01	0.00	< 0.01	0.99
AD	CAPS PM _{ex}	0.4	0.95	0.00	0.00	< 0.01	1.00
AS	CAPS PM _{ex}	1.0	1.00	0.00	0.00	< 0.01	1.00
ALL	CAPS PM _{ex}	NA	0.97	0.00	0.00	< 0.01	1.00

It is worth noting that for the particular instruments used in our study, the standard deviation for the extinction 205 data of the CAPS PM_{ssa} is larger than for the extinction data provided by the CAPS PM_{ex} (horizontal error bars). This finding is shown in the histogram of the extinction channel from one measurement level (in this case the used dataset refers to the 25 Mm⁻¹ target-level for AD aerosol) for both equipment (Figure 6). Thus, the precision of this particular CAPS PM_{ssa} is lower than the precision of the CAPS PM_{ex}. Regarding the precision of the CAPS PM_{ssa} in comparison with proven technologies, the standard deviation found in this study for both cases are comparable. The precision in the CAPS PM_{ex} and 210 PSAP-NEPH extinction measurements found in this study are very similar to the one found by Petzold et al. (2013).





Figure 4. Comparison result of the extinction channel of the CAPS PM_{ssa} with the CAPS PM_{ex} and the PSAP-NEPH for for the different aerosol types.





Figure 6 Frequency of extinction coefficient measurement for the CAPS PM_{ssa} and PM_{ex} systems at the nominal 25 Mm^{-1} (level 5) test point for AD.





3.2 Scattering Coefficient

215

The scattering channel of the CAPS PM_{ssa} was evaluated in comparison to the NEPH measurements for AD, BC, AS, and MIX (Table 1). The time series of scattering coefficient data for the various aerosol runs is shown in Figure 7. Supplemental Table A2 shows the average and 1- σ standard deviation obtained for the targeted scattering coefficient levels. There is no systematic error found neither in the average nor in the standard deviation of the measured values. The precision of both instruments for the measurement of scattering coefficient is very similar.



220 Figure 7. Time series of the measurements by the scattering channel.

Figure 8 shows the scatter plot of the 1-second average and standard deviation of the CAPS PM_{ssa} against NEPH. As it can be seen from Figure 8 and the data compiled in Table 4, the agreement with the NEPH measurements is excellent, with less than 8% difference in the slope, offset smaller than 2.00 Mm⁻¹ and correlation coefficient of 1.00 for all aerosol
types. The slope value and standard deviation as a function of SSA is shown in Figure 9. For the AD, BC and Mix cases, the NEPH data was corrected with the Massoli et al. (2009) approach. For the AS case both the Anderson et al. (1996) and Müller et al. (2009) were applied and the results given were practically the same, less than 2% in the slope and less than 1.00 Mm⁻¹ difference in the offset. For the overall measurement linear regression model, including all types of aerosols, the slope found was 1.01 (R²=1.00) for the comparison of the CAPS PM_{ssa} with the NEPH.

Table 4. Linear regression parameters including standard deviation of the mean, intercept, standard intercept, and R^2 for the comparison of the CAPS PM_{ssa} scattering channel with NEPH

Aerosol	Reference Instrument	SSA	m	Std m	b	Std b	R^2
AS	NEPH	1.00	1.02	0.00	-0.72	0.14	1.00
AD	NEPH	0.40	0.98	0.00	1.48	0.18	1.00
BC	NEPH	0.40	0.94	0.01	1.22	0.28	1.00
MIX	NEPH	0.60	1.07	0.01	-0.55	0.50	1.00
ALL	NEPH	NA	1.01	0.01	0.00	0.00	1.00









Figure 8. Comparison result of the scattering channel of the CAPS $\rm PM_{ssa}$ with the measurements from the NEPH for the different aerosol types.

Figure 9. Slope values of the linear regression as a function of expected aerosol SSA for CAPS PM_{ssa} and NEPH; uncertainty of the slopes is below the resolution of the symbols; see Table 4.

235 3.3 Absorption Coefficient

In spite of the fact that the CAPS PM_{ssa} is not capable of directly measuring the absorption coefficient, the values can be derived as the difference of the extinction and the scattering coefficients; see Equation (1). From the difference of the two CAPS PM_{ssa} channels the calculated absorption coefficients were compared to the direct measurement by the PSAP. In this analysis, when operating with a mixture of AS and AD, the PSAP data were treated using the correction from Ogren (2010). The time series for the measurement of the different aerosols are shown in Figure 10 whereas Supplemental Table

240 (2010). The time series for the measurement of the different aerosols are shown in Figure 10 whereas Supplemental T A3 shows the average and $1-\sigma$ standard deviation obtained for the targeted absorption coefficient levels.



