Development of a New Correction Algorithm Applicable to any Filter-Based Absorption Photometer

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- 8 Abstract

9 Among the various measurement approaches to quantify light absorption coefficient (B_{abs}), filterbased absorption photometers are dominant in monitoring networks around the globe. Numerous 10 correction algorithms have been introduced to minimize the artifacts due to the presence of the 11 12 filter in these instruments. However, from our recent studies conducted during the Fire Influence on Regional and Global Environments Experiment (FIREX) laboratory campaign, corrected filter-13 based B_{abs} remains biased high by roughly a factor of 2.5 when compared to a reference value 14 using a photoacoustic instrument for biomass burning emissions. Similar over-estimations of Babs 15 from filter-based instruments exist when implementing the algorithms on six months of ambient 16 data from the Department of Energy (DOE) Atmospheric Radiation Measurement (ARM) 17 Southern Great Plains (SGP) user facility from 2013 (factor of roughly 3). In both datasets, we 18 observed an apparent dependency on single scattering albedo (SSA) and absorption Ångström 19 exponent (AAE) in the agreement between Babs based on existing correction factors and the 20 reference B_{abs}. Consequently, we developed a new correction approach that is applicable to any 21 filter-based absorption photometer that includes light transmission from the filter-based instrument 22 as well as the derived AAE and SSA. For the FIREX and SGP datasets, our algorithm results in 23 good agreement between all corrected filter-based Babs values from different filter-based 24 instruments and the reference (slopes ≈ 1 and $R^2 \approx 0.98$ for biomass burning aerosols and slopes 25 \approx 1.05 and R² \approx 0.65 for ambient aerosols). Moreover, for both the corrected B_{abs} and the derived 26

27 optical properties (SSA and AAE), our new algorithms work better or at least as well as the two

common PSAP-based correction algorithms. The uncertainty of the new correction algorithm is estimated to be $\sim 10\%$, considering the measurement uncertainties of the operated instruments.

Therefore, our correction algorithm is applicable to any filter-based absorption photometer and

31 has the potential to "standardize" reported results across any filter-based instrument.

32 1. Introduction

Light-absorbing atmospheric aerosols directly affect the Earth's energy budget by absorbing solar 33 radiation, leading to a warming effect when they are suspended in the atmosphere and to the 34 melting of snow and ice following deposition (Bond and Bergstrom, 2006; Boucher, 2015; Horvath, 35 1993). For decades, scientists have conducted field experiments around the globe to investigate 36 37 how absorbing aerosols influence the atmospheric radiative balance and interact with clouds (e.g., Andrews et al. (2011); Cappa et al. (2016); Lack et al. (2008b); Rajesh and Ramachandran (2018); 38 Schwarz et al. (2008)). These experiments may be performed at fixed stations (e.g., observation 39 40 sites maintained by the Department of Energy (DOE) Atmospheric Radiation Measurement (ARM)

41 program or the National Oceanic and Atmospheric Administration (NOAA) Global Monitoring

Division (GMD)) or on mobile platforms (e.g., car trailer, aircraft, and ship), typically involving 42 43 the measurements of aerosol chemical, physical, and optical properties. Crucial to the quantification of the radiative forcing of absorbing aerosols are measurements of the absorption 44 45 coefficient (B_{abs}). For example, long-term monitoring of B_{abs} provides essential data to evaluate chemistry-climate model simulations (e.g., Chen et al. (2019); Vignati et al. (2010)), while 46 intensive measurements of Babs during short-term field campaigns allow for the investigation of 47 optical properties that govern features of aerosol forcing (e.g., McMeeking et al. (2014); Olson et 48 49 al. (2015)).

- A variety of instruments have been used to measure B_{abs} , which generally classified into two large categories: filter-based techniques and photoacoustic techniques (Lack et al., 2014; Moosmüller et al., 2009). The major difference between the two categories of technique is that B_{abs} is measured
- after the aerosols are deposited on the filter media in the filter-based instruments, while the aerosols are characterized within an air stream in the photoacoustic instruments. Compared to the filter-
- 55 based instruments, the photoacoustic instruments have the advantage of avoiding potential artifacts
- 56 due to the contact of aerosols with filters; therefore, they are often used as the reference instruments
- 57 in inter-comparison studies of aerosol absorption (e.g., Arnott et al. (2005); Davies et al. (2019);
- 58 Jiang et al. (2018); Li et al. (2019); Schmid et al. (2006); Sheridan et al. (2005)).
- Filter-based absorption photometers have been widely used at observational sites around the world
 due to their ease of operation and relatively low cost. Numerous instruments can be classified as
 filter-based absorption photometers including the Radiance Research Particle Soot Absorption
- 62 Photometer (PSAP), the NOAA Continuous Light Absorption Photometer (CLAP), the Brechtel
- 63 Manufacturing Tricolor Absorption Photometer (TAP), the Magee Scientific Aethalometer
- 64 (AETH), and the Thermo Scientific Multi-Angle Absorption Photometer (MAAP). Operationally,
- all of these instruments are similar in that aerosols are deposited onto a filter and the reduction in the transmission (T_{r}) of light by the particles (comptimes called attenuation (ATD)) is a set of the
- the transmission (Tr) of light by the particles (sometimes called attenuation (ATN)) is used to infer B_{abs}. Where the instruments may differ is that some are multi-wavelength (multi- λ) instruments
- B_{abs}. Where the instruments may differ is that some are multi-wavelength (multi- λ) instruments (e.g., 3λ-PSAP, CLAP, TAP, 7λ-AETH models), while others are not (e.g., 1λ-PSAP, other AETH
- 69 models, MAAP).
- One challenge with filter-based absorption photometers is that biases can arise due to the presence 70 of the filter. For example, light scattering by particles loaded onto the filter or by the filter itself 71 72 may affect the transmission of light (e.g., (Arnott et al., 2005; Bond et al., 1999)); non-absorbing material may result in absorption enhancement (e.g., (Cappa et al., 2008)); or organic vapors 73 74 adsorbed to the filter may itself absorb light (e.g., (Subramanian et al., 2007)). Consequently, 75 various correction algorithms exist to minimize these biases, but they are often specific only to 76 certain instruments. For example, some are applicable to the PSAP, CLAP, and TAP (e.g., (Bond et al., 1999; Müller et al., 2014; Ogren, 2010; Virkkula, 2010; Virkkula et al., 2005)), while others 77 78 are applicable to the AETH (e.g., (Arnott et al., 2005; Collaud Coen et al., 2010; Drinovec et al., 2017; Kirchstetter and Novakov, 2007; Schmid et al., 2006; Virkkula et al., 2007, 2015; 79 80 Weingartner et al., 2003)).
- Although the equations associated with these existing correction algorithms are different, they share some commonalities. For example, the filter-based absorption photometers are assessed using laboratory (e.g., ammonium sulfate, fullerene soot) or ambient aerosols during experiments which include reference measurements of B_{abs}. These reference measurements often include either
- direct photoacoustic B_{abs} or inferred B_{abs} as the difference between the extinction coefficient (B_{ext})

