Intercomparison of Photoacoustic and Cavity Attenuated Phase Shift Instruments: Laboratory Calibration and Field Measurements

Jialuo Zhang¹, Jun Chen¹*, Meng Wang¹, Mingxu Su¹, Wu Zhou¹, Ravi Varma², Shengrong Lou¹,³*

¹Shanghai Key Laboratory of Multiphase Flow and Heat Transfer in Power Engineering, School of Energy and Power Engineering, University of Shanghai for Science and Technology, Shanghai 200093, China
²Department of Physics, National Institute of Technology Calicut, Calicut 673601, Kerala, India
³State Environmental Protection Key Laboratory of the Cause and Prevention of Urban Air Pollution Complex, Shanghai Academy of Environmental Science, Shanghai 200070, China

Correspondence to: Jun Chen (j.chen@usst.edu.cn)
Shengrong Lou (lousr@saes.sh.cn)

Abstract: The study of aerosol optical properties is essential to understand its impact on the global climate. In our recent field measurement carried out in the Gehu area of southwest Changzhou City, a photoacoustic extinctiometer (PAX) and a cavity attenuated phase shift albedo monitor (CAPS-ALB) were used for online aerosol optical properties measurement. Laboratory calibration with gas and particle samples were carried out to correct disagreements of field measurements. During particle calibration, we adopted ammonium sulfate (AS) samples for scattering calibration of nephelometer parts of both the instruments, then combined with number-size distribution measurements into MIE model for calculating the value of the total scattering (extinction) coefficient. During gas calibration, we employed high concentration NO₂ for absorption calibration of PAX resonator, then further intercompared the extinction coefficient of CAPS-ALB with a cavity-enhanced spectrometer. The correction coefficient obtained from the laboratory calibration experiments was employed on the optical properties observed in the field measurements correspondingly, and showed good result in comparison with reconstructed extinction from the IMPROVE model. The intercomparison of the calibrated optical properties of PAX and CAPS-ALB in field measurement were in good agreement with slopes of 1.052, 1.024 and 1.046 for extinction, scattering and absorption respectively, which shows the reliability of measurement results and verifies the correlation between the photoacoustic and the cavity attenuated phase shift instruments.

Key words: Aerosol optical properties, instrument calibration, photoacoustic spectroscopy, cavity attenuated phase shift spectroscopy

1 Introduction

Atmospheric aerosols can directly affect the earth’s energy balance and cause global temperature changes by absorbing and scattering solar radiation(Horvath, 1993;Haywood and Shine, 1995;Penner et al., 2001). Therefore, considerable studies were undertaken to investigate the optical properties of aerosol particles from different regions(Baynard et al., 2007;Petzold...
et al., 2013; Moosmüller et al., 1998). The optical properties of regional aerosols depend on particle size distribution, mixing state and complex refractive index, thus online measurements are necessary (Nakayama et al., 2015; Schwartz et al., 2010). Furthermore, the calibration of instruments is a key step to ensure the reliability and quality of online measurement data of aerosol optical properties.