245 Figure 10. Time series of the measurements by the absorption channel.





The scatter plot for the average measured values from both methods for all levels is shown in Figure 11, whereas the results of the linear regression analysis are compiled in Table 5. The agreement between the methods is good, with deviations below 11% in the slope, and offsets less than 2.0 Mm^{-1} . The correlation coefficient is above 0.98 for all cases.

- For the full data set of CAPS PM_{ssa} and PSAP absorption coefficient data including all types of aerosols, the slope is 0.91 with a correlation coefficient of R^2 =0.98. Figure 11 demonstrates that for higher absorption coefficients, the two methods deviate more strongly than for lower absorption coefficients. This is mainly caused by the correction algorithm applied to the PSAP data (also seen on Figure 10); filter loading corrections are significantly larger for higher absorption coefficient levels than for lower absorption coefficient levels. If the three data points for higher absorption coefficient data ($\sigma_{ap} > 70$)
- 255 Mm^{-1}) are removed from the regression analysis, the slope value increases to 1.00 (R^2 =0.97), although with an offset of -3.64. This finding proves that, although the CAPS PM_{ssa} cannot directly measure aerosol light absorption, it provides a rather reliable measurement of the absorption coefficient of the sampled aerosol, at least for the small particle sizes and intermediate SSA values sampled in this study. The accuracy of absorption measurements by the two channels of the CAPS PM_{ssa} may be significantly reduced for weakly absorbing but large-sized and irregularly shaped mineral dust particles.

Table 5. Linear regression parameters including standard deviation of the mean, intercept, standard intercept, and R^2 for the comparison of the CAPS PM_{ssa} and the PSAP instruments.

Aerosol	Reference Instrument	m	Std m	b	Std b	R^2
AD	PSAP	0.89	0.01	0.00	0.00	1.00
BC	PSAP	0.90	0.00	0.00	0.00	0.99
MIX	PSAP	1.02	0.04	2.02	1.16	0.99
ALL	PSAP	0.91	0.02	0.00	0.00	0.98
$ALL \; (\sigma_{ap} < 70 \; Mm^{-1})$	PSAP	1.00	0.07	-3.64	2.33	0.97



265 Figure 11. Comparison result of the absorption indirect measurement by the CAPS PM_{ssa} with the measurements from the PSAP for AD, BC and Mixture.





3.4 Single Scattering Albedo Measurement

The ultimate property targeted by the CAPS PM_{ssa} is the aerosol single-scattering albedo. Figure 12 shows the average and standard deviation of the SSA measured by the CAPS PM_{ssa} and the applied proven technologies for each aerosol type containing a light-absorbing fraction, at the different extinction coefficient levels. The values for each level are also





Figure 12. Average and standard deviation of the measured Single Scattering Albedo as a function of extinction coefficient level for the different aerosols and technologies.

275

280

For the absorbing aerosols, we found maximum deviations between the different SSA values of 0.09, or 18%, with the deviations being randomly distributed around zero. For a single aerosol type, the SSA provided by the CAPS PM_{ssa} shows less scatter around the average value compared to the values derived from PSAP and NEPH data. The measurements by the CAPS PM_{ssa} are more robust in terms of stability in comparison with the values measured by the PSAP-NEPH combination, with an average of the standard deviation for the different aerosol types of 0.01 for the CAPS PM_{ssa} and 0.02 for the PSAP-NEPH combination. It is worth noting that even though there are differences found in the measurements, all measured SSA values fall within the range of values expected for each aerosol type.

Analysing the error propagation for the measured parameters (extinction and scattering coefficients), the increase of the uncertainty at the lower extinction coefficient levels is also visible for both CAPS PM_{ssa} and proven technologies; see Table 6 for details. From the experimental set-up, it was observed that the particle generation system was lightly unstable when operating at lower extinction/scattering levels, resulting in higher variations of the absolute values, which could explain such higher error propagation. This supports the previous findings that the CAPS PM_{ssa} accuracy is very good and comparable to the proven technologies.