- and the scattering coefficient (B_{scat}). Correction equations are developed by comparing data between the filter-based instrument and the reference instrument, where the equations often contain one term that accounts for filter loading effects and another that accounts for multiplescattering effects. Consequently, the correction equations frequently incorporate both Tr and either B_{scat} or the single-scattering albedo (SSA) to account for these effects. However, even when the
- 91 correction algorithms are applied, potential issues can remain such as:
- Corrected filter-based B_{abs} may remain biased high relative to a reference value of B_{abs} (e.g., (Arnott et al., 2003; Davies et al., 2019; Lack et al., 2008a; Li et al., 2019; Müller et al., 2011a)).
- Comparisons between the reference instrument and B_{abs} corrected by different algorithms can yield variable agreement (e.g., (Collaud Coen et al., 2010; Davies et al., 2019; Saturno et al., 2017)).
- 97 3. Corrected B_{abs} from different filter-based absorption photometers may not agree (e.g., (Davies et al., 2019; Müller et al., 2011a)).
- 99 4. Derived products (such as absorption Ångström exponents (AAE)) may differ based on the
 100 implemented correction algorithm (e.g., (Backman et al., 2014; Davies et al., 2019)).
- 101 5. The agreement between measurements of B_{abs} and estimates of B_{abs} by chemistry-climate
- models may vary based on the implemented correction algorithm (e.g., (Alvarado et al., 2016)).

The first three issues in this list may arise due to differences in aerosol optical properties between those used in deriving the correction equation and those associated with a given aerosol sample, and these issues can propagate through to the fourth issue. The final issue is arguably most important because evaluation of chemistry-climate models may be severely affected by the differences between different correction algorithms, which may inhibit the modeling community from providing accurate projections of future temperature and precipitation response.

In this work, we seek to address some of these issues. First, we evaluate the CLAP, TAP, and 109 PSAP using two common PSAP-based correction algorithms, namely Bond et al. (1999) as 110 updated by Ogren (2010) and Virkkula et al. (2005) as updated by Virkkula (2010). For brevity, 111 we refer to these corrections as "B1999" and "V2005" for Bond et al. (1999) and Virkkula et al. 112 (2005), respectively, incorporating their respective updates. In addition, we propose new 113 correction algorithms that are applicable to any filter-based absorption photometer (e.g., CLAP, 114 TAP, PSAP, and AETH) across multiple wavelengths by combining observed filter-based Babs 115 with B_{scat} (e.g., from a co-located nephelometer (NEPH)) and reference B_{abs} (e.g., from a co-116 located photoacoustic instrument). However, in reality (e.g., at long-term observatories), reference 117 values of B_{abs} are rare, and in some cases, complementary B_{scat} measurements may not exist; 118 consequently, we also provide methods to correct filter-based Babs data in these scenarios. To our 119 knowledge, this is the first study to simultaneously evaluate B1999 and V2005 corrections on 120 PSAP "successors" (i.e., CLAP and TAP) and to present a correction algorithm that is broadly 121 122 applicable to any filter-based absorption photometer. Regarding the latter, even if our correction algorithm has its own limitations, its use can nevertheless standardize the reporting of B_{abs} in long-123 term datasets. 124

125 2. Methodology

126 We developed the general form for our correction algorithms using CLAP and TAP measurements

- 127 collected from biomass burning (65 fires in total) during the Fire Influence on Regional to Global
- 128 Environments Experiment (FIREX) laboratory campaign in 2016. By using biomass burning

emissions, we considered a dataset spanning a broader range of aerosol optical properties (SSA at 129 130 652 nm: 0.14-0.98; AAE: 1.25-4.73) than has traditionally been used in developing these correction algorithms. We then conducted further evaluation and validation of the model using 131 132 ambient data, specifically using CLAP measurements from the DOE ARM Southern Great Plains (SGP) user facility in Lamont, OK, USA (02/01/13 to 07/09/13). Our algorithms were then 133 extended to the AETH data from the FIREX laboratory campaign and the PSAP data collected at 134