Ideally, the complete set of aerosol optical properties are required measuring simultaneously, including aerosol extinction, scattering and absorption coefficients, for aerosol optical closure studies, where the aerosol extinction coefficient is the sum of scattering coefficient and absorption coefficient. The integrating nephelometry (IN) is an effective, economical and widely recognized method for online obtaining aerosol scattering coefficient (Beuttell and Brewer, 1949; Heintzenberg and Charlson, 1996; Abu-Rahmah et al., 2006). Early on the systematic limitations of this technique were noted, that is so-called truncation error caused by technically impossible to cover the full range of the scattering angle, and which has mainly studied through numerical simulations with Mie model (S. Ensor and P. Waggoner, 1970; Anderson et al., 1996; Anderson and Ogren, 1998; Heintzenberg et al., 2006; Müller et al., 2009). The measurement techniques for the extinction coefficient of atmospheric aerosols mainly include cavity ring-down spectroscopy (CRDS), cavity attenuation phase shift (CAPS) and cavity enhanced absorption spectroscopy (CEAS). CRDS has extremely high detection accuracy and mature measurement system, which performed well in laboratory studies and field measurements (O'Keefe and Deacon, 1988; Baynard et al., 2007; Berden et al., 2010; Pettersson et al., 2004; Strawa et al., 2003). Related in its basic principle to CRDS, previously CAPS was used to calibrate the reflectivity of mirrors also applied to measure atmospheric nitrogen dioxide (Kebabian et al., 2005; Ge et al., 2013; Herbelin and McKay, 1981). It currently has been extended to the field of aerosol extinction coefficient measurement (Kebabian et al., 2007; Petzold et al., 2013). Massoli et al. (2010) gave a detailed description of CAPS results in the aerosol extinction coefficient measurements, including the first laboratory characterization and field deployment. Onasch et al. (2015) calibrated the optical path length error of CAPS with MIE model using monodisperse polystyrene spheres generated in the laboratory. Rather than single wavelength measurements, CEAS with broadband light source applied for atmospheric trace gas detection (Fiedler et al., 2003; Ball et al., 2004; Chen and Venables, 2011) was later extended to quantitative aerosol extinction (Varma et al., 2013; Zhao et al., 2014; Suhail et al., 2019). The filter-based methods are most commonly used for online measuring aerosol absorption coefficient (Horvath, 1997; Hansen et al., 1982; Petzold and Schönlinner, 2004). Considering aerosol morphology changes, multiple scattering and shielding effects, these methods require many correction factors that limits the quality of measurement results (Bond et al., 1999; Weingartner et al., 2003). Recently, the photoacoustic spectroscopy (PAS) technique (Terhune and Anderson, 1977; Bruce and Pinnick, 1977; Adams et al., 1990), a direct method that can be easily-calibrated, has been developed into a stable instrument in the field measurement of aerosol absorption (Moosmüller et al., 1998; Arnott et al., 1999; Lack et al., 2006; Lewis et al., 2008; Sharma et al., 2013; Nakayama et al., 2015). Arnott et al. (2000) calibrated their aerosol photoacoustic instrument by measuring the photoacoustic response in the presence of NO₂ and compared its result with aethalometer. Lack et al. (2006) used ozone with a known optical absorption level to calibrate the photoacoustic system with CRDS.

During our recent field campaign in Yangtze River Delta (YRD), the measurements of aerosol optical properties showed discrepancies from different instruments, among which the extinction, absorption, and scattering coefficients were measured
by CAPS, PAS, and IN respectively (Du et al., 2020). For investigation of the discrepancy between instruments and correction of the measurement data, this study carried out an aerosol optical properties intercomparison measurement. During calibration measurement, the extinction coefficient was calibrated with MIE model using mono-disperse particles and the absorption coefficient was calibrated with transmission method using an absorbing gas, while the scattering coefficient was calibrated with combination of above model and the method using no-absorbing particles. In addition, an Incoherent Broad-Band Cavity Enhanced Absorption Spectrometer (IBBCEAS) was used to measure extinction coefficient of NO$_2$ for comparing with CAPS. Then the correction factors obtained from the laboratory calibration experiments were employed on the data observed in the filed measurement correspondingly and compared with the reconstructed extinction of the interagency monitoring of protected visual environment (IMPROVE) model. Furthermore, the calibrated field measurement results from photoacoustic and cavity attenuated phase shift instruments were intercompared. For aerosol optical properties, different optical methods showed good agreement and closure correlation after calibration, which has been rarely studied in laboratory calibration and field measurement. In addition, the corrected field measurement data are more reliable for subsequent study of aerosol optical properties in YRD region.

2 Materials and Methods

2.1 Instrument description

During calibration experiments, the optical properties of aerosol were measured by a Cavity Attenuation Phase Shift-ALBedo monitor (CAPS-ALB) (Shoreline Science Reaserch, Japan) and a Photoacoustic Extintiometer (PAX) (Droplet Measurement Technologies, US). In addition, a Scanning Mobility Particle Sizer Spectrometer (SMPS) (Model 3938, TSI, US) was employed to measure the number-size distribution for MIE model, and an IBBCEAS setup was used to measure NO$_2$ concentration for extinction calculation. Above instrument details are summarized in Table 1.