290 Table 6. Absolute uncertainty of the SSA measurement for given aerosol types and applied instrument combinations





Aerosol	Instrument	Run 1	Run 2	Run 3	Run 4	Run 5
AS	CAPS PM _{ssa}	0.04	0.07	0.14	0.19	NA
AD	CAPS PM _{ssa}	0.05	0.05	0.14	0.34	0.46
nii)	PSAP-NEPH	0.06	0.06	0.14	0.38	0.45
BC	CAPS PM _{ssa}	0.05	0.07	0.11	0.17	NA
20	PSAP-NEPH	0.05	0.07	0.11	0.16	NA
Mix	CAPS PM _{ssa}	0.22	0.11	0.07	NA	NA
IVIIX	PSAP-NEPH	0.25	0.11	0.06	NA	NA

4 Summary and Outlook

An optical closure study has been performed using different types of aerosols (pure scattering, strongly absorbing, and mixture) to evaluate the performance and accuracy of the recently launched Cavity Attenuated Phase-Shift Single Scattering Albedo Monitor.

295

300

305

The results from the instrument intercomparison with proven technologies (CAPS PM_{ex} , NEPH, and PSAP) show a very good agreement for all aerosol types, with accuracy of 96% and 99% for the extinction coefficient and scattering coefficient channels, respectively, for all aerosol types. The small deviation of 4% observed in the extinction channel between the CAPS PM_{ssa} and PSAP-NEPH combination originates from the applied correction algorithm to the PSAP data, since it is a logarithmic function of the filter transmission leading to deviations in the dataset. For the evaluation of the performance for each aerosol individually, the extinction channel shows accuracy between 94% and 98%; and the scattering channel, between 94% and 98%. These values are very similar to those found by Petzold et al. (2013) for the CAPS PM_{ex} .

Regarding the application of the CAPS PM_{ssa} for the measurement of the absorption coefficient and singlescattering albedo, the instrument has shown good performance on both sides. The accuracy of the absorption coefficient

measurement by the CAPS PM_{ssa} in comparison with the PSAP was 91%, as obtained for the linear regression analysis for all investigated aerosol types and aerosol loadings. The large difference observed here comes from the correction scheme applied to the PSAP data at high loadings, as stated earlier. It is possible to observe that the higher deviations occur at high absorption coefficient, also where the transmission of the filter has a steeper decrease. Once the linear regression analysis excludes the points where the average absorption coefficient was higher than 70 Mm⁻¹, the slope approaches 100%

310 agreement between the two technologies. For the measurement of SSA, the CAPS PM_{ssa} showed a very good stability for all measured σ_{ep} levels, better than the PSAP-NEPH combination. The measured values are within what is expected for the different types of aerosols (0.4 for strongly absorbing aerosols and 1.0 for purely scattering aerosols).

The results reported from our study demonstrate that the CAPS PM_{ssa} is a very robust and reliable instrument for the direct measurement of the scattering and extinction coefficient, as well as for the indirect measurement of the 315 absorption coefficient and single scattering albedo.

5 Author Contributions

JP, UB, and AP designed the study and prepared the manuscript, with contributions from all co-authors. AF and TO provided technical details of the instrumentation and contributed to the interpretation of the study results.





6 Competing Interests

320 The authors declare that they have no conflict of interest.

7 Acknowledgements

Parts of this work was funded by the EU FP7 project IGAS (Grant Agreement No. 312311), the Federal Ministry of Education and Research, Germany, in IAGOS D (Grant Agreement No. 01LK1301A), EU H2020 Project ENVRIPUS (Grant No. 654182) and HITEC Graduate School for Energy and Climate.

325 8 References

365

Anderson, T. L., Covert, D. S., Marshall, S. F., Laucks, M. L., Charlson, R. J., Waggoner, A. P., Ogren, J. A., Caldow, R., Holm, R. L., Quant, F. R., Sem, G. J., Wiedensohler, A., Ahlquist, N. A., and Bates, T. S.: Performance characteristics of a high-sensitivity, three-wavelength, total scatter/backscatter nephelometer, Journal of Atmospheric and Oceanic Technology, 13, 967-986, 1996.