- the SGP site to verify the generalizability of the algorithms. 135
- 136 2.1. The FIREX campaign
- 137 2.1.1. Experimental setup

In October and November of 2016, we participated in the laboratory portion of the FIREX 138 139 campaign to investigate the wildfire smoke and their impact on the atmosphere. During the campaign, over 100 burns took place at the U.S. Forest Service's combustion facility at the Fire 140 Sciences Laboratory (FSL). The fuels burned in this study are representative of western US 141 ecosystems, such as spruce, fir, various pines, and "chaparral" biome (e.g., manzanita, chamise). 142 143 (See Koss et al. (2018) and Selimovic et al. (2018) for more details).

144 A typical burn lasted for 1-3 hours depending on the smoke sampling strategies (e.g., stack burns versus room burns). During each burn, one or multiple "snapshots" of smoke (typical B_{abs} at 652 145 nm ranged from 100 to 1200 Mm⁻¹) were transferred from the combustion room at FSL into a 146 mixing chamber (210 L) through a long transfer duct (30 m in length, 8" in diameter). The smoke 147 148 was then diluted by filter air (~230 LPM) in the chamber. Once the concentration in the chamber was stable (detected by the Photoacoustic Extinctioneter (PAX) which was operated continuously 149 through all fires), the smoke was passed to a suite of instruments to obtain aerosol and gas phase 150 parameters. This chamber also served as an intermediate between the transfer duct and the 151 instrumentation to minimize potential biases that arose due to different sample flow rates and 152 sample locations of the instruments. A more detailed description of our experiments can be found 153

- in Li et al. (2019). 154
- 2.1.2. Measurements of aerosol optical properties 155
- 156 During the campaign, five instruments provided measurements of B_{abs} (CLAP, NOAA GMD; TAP,
- Brechtel Manufacturing Inc. (BMI); Aethalometer (Model AETH-31), Magee Scientific; and two 157
- PAXs (Model PAX-870 and PAX-405), Droplet Measurement Technologies) and two instruments 158
- provided measurements of B_{scat} (PAX-870 and PAX-405). The instruments included in the present 159
- work are summarized in Table 1. 160
- Both CLAP and TAP provide B_{abs} measurements of the particles deposited on a filter, similar to 161
- PSAP. Different from PSAP, there are multiple filter spots (8 sample spots and 2 reference spots) 162
- cycling of one filter in CLAP and TAP, enabling the instruments to run continuously through two 163
- or three burns without changing filter. In the CLAP and TAP, sample illumination is provided by 164 165
- LEDs operated at three wavelengths (467, 528, and 652 nm). Here, we apply both B1999 and V2005 to CLAP and TAP data, similar to previous work (e.g., (Backman et al., 2014; Davies et 166
- 167 al., 2019)).
- 168 The key differences between the CLAP and TAP during the FIREX campaign include:

- The spot change of the CLAP was manually performed when Tr reached approximately 0.5
 (or ATN decreased to ~69), while the TAP advanced to a new spot automatically with a Tr
 threshold set to be 0.5.
- 172 2. The spot area, flow rate, and LED-detected wavelengths differed slightly (Table 1).
- The CLAP recorded B_{abs} every one minute, while the TAP recorded B_{abs} every ten seconds. To enable the following analysis, we compute the 1-minute averages of TAP-derived parameters.
- 4. For the first portion of the campaign (the first 17 days of the 45-day campaign), Pallflex E70-
- 2075S filters were used in the CLAP while Azumi filters (model 371M, Azumi Filter Paper
 Co., Japan) were used in the second portion of the campaign (due to a lack of availability of
- the Pallflex filters). The TAP was equipped exclusively with the Azumi filters throughout the campaign. We apply the filter correction recommended in Ogren et al. (2017) to the CLAP and convert from Pallflex to Azumi filters.
- 181 5. BMI substantially re-engineered the CLAP in their development of the TAP.
- 182 These differences resulted in variable agreement between the CLAP and TAP during FIREX;
- however, the two instruments did largely agree within experimental uncertainty (e.g., see Fig. S8
- and Fig. S13 in Li et al. (2019)).
- 185 A PAX measures B_{abs} and B_{scat} simultaneously for suspended particles using a modulated diode 186 laser. We use these photoacoustic absorption measurements as the reference to evaluate the filter-
- 186 laser. We use these photoacoustic absorption measurements as the reference to evaluate the filter-187 based B_{abs} and develop our correction algorithms. To enable the evaluation of CLAP and TAP 188 which operate at different wavelengths than the PAXs, we interpolate the measurements of B_{abs}
- and B_{scat} to the wavelengths of 467, 528, and 652 nm using the values of AAE and scattering
- Ångström exponents (SAE), similar to Backman et al. (2014) and Virkkula et al. (2005).
 Theoretically, AAE and SAE fit absorption and scattering as power law functions of wavelength
- 191 Theoretically, AAE and SAE fit absorption and 192 (Borgetrom et al. 2007)
 - **192** (Bergstrom et al., 2007).
 - Due to the numerous correction algorithms for the Aethalometer (e.g. (Arnott et al., 2005; Collaud
 Coen et al., 2010; Kirchstetter and Novakov, 2007; Saturno et al., 2017; Schmid et al., 2006;
 Virkkula et al., 2007; Weingartner et al., 2003)), we do not evaluate these in the present work to

limit the scope. In fact, the majority of our focus is the B1999 and V2005 corrections to TAP and