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Parameters</th>
<th>Time resolution</th>
<th>Flow, Lmin$^{-1}$</th>
<th>Wavelength, nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAPS-ALB</td>
<td>Extinction coefficient, Scattering coefficient [Mm$^{-1}$]</td>
<td>1 s</td>
<td>0.85</td>
<td>530</td>
</tr>
<tr>
<td>PAX</td>
<td>Absorption coefficient, Scattering coefficient [Mm$^{-1}$]</td>
<td>1 s</td>
<td>1</td>
<td>532</td>
</tr>
<tr>
<td>SMPS</td>
<td>Number size distribution [cm$^{-3}$]</td>
<td>5 min</td>
<td>0.3</td>
<td>-</td>
</tr>
<tr>
<td>IBBCEAS</td>
<td>NO$_2$ concentration [ppb]</td>
<td>1 min</td>
<td>0.6</td>
<td>355-380</td>
</tr>
</tbody>
</table>

Aerosol sample flow was drawn into the PAX using an external vacuum pump, then split between the wide-angle integrating reciprocal nephelometer and photoacoustic resonator for simultaneous online measurements of light scattering.
coefficient and absorption coefficient. In the photoacoustic cavity, the laser beam passing through the sample stream was modulated at the resonant frequency of the cavity, and the light-absorbing molecules were heated and quickly transferred the heat to the receiving end of the instrument, the pressure wave generated by periodic heating was detected by a sensitive microphone. The calculation formula of absorption coefficient \((b_{abs}^{obs})\) is as follows (Rosencwaig, 1980):

\[
b_{abs}^{obs} = \frac{P_{mic} \cdot A_{res} \cdot \pi^2 \cdot f_{res}}{P_L \cdot (\gamma - 1) \cdot Q}
\]

Where, \(P_{mic}\) is the pressure at the microphone at the resonant frequency \(f_{res}\), \(P_L\) is laser power, \(A_{res}\) is the geometric cross-section of the resonator, \(\gamma\) is the ratio of specific heat at constant pressure and volume, \(Q\) is the quality factor of the resonator that calculated from temperature, pressure, and relative humidity (\(RH\)).

The wide-angle integrating reciprocal nephelometer with a scattering integration angle of 6-174° range used in PAX, which detects scattering light from a parallel beam through a cosine-weighted detector. The detector located in the center of the cavity is fiber coupled to a photo-multiplier tube (PMT), where the measured laser power is proportional to the total scattering cross section. The expression for determining scattering coefficient \((b_{sca}^{obs})\) is given by (Abu-Rahmah et al., 2006):

\[
b_{sca}^{obs} = \frac{P_{PMT}}{P_L}
\]

Where \(P_{PMT}\) is the value of the PMT signal with scattering background subtracted, \(P_L\) is measured laser power. The scattering background was measured during the zeroing process of the instrument operation.

In addition, the extinction coefficient \((b_{ext}^{obs})\) considered as theoretical value in recommended calibration method of PAX that can be obtained by measuring the intensity of transmitted light with a photodetector combined with Lambert Beer's law as follow:

\[
b_{ext}^{obs} = \frac{\ln (I_0/I)}{L} \cdot 10^6 [\text{Mm}^{-1}]
\]

Where, \(I_0\) and \(I\) are the laser intensity with or without extinction substances, respectively. \(L\) is the path length of the laser beam through the cavity in meters, here is 0.354 m. \(10^6\) is a conversion factor to express extinction in \(\text{Mm}^{-1}\).

The CAPS-ALB using an internal vacuum pump to introduce aerosol flow into the sample cell to measure the extinction coefficient and scattering coefficient simultaneously. Nearly 1° truncation angle integrating sphere integrating nephelometer (ISIN) has been employed in CAPS-ALB. The integrating sphere with attached truncation reduction tubes located around the sample cell and PMT are equipped to collect scattering light, which effectively reduces the angle truncation error (Varma et al., 2003). As a typical kind of reciprocal nephelometer, its scattering coefficient \((b_{sca}^{obs})\) can also be calculated using Eq (2).

The extinction measurement system of CAPS-ALB utilizing a visible light-emitting diode (LED) with the luminescence as a light source and a sample cell incorporating two high reflectivity mirrors centered at the wavelength of the LED and a vacuum photodiode detector. The extinction coefficient of CAPS-ALB \((b_{ext}^{obs})\) is obtained by measuring the light attenuation of the visible long optical path with a vacuum photodiode, and detecting the phase shift of the square wave frequency modulation heterodyne detection of the light source, its expression as follow:
where $\cot$ is the cotangent, $c$ is the speed of light, $f$ is the LED modulation frequency, $T$ and $P$ are the sample temperature and pressure, respectively. The amount of phase shift ($\theta$) is a function of fixed instrument properties such as cell length, mirror reflectivity and modulation frequency, and of the presence of aerosols (Kebabian et al., 2007). The term $\cot \theta_0$ is obtained from a periodic baseline measurement (using particle-free air). It is worth mentioning that the effective optical-path error in the sample cell of CAPS-ALB, which caused by the purge airflow of the mirror limits the space of the aerosol samples, has been initially corrected in the internal calculation process. The original correction factor was 0.7 that close to the value reported by Onasch et al. (2015), which generally calibrated with MIE model calculation.