- 330 Anderson, T. L. and Ogren, J. A.: Determining aerosol radiative properties using the TSI 3563 integrating nephelometer, Aerosol Science and Technology, 29, 57-69, 1998. Andrews, E., Ogren, J. A., Bonasoni, P., Marinoni, A., Cuevas, E., Rodriguez, S., Sun, J. Y., Jaffe, D. A., Fischer, E. V., Baltensperger, U., Weingartner, E., Coen, M. C., Sharma, S., Macdonald, A. M., Leaitch, W. R., Lin, N. H., Laj, P., Arsov, T., Kalapov, I., Jefferson, A., and Sheridan, P.: Climatology of aerosol radiative properties in the free troposphere, Atmos
- 335 Res, 102, 365-393, 2011. Arnott, W. P., Walker, J. W., Moosmuller, H., Elleman, R. A., Jonsson, H. H., Buzorius, G., Conant, W. C., Flagan, R. C., and Seinfeld, J. H.: Photoacoustic insight for aerosol light absorption aloft from meteorological aircraft and comparison with particle soot absorption photometer measurements: DOE Southern Great Plains climate research facility and the coastal stratocumulus imposed perturbation experiments, Journal of Geophysical Research-Atmospheres, 111, 2006.
- 340 Bond, T. C., Anderson, T. L., and Campbell, D.: Calibration and intercomparison of filter-based measurements of visible light absorption by aerosols, Aerosol Science and Technology, 30, 582-600, 1999. Boucher, O., Randall, D., Artaxo, P., Bretherton, C., Feingold, G., Forster, P., Kerminen, V.-M., Kondo, Y., Liao, H., Lohmann, U., Rasch, P., Satheesh, S. K., Sherwood, S., Stevens, B., and Zhang, X. Y.: Clouds and Aerosols. In: Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the
- 345 Intergovernmental Panel on Climate Change, Stocker, T. F., Qin, D., Plattner, G.-K., Tignor, M., Allen, S. K., Boschung, J., Nauels, A., Xia, Y., Bex, V., and Midgley, P. M. (Eds.), Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 2013. Corbin, J. C., Pieber, S. M., Czech, H., Zanatta, M., Jakobi, G., Massabo, D., Orasche, J., El Haddad, I., Mensah, A. A.,
- Stengel, B., Drinovec, L., Mocnik, G., Zimmermann, R., Prevot, A. S. H., and Gysel, M.: Brown and Black Carbon Emitted
 by a Marine Engine Operated on Heavy Fuel Oil and Distillate Fuels: Optical Properties, Size Distributions, and Emission Factors, Journal of Geophysical Research-Atmospheres, 123, 6175-6195, 2018.
 Uner, T. T., Xu, W. O., Li, L. Fraedree, A. Zhen, L. Wang, O. O. Chen, C. Zheng, Y. L. Wang, Z. F. Fu, B. O., Lin, X.
- Han, T. T., Xu, W. Q., Li, J., Freedman, A., Zhao, J., Wang, Q. Q., Chen, C., Zhang, Y. J., Wang, Z. F., Fu, P. Q., Liu, X. G., and Sun, Y. L.: Aerosol optical properties measurements by a CAPS single scattering albedo monitor: Comparisons between summer and winter in Beijing, China, Journal of Geophysical Research-Atmospheres, 122, 2513-2526, 2017.
- Haywood, J. M. and Shine, K. P.: The effect of anthropogenic sulfate and soot aerosol on the clear-sky planetary radiation budget, Geophysical Research Letters, 22, 603-606, 1995.
 Heintzenberg, J. and Charlson, R. J.: Design and applications of the integrating nephelometer: A review, Journal of Atmospheric and Oceanic Technology, 13, 987-1000, 1996.
- Heintzenberg, J., Wiedensohler, A., Tuch, T. M., Covert, D. S., Sheridan, P., Ogren, J. A., Gras, J., Nessler, R., Kleefeld,
 C., Kalivitis, N., Aaltonen, V., Wilhelm, R. T., and Havlicek, M.: Intercomparisons and aerosol calibrations of 12 commercial integrating nephelometers of three manufacturers, Journal of Atmospheric and Oceanic Technology, 23, 902-914, 2006.
 Lether D. A., Leuriser, F. B., Bernerd, T., Pitterner, A., and Pericherland, A. B., Assential characteristic measurement using a statement of the statement

Lack, D. A., Lovejoy, E. R., Baynard, T., Pettersson, A., and Ravishankara, A. R.: Aerosol absorption measurement using photoacoustic spectroscopy: Sensitivity, calibration, and uncertainty developments, Aerosol Science and Technology, 40, 697-708, 2006.

Massoli, P., Kebabian, P. L., Onasch, T. B., Hills, F. B., and Freedman, A.: Aerosol Light Extinction Measurements by Cavity Attenuated Phase Shift (CAPS) Spectroscopy: Laboratory Validation and Field Deployment of a Compact Aerosol Particle Extinction Monitor, Aerosol Science and Technology, 44, 428-435, 2010.