197 CLAP. However, we still test the performance of the new algorithms on the AETH to explore its

- 198 applicability to that instrument.
- 199 2.2. Measurements of aerosol optical properties at the SGP observatory
- The ambient data used in this manuscript are the ground-based aerosol data measured at the SGP observatory from 02/01/13 to 07/09/13 (archived at https://www.archive.arm.gov/discovery/). For evaluation purposes, we randomly select a range of dates during which the observations are valid (without incorrect, suspect, and missing data) and the PASS is operated after laser (532 nm) upgrade.
- At the site, an impactor was used to switch the sampling between two cutoffs (particle diameter
- 206 <10 μ m (PM10) in the first 30 minutes of each hour and <1 μ m (PM1) in the latter 30 minutes of
- each hour). The aerosols exiting from the impactor were dried to RH less than 40% and passed to
- a CLAP, a PSAP, and two NEPHs. Moreover, a three-wavelength photoacoustic soot spectrometer
- 209 (PASS-3) was operated at the site and measured B_{abs} and B_{scat} of the aerosols, but these aerosols
- did not pass through the impactor (e.g., characterizing total suspended particles (TSP)). Typical
- B_{abs} and B_{scat} reported at the site ranged from 0 to 10 Mm^{-1} and 0 to 50 Mm^{-1} at 550 nm, respectively

(e.g., (Sherman et al., 2015)). Although the site is rural (clean background air), long-term transport
 aerosols (such as mineral dust, absorbing organic aerosols, and secondary organic aerosols (SOA))

214 may affect the local aerosol properties (Andrews et al., 2019).

215 We preprocess the SGP data in three steps. First, due to the systematic difference of aerosol sizes

between PASS-derived and filter-based absorption, we only include the PM10 observations, inherently assuming that any differences in the optical properties of PM10 and TSP are negligible.

Then, we smooth the 1-second data into 10-minute averages. Thirdly, we estimate the detection

- 219 limits at each of the three wavelengths in the PASS-3 using the data measured during the
- 220 "background zero" periods (Allan, 1966) and discard the observations which are below the
- detection limits. With a 10-min-averaging-time, the detection limits (3σ) for the PASS-3 are 0.78
- 222 Mm⁻¹ (405 nm), 2.01 Mm⁻¹ (532 nm), and 0.30 Mm⁻¹ (781 nm). For the filter-based instruments,
- the detection limits are based on previous studies (See Table 1). Moreover, we only retain the
- observations that satisfy B_{abs} (405 nm) > B_{abs} (532 nm) > B_{abs} (781 nm) (or AAE>0), similar to Fischer and Smith (2018). As with the PAX data from the laboratory, we adjust the PASS-derived
- Fischer and Smith (2018). As with the PAX data from the laboratory, we adjust the PASS-der
 B_{abs} to 467, 528, and 652 nm using the inferred AAE values for each 10-minute average.

Instrument	Flow rate (LPM)	Spot area (cm²)	Type of filter	Measured parameters	Response time	Measurement uncertainty	Detection limit (3σ, Mm ⁻¹)
PAX-870	1.0	-	-	B _{abs} and B _{scat} (870 nm)	1s	~11% (B _{abs}) ~17% (B _{scat}) (Nakayama et al., 2015)	0.47 (B _{abs}) 0.66 (B _{scat}) ^a
PAX-405	1.0	-	-	B_{abs} and B_{scat} (405 nm)	1s	4% (B _{abs}) 7% (B _{scat}) (Nakayama et al., 2015)	0.27 (B _{abs}) 0.60 (B _{scat}) ^a
PASS-3 b	1.0	-	-	B _{abs} and B _{scat} (405, 532, and 781 nm)	1s	4 %, 8 %, and 11 % (B _{abs}) (Nakayama et al., 2015)	0.78 (405 nm) 2.01 (532 nm) 0.30 (781 nm) ª
NEPH Þ	7.5	-	-	B _{scat} (450, 550, and 700 nm)	1s	10% (Anderson et al., 1996)	0.29 (450 nm) 0.11 (550 nm) 0.17 (700 nm) (5-min average) (Müller et al., 2011b)
CLAP	0.83 ± 0.02 (FIREX) 0.945 (SGP)	0.199 (FIREX) 0.195 (SGP)	Pallflex E70-2075S and Azumi filter (model 371M) °	B _{ATN} and Tr ^d (467, 529, and 653 nm)	60s	30% (Ogren et al., 2017)	0.6 (1-min average), 0.12 (10-min average) (Ogren et al., 2017)
ТАР	1.26 ± 0.01	0.253	Azumi filter (model 371M) ⁰	B _{ATN} and Tr ^d (467, 528, and 652 nm)	10s	30% (Laing et al., 2016)	2.67 (467 nm) 4.11 (528 nm) 2.13 (652 nm) (30-s average) (Davies et al., 2019)
AETH	2.4	0.5	quartz fiber sampling tape	B _{A™} and Tr (370, 470, 520, 590, 660, 880, and 950 nm)	120s	10% (Sedlacek, 2016)	0.1 (Sedlacek, 2016)
PSAP	1.0	0.178	Pallflex E70-2075W	B _{ATN} and Tr ^d (470, 522,and 660 nm)	60s	~15% (Bond et al., 1999)	0.3 (Springston, 2016)

227 Table 1 Summary of specifications for instruments relevant to this work.

^a The detection limits of PAX and PASS-3 are determined by Allan deviation analysis (Allan, 1966) of B_{abs} during "background zero".