Our IBBCEAS device developed in-house was used to measure gas concentration in the NO$_2$ comparison experiment (Chen and Venables, 2011). The IBBCEAS measures the light intensity change of the light source through the optical cavity, then inverts the concentration of the gaseous samples. When a pair of high-reflectivity plano-concave mirrors with a reflectivity of $R$ are composed of an optical cavity with a length of $L$ that is illuminated by continuous broadband incoherent light, the output light intensity $I$ is equal to the sum of the output light intensity of each order. Combined with Lambert Beer's law, the expression for extinction coefficient $b_{\text{ext-CEAS}}$ at measured wavelength as follow (Fiedler et al., 2003; Ball et al., 2004):

$$b_{\text{ext-CEAS}}(\lambda) = \left( \frac{I_0(\lambda)}{I(\lambda)} - 1 \right) \left( \frac{1 - R(\lambda)}{L} \right) = \sum_i \sigma_i(\lambda) N_i$$

Here, $I_0$ is light intensity without absorbing matter, $\sigma_i$ and $N_i$ are absolute extinction cross section and concentration of species i. $I_0$, $I$, $R$, $\sigma_i$ and $N_i$ are functions of wavelength. Therefore, for different detection wavelengths, the extinction coefficient cannot be compared directly. A simple method is to establish a relationship with the species concentration. Fitting the extinction cross-section $\sigma_i$ to the extinction coefficient ($b_{\text{ext-CEAS}}$), the concentration of the measured gas $N_i$ can be inverted. Noting that the reflectivity $R$ of the cavity mirrors in IBBCEAS has been calibrated before our experiments, so the result of IBBCEAS can be considered as absolute value.

The number size distribution for MIE model calculation was obtained from SMPS, which is consists of an Electrostatic Classifier (Model 3082, TSI, US) and a Condensation Particle Counter (CPC) (Model 3750, TSI, US). The Electrostatic Classifier was used with a Long Differential Mobility Analyzer (LONG DMA) (Model 3081, TSI, US), its particle-size selection range is 14.1~736.5 nm, with a sample flow of 0.3 L min$^{-1}$ and a sheath flow of 3 L min$^{-1}$. The aerosol sample passes through the radioactive neutralizer to be charged, then enters the DMA to selects particles of different particle sizes by changing the voltage. The number of selected particles is counted to after the process of hygroscopic growth in CPC, which has an uncertainty of within ±10 % in measuring particle concentration (Petzold et al., 2013).

### 2.2 Experimental

Based on the above principles, we adopted the following experimental procedures to compare PAX and CAPS-ALB as Fig. 1 shows. The blue solid line represents the process of particle calibration, the red solid line represents the procedures of
gas calibration. All joints have been leak tested to ensure tightness.

Figure 1: Experimental schematics (a) Particle calibration (b) Gas calibration.