Massoli, P., Murphy, D. M., Lack, D. A., Baynard, T., Brock, C. A., and Lovejoy, E. R.: Uncertainty in Light Scattering
 Measurements by TSI Nephelometer: Results from Laboratory Studies and Implications for Ambient Measurements,
 Aerosol Science and Technology, 42, 1064-1074, 2009.

Moosmüller, H., Varma, R., and Arnott, W. P.: Cavity ring-down and cavity-enhanced detection techniques for the measurement of aerosol extinction, Aerosol Science and Technology, 39, 30-39, 2005.

 Müller, T., Nowak, A., Wiedensohler, A., Sheridan, P., Laborde, M., Covert, D. S., Marinoni, A., Imre, K., Henzing, B.,
 Roger, J.-C., dos Santos, S. M., Wilhelm, R., Wang, Y.-Q., and de Leeuw, G.: Angular Illumination and Truncation of Three Different Integrating Nephelometers: Implications for Empirical, Size-Based Corrections, Aerosol Science and Technology, 43, 581-586, 2009.

Ogren, J. A.: Comment on Calibration and Intercomparison of Filter-Based Measurements of Visible Light Absorption by Aerosolso, Aerosol Science and Technology, 44, 589-591, 2010.

- Ogren, J. A., Wendell, J., Andrews, E., and Sheridan, P. J.: Continuous light absorption photometer for long-term studies, Atmospheric Measurement Techniques, 10, 4805-4818, 2017.
 Onasch, T. B., Massoli, P., Kebabian, P. L., Hills, F. B., Bacon, F. W., and Freedman, A.: Single Scattering Albedo Monitor for Airborne Particulates, Aerosol Science and Technology, 49, 267-279, 2015.
- Perim de Faria, J., Bundke, U., Berg, M., Freedman, A., Onasch, T. B., and Petzold, A.: Airborne and laboratory studies of
 an IAGOS instrumentation package containing a modified CAPS particle extinction monitor, Aerosol Science and
 Technology, doi: 10.1080/02786826.2017.1355547, 2017. 1-14, 2017.

Petzold, A., Onasch, T., Kebabian, P., and Freedman, A.: Intercomparison of a Cavity Attenuated Phase Shift-based extinction monitor (CAPS PMex) with an integrating nephelometer and a filter-based absorption monitor, Atmospheric Measurement Techniques, 6, 1141-1151, 2013.

- 390 Petzold, A. and Schönlinner, M.: Multi-angle absorption photometry—a new method for the measurement of aerosol light absorption and atmospheric black carbon, Journal of Aerosol Science, 35, 421-441, 2004. Sheridan, P. J., Andrews, E., Ogren, J. A., Tackett, J. L., and Winker, D. M.: Vertical profiles of aerosol optical properties over central Illinois and comparison with surface and satellite measurements, Atmospheric Chemistry and Physics, 12, 11695-11721, 2012.
- 395 Virkkula, A.: Correction of the Calibration of the 3-wavelength Particle Soot Absorption Photometer (3 PSAP), Aerosol Science and Technology, 44, 706-712, 2010.





9 Appendix A

400

SUPPLEMENTAL INFORMATION

Table A1. Extinction coefficient mean and $1-\sigma$ standard deviation of the mean measured by the CAPS PM_{ssa} extinction channel and proven technologies

			Run 1	Run 2	Run 3	Run 4	Run 5
	CADE DM	Av	54.62	127.43	311.65	198.31	NA
	CAPS PM _{ssa}	Std	0.29	0.66	1.04	1.50	NA
AS		Av	53.39	124.78	306.40	195.94	NA
	CAPS PM _{ex}	Std	0.21	0.41	0.68	1.01	NA
	CADS DM	Av	221.04	105.98	66.16	26.25	8.84
\sim	CAFS FM _{ssa}	Std	1.34	0.23	0.22	0.20	0.08
AI	CAREDM	Av	210.15	100.22	63.08	24.93	8.66
	CAPS PM _{ex}	Std	1.53	0.22	0.16	0.14	0.05
	CAPS PM	Av	198.00	150.09	104.15	56.88	28.85
\sim	CAI 5 I M _{ssa}	Std	0.20	0.17	0.39	0.53	0.37
AI	DCAD NEDU	Av	187.37	135.55	102.30	51.34	26.78
	PSAP-NEPH	Std	0.33	0.23	0.36	0.61	0.34
	CADS DM	Av	136.77	76.16	50.99	27.73	NA
٢)	CAI 5 I M _{ssa}	Std	0.26	0.20	0.22	0.13	NA
B(DCAD NEDU	Av	134.98	81.59	48.51	26.28	NA
	PSAP-NEPH	Std	0.22	0.18	0.16	0.29	NA
	CADS DM	Av	23.05	63.14	100.94	NA	NA
×	CAI 5 I M _{ssa}	Std	0.17	0.25	0.20	NA	NA
Mi	DCAD NEDU	Av	21.28	58.47	90.83	NA	NA
	PSAP-NEPH	Std	0.19	0.23	0.18	NA	NA