^b During the analysis of the data collected at the SGP, we use B_{abs} derived by the PASS and B_{scat} derived by the NEPH to yield the coefficients in the algorithms.

^c Two types of filters were used during the FIREX campaign (See Sect. 2.1.2).

^d The operating wavelengths of CLAP, TAP, and PSAP are stated slightly different by the instrument manufactures. We simply use 467, 528, and 652 nm throughout

this manuscript.

233 2.3. Calibrations

249

Following Bond et al. (1999) and Ogren et al. (2010), the filter-based instruments were calibrated and corrected for sample area, flow rate, and filter type (see Li et al. (2019) for the FIREX data and Sherman et al. (2015) for the SGP data). Other than that, we did not do any verification beyond the manufacturer's calibration for the filter-based instruments. The SGP nephelometer measurements were corrected for truncation effects (Sherman et al., 2015). The FIREX photoacoustic measurements were calibrated by ammonium sulfate aerosol and fullerene soot (Li et al., 2019).

241 2.4. The correction algorithms

242 In filter-based instruments, the light intensities transmitted through the sample spot and blank spot

of the filter are recorded as I_s and I_b , respectively. The logarithmic ratio of the two intensities at time t is defined as ATN using the Beer-Lambert law:

245
$$ATN(t) = -100 \times ln \frac{I_s(t)}{I_b(t)}$$
1

where ATN = 0 when beginning a new filter spot (t = 0).

247 The ATN can be related to Tr by normalizing I_s/I_b at time t relative to I_s/I_b at the start of a new 248 filter spot (t = 0):

$$Tr(t) = \frac{I_s(t)/I_b(t)}{I_s(0)/I_b(0)} = exp(\frac{-ATN(t)}{100})$$
2

- The change of ATN over a time interval (Δt) for the instrument operated at a volume flow rate of Q and spot area of A yields the attenuation coefficient (B_{ATN}) for that time interval:
- $B_{ATN} = \frac{A}{Q \times \Delta t} \times \Delta ATN$ 3

 B_{ATN} is finally converted to B_{abs} by applying correction algorithms. The general form of the correction algorithms presented for the PSAP in Bond et al. (1999) and Virkkula et al. (2005) can be summarized as:

 $B_{abs} = B_{ATN} \times f(Tr) - C_1 \times B_{scat}$

4

where f(Tr) is some function of Tr (that may vary between approaches), correcting for the filter loading effect. C₁ is a constant that may vary with wavelength; specifically, it is a penalty for the light scattering by the particles collected on the filter which may contribute to the quantification of ATN. In most atmospheric and laboratory studies, B_{scat} is measured independently, typically using a co-located NEPH.

- Besides the above mentioned B1999 and V2005 corrections, the constrained two-stream (CTS) correction proposed by Müller et al. (2014) can be also applied on PSAP-similar instruments. The CTS correction was developed based on the relationship between absorption optical depth and B_{abs}. However, it is not straightforward to reformulate the CTS correction as a function of Tr as those presented in Eq. (5) and Eq. (6). Thus, we exclude the B_{abs} results corrected by the CTS correction
- in the following analysis. For those who are interested in the B_{abs} results corrected by the CTS
- correction, we provide the correction results of our SGP-CLAP data in Fig. S9.

269 2.4.1. The B1999 correction

Bond et al. (1999) was the first study to present the correction algorithm for filter-based instruments. This empirical correction was originally developed for the PSAP operated at 550 nm using various mixtures of laboratory-generated nigrosin (SSA ≈ 0.5) and ammonium sulfate (SSA ≈ 1) with B_{abs} ranged from 0 to 800 Mm⁻¹.

After calibrating the flow rate and spot area of the PSAP, Bond et al. (1999) derived $C_1 = 0.016$ and defined f(Tr) as follows:

5

276 $f(Tr)_{B1999} = \frac{1}{C_2 \times Tr + C_3}$

Following an adjustment by Ogren (2010), C_2 and C_3 in Eq. (5) were updated to 1.55 and 1.02, respectively. These are the values used in the present work (Table 2) for B1999. Moreover, Ogren (2010) stated that the correction forms of Eq. (4) and Eq. (5) were valid for any wavelength, while additional experiments were needed to establish the equation parameters for the wavelengths other than 574 nm.

282 2.4.2. The V2005 correction

Virkkula et al. (2005) developed a correction algorithm for both three-wavelength PSAP (467, 530, and 660 nm) and one-wavelength PSAP (574 nm) using the same functional form as Eq. (4). Since the operating wavelengths of the photoacoustic instruments and the NEPH were different from those of the PSAP, the measured photoacoustic B_{abs} and B_{scat} was extrapolated or interpolated to 467, 530, and 660 nm, using inferred AAE and SAE respectively. In this study, the authors used various mixtures of kerosene soot, ammonium sulfate, and polystyrene latex (SSA ranged from 0.2 to 0.9) with B_{abs} ranging from 0 to 800 Mm⁻¹ at 530 nm.