(a) Particle calibration

For systematic errors, such as angle truncation, laboratory generated nebulized ammonium sulfate (AS) (AR 99%, Aladdin Chemical) aerosols were used to calibrate and test the nephelometers of CAPS-ALB and PAX using same experimental procedure as follow. AS aqueous solution was nebulized by an atomizer (Model 9032, TSI, US) with filter air at a constant inlet pressure of 20 psi, which can generate a stable outlet flow rate of ~ 5 L min⁻¹. As shown by the blue solid line in the Fig. 1(a), the nebulized aerosol flow was diluted with filtered air to adjust its concentration, and then dried using diffusion dryers with silica gel that reduced the sample RH to ~10 % before delivery to the instruments, where the excess airflow was discharged by bypass. Only opening PAX or CAPS-ALB valves, the dry aerosol flow was connected to the instruments sampling port for at least ~5 min until the measured value stabilizes. The entire flow system used conductive silicone tubing’s and reduced bending to minimize the loss of particles during aerosol transportation. For high-concentration of non-absorbing AS aerosol with refractive index of 1.53+0.00i, the absorption effect can be ignored. Therefore, the scattering calibration factor ($f_{sca}^{obs}$) was calculated by comparing the measured extinction coefficient ($b_{ext}^{obs}$) and scattering coefficient ($b_{sca}^{obs}$) (Lewis et al., 2008; Cross et al., 2010).
For the purpose of estimating the scattering or extinction coefficients measured in above experiments and further correcting the absolute total scattering (extinction) coefficient, we performed an additional calibration using polystyrene latex (PSL) spheres with Mie model. This model is a rigorous analytical solution of the scattering field distribution of monochromatic light illuminates on spherical particles (Born and Wolf, 1999). Thus, assuming the particles to be round, it is considered feasible to apply Mie model to retrieve the number size distribution for calculating the total scattering coefficient of atmosphere aerosol. The scattering and extinction efficiency factor $Q_{\text{sca}}$ and $Q_{\text{ext}}$ can be calculated from the function of particle complex refractive index, light source wavelength and size distribution (Wu et al., 2018; Bohren and Huffman, 1983). By integrating the particle cross-sectional area $\pi D^2/4$, particle number concentration $N(D)$, and $Q_{\text{sca/ext}}$ on the particle diameter $D$ distribution, yields the calculated scattering and extinction coefficient $b_{\text{sca/ext}}^{\text{MIE}}$ as follow expression:

$$b_{\text{sca}}^{\text{MIE}} = \int_0^\infty Q_{\text{sca}} \cdot \frac{\pi D^2}{4} \cdot N(D) \cdot dD$$  \hspace{1cm} (6)

$$b_{\text{ext}}^{\text{MIE}} = \int_0^\infty Q_{\text{ext}} \cdot \frac{\pi D^2}{4} \cdot N(D) \cdot dD$$  \hspace{1cm} (7)

The experiments incorporating mono-disperse PSL spheres with complex refractive index 1.60+0.00i and diameter of 350±6 nm (Thermo Scientific) were carried out follow the calibration procedures of scattering calibration (replace PSL with AS). Opened the SMPS valve and connected the diluted dry aerosol flow to the its sampling port, then continuously measured together with CAPS-ALB or PAX for ~20 min for collecting at least three sets of effective data of particle-size distribution at each concentration. By comparing the MIE calculated average with the measured value for multiple concentrations, the MIE model correction factor ($f_{\text{sca}}^{\text{PAX-MIE}}, f_{\text{ext}}^{\text{CAPS-MIE}}$) can be determined.

b. Gas calibration

As noted in previous studies (Arnott et al., 2000), the PAS resonator acoustic calibration used sufficiently high concentrations of absorbing gas to generate a huge absorption, so that the Rayleigh scattering was negligible. Therefore, our experiment adopted high concentration NO$_2$ for absorption calibration and determined the absorption correction factor ($f_{\text{abs}}$) from comparison of measured absorption ($b_{\text{abs}}^{\text{obs}}$) and extinction ($b_{\text{ext}}^{\text{obs}}$) coefficients without knowing NO$_2$ concentration. In the case of only PAX valve opened, by diluting 200 ppm NO$_2$ in different dilution ratios, the filtered air and NO$_2$ mixture were introduced to PAX for ~5 min, in which the flow of filtered air and NO$_2$ were controlled by the mass flow controller to specified proportion, respectively. The entire flow system used Teflon tubes to minimize NO$_2$ loss and contaminations, and a bypass was set to ensure the stability of the sample flow and pressure.

Subsequently, considering the possible particulate loss of CAPS-ALB calibration, IBBCEAS and CAPS-ALB was used to measure NO$_2$ samples simultaneously for comparing the measured extinction coefficient in gaseous way. This experiment was carried out based on the experimental procedure for PAX absorption calibration, though closing PAX route and simultaneously opening the valves of CAPS and IBBCEAS. Based on the limitation of IBBCEAS the NO$_2$ concentration was controlled below 1 ppm and each concentration was maintained for at least about 15 min until the measured value stabilizes.
By multiplying the NO$_2$ concentration measured by IBBCEAS and the NO$_2$ extinction cross section from previous study of Voigt et al. (2002) at the CAPS-ALB detection wavelength. For reasonable comparison in extinction coefficient of IBBCEAS and CAPS-ALB, the spectral resolution of two instruments was need to be synchronized. CAPS-ALB uses LED as the light source and 10-nm wide optical filter to define the measurement range, but its specific band range hasn’t been found, here we presumed that to be 525-535 nm. Therefore, when calculating extinction coefficient of IBBCEAS from measured NO$_2$ concentration and its absorption cross section at the specific wavelength, the average value of the NO$_2$ absorption cross section in the range of wavelength 525nm to 535nm was applied. Then the conversion result of the extinction coefficient ($b_{ext-CEAS}$) measured by IBBCEAS was obtained. Thus, the extinction correction factor ($f_{ext-CAPS-CEAS}$) from comparison with IBBCEAS can be determined.