Table A2. Scattering coefficient mean and $1-\sigma$ standard deviation of the mean measured by the CAPS
PM _{ssa} and NEPH

			Run 1	Run 2	Run 3	Run 4	Run 5
	CADE DM	Av	131.79	92.57	54.29	12.31	NA
S	CAPS PM _{ssa}	Std	0.11	0.16	0.08	0.06	NA
A	NEDU	Av	133.22	93.22	54.18	11.77	NA
	NEPH	Std	0.11	0.10	0.08	0.04	NA
	CADS DM	Av	78.29	59.42	41.18	21.98	10.32
D	CAPS PlvI _{ssa}	Std	0.11	0.10	0.16	0.22	0.15
A	NEDU	Av	78.50	59.86	41.70	22.93	11.87
	NEI II	Std	0.12	0.12	0.17	0.22	0.17
	CADS DM	Av	54.33	30.54	20.58	10.66	NA
U	CAFS FM _{ssa}	Std	0.14	0.11	0.11	0.08	NA
B	NEDLI	Av	52.71	29.81	20.91	11.31	NA
	NEFT	Std	0.14	0.11	0.12	0.08	NA
	CADS DM	Av	11.66	32.52	51.09	NA	NA
ix.	CAPS PM _{ssa}	Std	0.11	0.14	0.14	NA	NA
Σ	NEDH	Av	11.32	34.05	54.43	NA	NA
	NEPH	Std	0.11	0.14	0.12	NA	NA

Table A3. Absorption coefficient mean and 1- σ standard deviation of the mean measured by the CAPS PM_{ssa} (extinction minus scattering) and PSAP

			Run 1	Run 2	Run 3	Run 4	Run 5
		Av	78.69	43.78	29.73	16.57	NA
٢)	CAPS PM _{ssa}	Std	0.18	0.13	0.14	0.09	NA
B(56.15	Av	70.13	44.27	23.85	12.74	NA
	PSAP	Std	0.19	0.16	0.12	0.09	NA
	G + DG D) (Av	119.75	90.76	62.02	25.40	18.53
\circ	CAPS PM _{ssa}	Std	0.14	0.13	0.24	0.32	0.23
Ρ	DGAD	Av	108.92	75.97	60.09	20.16	14.92
	PSAP	Std	0.31	0.19	0.23	0.40	0.20
		Av	10.09	26.09	42.44	NA	NA
×	CAPS PM _{ssa}	Std	0.10	0.16	0.11	NA	NA
Mi	DG I D	Av	11.95	29.42	45.03	NA	NA
	PSAP	Std	0.18	0.17	0.14	NA	NA





Table A4. Single Scattering Albedo average value and standard deviation for CAPS PM_{ssa} and proven technologies

	Scat/Ext		Run 1	Run 2	Run 3	Run 4	Run 5
70	CADE DM	Av	0.99	0.99	1.01	1.09	NA
A	CAPS PM _{ssa}	Std	0.02	0.02	0.03	0.11	NA
	CADS DM	Av	0.40	0.40	0.40	0.39	0.36
\circ	CAFS FIM _{ssa}	Std	0.01	0.01	0.01	0.02	0.05
AI	DCAD NIEDH	Av	0.42	0.44	0.41	0.45	0.44
	PSAP-NEPH	Std	0.02	0.02	0.02	0.04	0.07
	CADS DM	Av	0.40	0.40	0.40	0.38	NA
٢)	CAI 5 I M _{ssa}	Std	0.01	0.02	0.02	0.04	NA
B(DSAD NEDH	Av	0.39	0.37	0.43	0.43	NA
	I SAI -NEI II	Std	0.02	0.02	0.03	0.05	NA
	CAPS PM	Av	0.51	0.52	0.51	NA	NA
Mix	CAI 5 I W _{ssa}	Std	0.06	0.03	0.02	NA	NA
	PSAP_NEPH	Av	0.53	0.58	0.60	NA	NA
	I SAI -NEI II	Std	0.13	0.05	0.04	NA	NA