Different from the f(Tr) in the B1999 correction which was a reciprocal function of Tr, the f(Tr)
 presented in V2005 was a multivariate linear function of the natural logarithm of Tr and SSA
 (including an interaction term between the two):

- 293 $f(Tr(\lambda), SSA(\lambda))_{V2005} = C_4 + C_5 \times (C_6 + C_7 \times SSA(\lambda)) \times ln(Tr(\lambda))$ 6
- where the parameters in Eq. (6) vary with wavelengths. The parameters in V2005 were updated in Virkkula (2010) by correcting for flowmeter calibration (Table 2).
- Due to the unknown values of SSA before deriving B_{abs} , Virkkula et al. (2005) provided a solution through an iterative procedure. In the iteration, B_{abs} is first calculated using the B1999 correction (e.g., Eq. (4) and Eq. (5)) and is then used to compute the initial guess of SSA for use in Eq. (6).
- 299 The B_{abs} and SSA can be updated using Eq. (4) and Eq. (6) until convergence is reached.
- 300 2.4.3. Refitting the coefficients in B1999 and V2005

With the reference measurements of B_{abs} from the photoacoustic instruments, we are able to refit the coefficients in the B1999 and V2005 corrections (C₂ to C₇ in Eq. (5) and Eq. (6)) using our data. Specifically, we use the Levenberg-Marquardt algorithm (1944) to iteratively fit the coefficients until the chi-square of the coefficients are minimized. The fitting is implemented using the "Curvefit" function in Igor Pro. It is worth noting that the derived coefficients may only be valid for the SGP and FIREX data. For aerosol properties different from our study, the optimal

307 coefficients are likely to be different from the ones reported here. Hereafter, the B1999 and V2005

results with refitted coefficients are referred to as "updated B1999" and "updated V2005", respectively.

310 2.4.4. The new correction

We develop a set of new correction algorithms with the same general form as Eq. (4) using the biomass burning emissions from 65 different burns during the FIREX laboratory study, providing a broader range of aerosol optical properties and aerosol concentrations than previous work. This was motivated by the disagreement that remained between filter-based and photoacoustic instruments, even after applying B1999 to the data (e.g., see Li et al. (2019) Fig. 4 and our Fig. 2 below). These differences may persist because we were effectively extrapolating the B1999 correction equation to values outside the range for which it was developed.

This new correction is developed based on multiple linear regression techniques with three dependent variables of ln(Tr), SSA, and AAE and one independent variable of B_{abs}/B_{ATN} (Eq. (7) -(9)). As with other correction equations, this model takes into account the influence of scattering and weakly-absorbing materials, but we have the additional aim of developing a model that is applicable to any filter-based absorption photometer.

Similar to the B1999 and V2005 corrections, this new model starts with the general form of Eq. (4), re-written here to define B_{scat} in terms of SSA and B_{abs} .

$$\boldsymbol{B}_{abs}(\boldsymbol{\lambda}) = \boldsymbol{B}_{ATN}(\boldsymbol{\lambda}) \times f(Tr(\boldsymbol{\lambda})) - C_1 \times \frac{SSA(\boldsymbol{\lambda})}{1 - SSA(\boldsymbol{\lambda})} \times \boldsymbol{B}_{abs}(\boldsymbol{\lambda})$$

$$7$$

- Re-arranging this equation to move all B_{abs} terms to the left-hand side yields:
- 327 $\boldsymbol{B}_{abs}(\boldsymbol{\lambda}) = \boldsymbol{B}_{atn}(\boldsymbol{\lambda}) \times g(\boldsymbol{Tr}(\boldsymbol{\lambda}), \boldsymbol{SSA}(\boldsymbol{\lambda}))$ 8

328 where
$$g(Tr(\lambda), SSA(\lambda)) = f(Tr(\lambda)) \times \frac{1 - SSA(\lambda)}{1 - (1 - C_1) \times SSA(\lambda)}$$

We define a new function "g" that can be used in Eq. (8). Specifically, we construct a multivariate linear model for "g", introducing AAE as a dependent variable and including interaction terms between SSA, AAE, and ln(Tr):

$$g(Tr(\lambda), SSA(\lambda), AAE) = G_0 + G_1 \times ln(Tr(\lambda)) + G_2 \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times AAE + G_4 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times ln(Tr(\lambda)) \times ln(Tr(\lambda)) \times SSA(\lambda) + G_3 \times ln(Tr(\lambda)) \times ln(Tr(\lambda))$$

325

$$G_5 \times SSA(\lambda) \times AAE + G_6 \times ln(Tr) \times AAE + G_7 \times SSA(\lambda) \times AAE \times ln(Tr(\lambda))$$
 9

Equation (9) suggests that different combinations of SSA, AAE and ln(Tr) can result in the same 334 value of "g" (i.e., B_{abs}/B_{ATN}); likewise, a given value of B_{abs}/B_{ATN} may have infinitely many points 335 with distinct slopes passing through it (Fig. S3). Although Eq. (9) is developed based on statistical 336 approaches, we attempt to relate this statistical model to physical effects. The coefficients $G_1 - G_3$ 337 are fairly straightforward, as these account for the influence of filter loading (G₁), relative light 338 scattering by the aerosols (G_2) , and the brownness of the aerosols (G_3) . The interaction terms (G_4) 339 $-G_7$) are more difficult in assigning a physical meaning; however, the interaction between filter 340 loading and relative light scattering (G_4) is might be interpreted as an absolute light scattering by 341 the aerosols on the filter, while the interaction between filter loading and aerosol brownness (G_6) 342 is somewhat analogous to G₄. The three-way interaction between filter loading, scattering and 343 brownness of aerosols (G_7) is required because of the three two-way interaction terms. 344