3. Results and Discussion

3.1 PAX calibration results

In our calibration experiments for PAX, with assumption of linearity in calibration down to the detection limit of the instruments (Arnott et al., 2000), the high concentration of absorbing gas and scattering particles generated a huge absorption and scattering effect that weaken the interference of noise for corrected the response curve of the PAX photoacoustic resonator and nephelometer respectively.

Fig. 2(a) shows the relationship between the absorption coefficient ($b_{abs}^{obs}$) and the extinction coefficient ($b_{ext}^{obs}$) in the NO$_2$ measurement results. The slope of fitted line in Fig. 2(a), which represent $f_{abs}^{obs}$, were determined to be $0.961 \pm 0.019$ with correlation factor $R^2 \sim 0.985$. The calibration result showed that the absorption measurement of PAX only needs slight correction and has high accuracy. Fig. 2(b) presents typical correlation plots comparing the extinction coefficient from transmissivity ($b_{ext}^{obs}$) for AS samples (The black solid dot) and MIE model calculation ($b_{ext}^{MIE}$) for PSL samples (The red solid dot) with the measured scattering coefficient ($b_{ sca}^{obs}$) respectively in PAX scattering calibration, where the extinction and scattering are theoretically equivalent due to negligible absorption.
Figure 2: PAX calibration results: (a) Comparison of the measured extinction and absorption coefficient. (b) Comparison of the measured and MIE-model calculated extinction coefficient with the measured scattering coefficient.

In Fig. 2(b), the slope of the black solid line indicates the measured scattering correction factor ($f_{\text{sca}}^{\text{obs}}$) that was determined to be $0.970 \pm 0.046$ with correlation factor $R^2 \sim 0.924$. Moreover, we calculated the absolute extinction coefficient with MIE model for further correction. Here, limited by the detection range, another set of coordinate system was used for comparison. The slope of the red solid line that indicates the MIE model scattering correction factor ($f_{\text{sca}}^{\text{PAX-MIE}}$) were determined to be $0.980 \pm 0.039$ with correlation factor $R^2 \sim 0.984$. The scattering correction factors from transmission method and MIE model were within acceptable range of the truncation error, and had only $\sim 1\%$ discrepancy in different measurement range, showing that a good agreement between the two methods and the reliability of PAX scattering calibration result.

### 3.2 CAPS-ALB calibration results

In the CAPS-ALB calibration experiment, we first utilized PSL spheres to correct its extinction coefficient through MIE model calculation, and then employed AS samples to correct its scattering coefficient comparing the calibrated extinction coefficient. In addition, we used self-developed experimental IBBCEAS device for further verifying the correction factor.
calculated by MIE model.

Fig. 3(a) shows correlation of extinction measured by CAPS-ALB ($b_{\text{ext}}^{\text{obs}}$) and extinction calculated by MIE model ($b_{\text{ext}}^{\text{MIE}}$) for 350nm mono-disperse PSL spheres. The slope in Fig. 3(a) represents extinction correction factor ($f_{\text{ext}}^{\text{CAPS-MIE}}$) were determined to be 0.983±0.018 with correlation factor $R^2$~0.999. It shows that the good accuracy of original calibration factor for the effective optical path error, only slight adjustment was required. The other factors that might affect the extinction calibration is the uncertainty of the aging effects of LED and detectors (PMT and vacuum photodiode), which has different effects according to cell geometry.

Correlation plots comparing scattering coefficient ($b_{\text{sca}}^{\text{obs}}$) and extinction coefficient ($b_{\text{ext}}^{\text{CAPS}}$) for AS samples measured by CAPS-ALB are shown in Fig. 3(b). According to its linear fitting result, $f_{\text{sca}}^{\text{obs}}$ were determined to be 1.016±0.002 with correlation factor $R^2$~0.996. It showed that the measured scattering coefficient has high accuracy, and verified the reliability of extinction correction factor of CAPS-ALB ($f_{\text{sca}}^{\text{CAPS-MIE}}$).