To further this physical interpretation of our statistical model (Eq. 9), we explore the relationship 345 between $\frac{B_{abs}}{B_{ATN}}$ and ln(Tr), which essentially follows a "y = m·x + b" form, where y is $\frac{B_{abs}}{B_{ATN}}$ and x is 346 ln(Tr). The slope (m) is defined as $G_1 + G_4 \times SSA + G_6 \times AAE + G_7 \times SSA \times AAE$, and the intercept 347 (b) is defined as $G_2 \times SSA + G_3 \times AAE + G_5 \times SSA \times AAE$. Therefore, different combinations of SSA 348 and AAE modulate this relationship between $\frac{B_{abs}}{B_{ATN}}$ and ln(Tr). For example, loading "black" 349 particles on the filter (e.g., AAE ~1 and SSA ~0.3) tends to produce larger values of $\frac{B_{abs}}{B_{ATN}}$, while 350 loading "white" particles on the filter (e.g., AAE ~3 and SSA ~0.9) tends to produce smaller values 351 of $\frac{B_{abs}}{B_{ATN}}$ (see Fig. S3 of this work and Fig. 4 in Virkkula et al.(2005)). This relationship becomes 352 more complex when considering, e.g., mixed sulfate and black carbon particles; SSA can be high 353 while AAE is low, and the corresponding $\frac{B_{abs}}{B_{ATN}}$ can be variable (also see Fig. S3). Therefore, in 354 order to properly compensate for the effects of loading and aerosol optical properties, a multiple 355 356 linear regression with interaction terms is introduced in Eq. (9).

A detailed description of the procedure for the model development (e.g., variable transformation (from Tr to ln(Tr)), variable selection using best-subsets and stepwise approaches, and model assessment) is provided in the Supplementary Material. We evaluate the model by plotting $\frac{B_{abs}}{B_{ATN}}$ against aerosol properties not included in Eq. (9) (such as relative humidity and aerosol geometric mean diameter, which have been previously reported to bias corrections of filter-based B_{abs}, (Moteki et al., 2010; Nakayama et al., 2010; Schmid et al., 2006)). The results are presented in Fig. S5-S7.

As in V2005, iteration is required in our algorithm because B_{abs} is dependent on knowledge of SSA and AAE, which themselves are dependent on B_{abs} . We propose the following iterative process to update SSA and AAE in the model.

- 1. Initialize AAE from B_{ATN} across the three wavelengths ($B_{ATN} \sim \lambda^{-AAE}$) and initialize SSA for each wavelength using B_{ATN} from the filter-based absorption photometer and B_{scat} from a co-
- 369 located NEPH, i.e., $SSA(\lambda) = \frac{B_{scat}(\lambda)}{B_{scat}(\lambda) + B_{ATN}(\lambda)}$.
- Yield an initial set of coefficients G₀ through G₇ for each wavelength to calculate g(Tr, SSA, AAE) in Eq. (9), using one of the Algorithms described in Sect. 2.5.
- 372 3. Calculate B_{abs} for each wavelength using Eq. (8).
- 4. Update AAE and SSA using B_{abs} calculated in Step 3.
- 5. Derive a new set of coefficient values.
- 375 6. Iterate Steps 3-5 until converged.

Study	Aerosol source	SSA subset	Range of B _{abs} (Mm ⁻¹)	Filter-based instrument for B _{abs}	Reference instrument for B _{abs} ^a	Instrument for B _{scat} ^a	Coefficient values in the correction algorithm ^{b, c}		
The study in B1999	Lab-generated aerosols, including various mixtures of nigrosin and ammonium sulfate	0.5-1 (550 nm)	0-800 (550 nm)	One-λ PSAP (550 nm)	The difference between extinction (OEC) and scattering coefficient (NEPH) ^d	NEPH (450, 550, and 700 nm)	$\begin{array}{l} C_1 = 0.016 \pm 0.023 \; (550 \; nm) \\ C_2 = 1.55 \pm 0.25 \; (550 \; nm) \\ C_3 = 1.02 \pm 0.17 \; (550 \; nm) \end{array}$		
	Lab-generated				The average of the		467 nm	530 nm	660 nm
The laboratory study in V2005	aerosols, including various mixtures of kerosene soot, ammonium sulfate, and polystyrene latex	0.2-0.9 (530 nm)	0-800 (530 nm)	One-λ PSAP (550 nm), three-λ PSAP (467, 530, 660 nm)	PA (532 nm and 1064 nm) and the difference between extinction (OEC) and scattering coefficient (NEPH) d	NEPH (450, 550, and 700 nm)	$C_1 = 0.015$ $C_4 = 0.377 \pm 0.013$ $C_5 = -0.640$	$C_1 = 0.017$ $C_4 = 0.358$ ± 0.011 $C_5 = -0.640$	$C_1 = 0.022$ $C_4 = 0.352 \pm 0.013$ $C_6 = -0.674 \pm 0.013$
The ambient study in V2005	Ambient aerosols measured during RAOS and NEAQS ^e	0.75-1 (530 nm)	0-15 (530 nm)	One- λ PSAP (550 nm), three-λ PSAP (467, 530, 660 nm)	PA (532 nm and 1064 nm)	NEPH (450, 550, and 700 nm)	$\begin{array}{c} \text{C}_{6} = 0.640 \\ \pm 0.007 \\ \text{C}_{6} = 1.16 \pm \\ 0.05 \\ \text{C}_{7} = -0.63 \pm \\ 0.09 \end{array}$	$\begin{array}{c} \text{C}_{6} = 0.043 \\ \pm 0.007 \\ \text{C}_{6} = 1.17 \pm \\ 0.03 \\ \text{C}_{7} = -0.71 \pm \\ 0.05 \end{array}$	$C_{6} = 1.14 \pm 0.11$ $C_{7} = -0.72 \pm 0.16$
FIREX f	Biomass burning aerosols under relatively controlled laboratory conditions	0.2 -1 (550 nm)	38-1800 (550 nm)	CLAP (467, 529, 652 nm), TAP (467, 528, 653 nm), AETH (370, 470, 520, 590, 660, 880, 950 nm)	PAX (405 nm and 870 nm)	PAX (405 nm and 870 nm)	See Table 4 and Tables S6-S		s S6-S10
SGP (02/01/13 to 07/09/13) ^{f, g}	Ambient aerosols collected at the SGP user facility in Lamont, OK	0.75-1 (530 nm)	0-8 (550 nm)	CLAP (461, 522, 653 nm), PSAP (470, 522, 660 nm)	PASS (405, 532, and 781 nm)	NEPH (450, 550, and 700 nm)			