Fig. 3(c) presents the comparison between measured extinction coefficient of CAPS ($b_{\text{ext}}^{\text{CAPS}}$) and IBBCEAS ($b_{\text{ext}}^{\text{CEAS}}$) for NO$_2$ samples. The slope of Fig.3(c) represents extinction correction factor ($f_{\text{ext}}^{\text{CAPS-CEAS}}$) were determined to be 0.946±0.007 with correlation factor $R^2$~0.998. The experimental correction factor of IBBCEAS ($f_{\text{ext}}^{\text{CAPS-CEAS}}$) was consistent with the theoretical correction factor of the MIE model ($f_{\text{ext}}^{\text{CAPS-MIE}}$) within an acceptable error range of 4 %, proving that the reliability of MIE model calculation and the applicability of CAPS-ALB calibration, no matter whether choosing gas or particle ways.
3.3 Calibrated field measurement

The field measurements were carried out in the Gehu area of southwest Changzhou City, Jiangsu Province (31°63′ N, 119°90′ E) from 25 May to 27 June before the rainy season in 2019. Changzhou has a location in the center of the Yangtze River Delta and has a subtropical monsoon climate. The measurement site was surrounded by 60% of ecological wetlands and green gardens, and 20% of territorial waters, which results represented the regional ambient conditions of the Yangtze River Delta before the rainy season. The sampling point was located on the top floor of a building at the height of 15 m above ground.
and all sampling tubes used a cyclone size cutter (URG, 2.5 μm, 5 lpm).

The correction factor obtained from the laboratory calibration experiments was employed on the optical properties observed in the filed measurement correspondingly. For comparison, the IMPROVE model was applied to identify aerosol light extinction contribution of major chemical components during field measurement. The IMPROVE model was established by analyzing the data from the long-term monitoring of aerosol mass concentration carried out in multi-site of the Inter-agency Monitoring of PROtected Visual Environments network in the United States. The IMPROVE model reconstructs extinction coefficient using the mass concentration of aerosol chemical components and their mass extinction efficiency, which has been used worldwide for estimating the aerosol extinction coefficient (Pitchford et al., 2007; Tao et al., 2014). The major chemical components in this study including water-soluble inorganic ions, organic carbon (OC), and elemental carbon (EC) were analyzed and quantified by a monitor for aerosols and gases in ambient air (MARGA) (ADI 2080, Metrohm, Switzerland) and an OC/EC analyzer (Model RT-4, Sunset, US). The simplified general formula of IMPROVE model used in reconstruction of total scattering (extinction) coefficient ($b_{ext}^{IMPROVE}$) can be expressed as (Xia et al., 2017):

$$b_{ext}^{IMPROVE} = 2.2 \times f_s(RH) \times [Small \ (NH_4)_2SO_4] + 4.8 \times f_i(RH) \times [Large \ (NH_4)_2SO_4]$$

$$+ 2.4 \times f(RH) \times [Small NH_4NO_3] + 5.1 \times f(RH) \times [Large NH_4NO_3] + 2.8 \times [Small OM]$$

$$+ 6.1 \times [Large OM] + 1.7 \times f_{ss} \times [SS] + 1.0 \times [FS] + 0.6 \times [CM] + 8.28 \times [EC]$$

where $[X]$ represent the mass concentration of aerosol chemical component $X$, μg/m$^3$; Ammonium Sulfate [(NH$_4$)$_2$SO$_4$] = 1.375 [SO$_4^{2-}$]; Ammonium Nitrate [NH$_4$NO$_3$] = 1.29 [NO$_3^-$]; Organic Matters [OM] = 1.6 [OC]; Sea Salt [SS] = 1.8 [Cl$^-$]; Fine Soil [FS] = 2.2 [Al] + 2.49 [Si] + 1.94 [Ti] + 1.63 [Ca] + 2.42 [Fe]; Coarse Mass [CM] = [PM10] − [PM2.5]; $f_s(RH)$, $f_i(RH)$, and $f_{ss}$ represent RH growth curves of sulfate, nitrate, and SS (Jung et al., 2009). Due to the lack of soil element information, Ca$^{2+}$ was assumed to account for 5% of the concentration of fine soil mass based on previous studies, thus [FS] = 20[Ca$^{2+}$] (Amato and Hopke, 2012).

Considering unavailable period of aerosol composition measurement (Due to status of MARGA), only from 1 to 6 June were selected for the comparison. Fig. 4 (a) and (b) showed intercomparison of the measured extinction coefficient of PAX and CAPS-ALB with IMPROVE-calculated extinction coefficient, the linear fitting slopes are 1.182 and 1.183 with the correlation factor $R^2$ of 0.807 and 0.824, respectively. Comparing the correlation factor, it is in good agreement with Shanghai (0.83) and Hangzhou (0.81) in previous studies (Wang et al., 2016). Thus, it can be concluded that the IMPROVE model has good applicability in Gehu area. Here, the extinction of PAX was the sum of the measured absorption and scattering. In addition, Fig. 4(c) showed a timing diagram of the extinction coefficient from PAX, CAPS-ALB measurement and IMPROVE model calculation. It showed a good agreement between the measured and theoretical value and proved the reliability of our measurement data.
Figure 4: Intercomparison of the measured extinction coefficient of (a) PAX and (b) CAPS-ALB with IMPROVE-calculated extinction coefficient during field measurement (1-6 June 2019), and (c) the timing diagram of the extinction coefficient from PAX, CAPS-ALB measurement and IMPROVE model calculation.

Then the CAPS-ALB and PAX corresponding optical properties of field measurement were compared respectively in the case of calibrated and uncalibrated as Fig. 5(a), (b) and (c) showed. Here, the extinction coefficient of PAX has been mentioned above as well as the absorption of CAPS-ALB was the difference between the measured extinction and scattering. The linear fitting slope was 1.052, 1.024 and 1.046 from comparison of PAX and CAPS calibrated extinction, scattering and absorption coefficient, with the correlation factor $R^2$ as 0.936, 0.924 and 0.772. Comparing the calibrated and uncalibrated results, only slight corrections existed in the extinction and scattering coefficients, while the discrepancy in the absorption coefficient has been corrected from ~30% to less than 5%. It can be considered that the optical properties measured from PAX and CAPS-ALB with different measurement principles had a good agreement, which in turn proved the reliability of our laboratory calibration results and the closure correlation of CAPS-ALB and PAX measurements.

In addition, through deleting the time points of instruments data under zero calibration and abnormal working conditions, the overall trend of calibrated extinction, scattering and absorption coefficients during the measurement period (from 25 May to 27 June) were obtained as shown in Fig. 5(d), (e) and (f). For the aerosol optical properties of the measurement region, it showed a dominated contribution of scattering effect to the extinction coefficient, and a low levels of absorption coefficient.

The different internal structure of the nephelometers, even using the same principle, caused the slight difference in the
measured scattering coefficient. While the relatively small amount of absorption coefficient of CAPS-ALB derived from extinction subtracted scattering coefficient has been greatly affected. Therefore, the absorption coefficient which is difficult to quantify, was verified by CAPS-ALB via correcting the scattering coefficient and the relationship of optical properties.

4. Conclusion

In this work we carried out aerosol optical properties inter-comparison measurements from photoacoustic and cavity
attenuated phase shift instruments. The instruments were calibrated via laboratory experiments and the corrected field measurement data have also been intercompared. Thus, following points can be concluded:

(1) The laboratory results showed that disagreements exist between the two instruments before calibration. The scattering coefficient part plays a crucial role as the bridge in constructing correlation of both instruments. Then the corrected extinction and absorption coefficients from both instruments were intercompared well.

(2) The intercomparison of calibrated absorption and extinction coefficients in a field measurement using photoacoustic and cavity attenuated phase shift instruments showed good agreement. Therefore, laboratory calibrations were used for corrections for ensuring the quality of field data and further analysis of radiative study.

**Data availability**

The raw data from the experiments are available upon request (j.chen@usst.edu.cn).

**Author Contribution**


**Competing Interests**

The authors declare that they have no conflict of interest.

**Financial Support**

This work was supported by the National Natural Science Foundation of China (Grant Nos. 91544225, 51776129), the National Key Research and Development Program of China (Grant Nos. 2018YFC0213800, 2017YFC0211500) and the Open Fund Project of State Key Laboratory of Loess and Quaternary Geology (SKLLQG).

**References**