Table 2 Overview of the studies of B1999 and V2005 and the description of our experiments. 376

^a The operating wavelengths are based on the manufacturer specifications.

^b The coefficients provided in Table 2 are the values presented in Ogren (2010) and Virkkula (2010), which are updated from Bond et al. (1999) and Virkkula et al. (2005), 379 respectively.

380 ^c We reformulate the correction equations in the original publications to agree with Eq. (4)-(6) in this manuscript. C_1 to C_7 are the coefficients in the present work.

381 ^d OEC is optical extinction cell and PA is the instrument using photoacoustic technique.

382 ^e RAOS and NEAQS are Reno Aerosol Optics Study and New England Air Quality Study, respectively.

383 ^f The relative humidity (%) of the sampled aerosols from the FIREX and SGP study is 29.3 ± 3.3 and 26.1 ± 13.94 , respectively.

^g The scatterplot of AAE and SAE for the SGP data can be found in Fig. S15. Our results of AAE and SAE are compared to the values reported for different NOAA Earth 384

385 System Research Laboratory (ESRL) observational sites in Schmeisser et al. (2017).

377 378

386 2.5. Application of correction algorithms

In developing a procedure for applying our algorithm, we envision three potential scenarios:

3881. Algorithm A: The filter-based instrument is co-located with a NEPH and reference instrument389providing B_{abs} . This scenario facilitates the computation of G_0 through G_7 in Eq. (9) (step 2 in390the iterative process) as well as the derivation of new coefficients for existing correction391algorithms. This scenario can also enable the develop of a new a set of coefficients that may392be more appropriate for aerosol sources that we do not consider here.

- Algorithm B: The filter-based instrument is co-located with a NEPH but not a reference instrument providing B_{abs}, which is perhaps the most likely scenario (at least at many long-term monitoring sites). This scenario requires an initial guess of the coefficients; we provide sets of these in Table 4 below for different filter-based instruments and aerosol sources.
- 397 3. Algorithm C: The filter-based instrument is deployed with neither a co-located NEPH nor a 398 reference instrument providing B_{abs} . This scenario is the most challenging, because there are 399 no measurements of B_{scat} to compute SSA; to address this issue, we propose the use of a non-400 linear relationship between SSA and AAE (AAE = $a + b \times SSA^c$) to provide an initial guess of 401 SSA in the iterations.
- 402 To aid in decision-making between algorithms, we developed a flow chart for selecting appropriate

403 correction algorithm for CLAP, TAP, and PSAP (Fig. 1). Furthermore, Igor Pro (WaveMetrics,

404 Inc.) and Python (Version 3.7.5) based program for selecting and implementing our correction

405 algorithms can be found at https://doi.org/10.5281/zenodo.3742342.



406

- **Figure 1.** The flow-chart for the application of correction algorithms on PSAP, CLAP, and TAP.
- 408 Similar logic is followed for the AETH.
- 409 3. Results and discussion
- 410 3.1. Application of the previous algorithms on different aerosols

411 We first consider the application of the B1999 and V2005 corrections on different combinations

- 412 of aerosol type and filter-based absorption photometer. Specifically, we apply the two corrections
- to the biomass burning data from the FIREX laboratory campaign (CLAP and TAP) as well as six

months of ambient data from the SGP site (CLAP and PSAP). In doing so, we use the "default" 414 415 coefficients recommended in B1999 and V2005 as well as "updated" coefficients that are estimated via regression techniques. It is important to keep in mind that the updated coefficient 416 417 values of B1999 and V2005 (Table S7) are only valid for the aerosols investigated in this study. Future experiments are needed to systematically determine how the coefficients in B1999 and 418 V2005 may change for different aerosol types. We focus on the results of the CLAP in the main 419 420 text, because a CLAP is the only instrument common to deployments for both FIREX and SGP. The results of the TAP from FIREX and the PSAP from the SGP site can be found in the 421 Supplementary Material (Table S5 and Fig. S11). 422

- 423 Our inter-comparison between the corrected CLAP-derived B_{abs} and reference B_{abs} for the FIREX
- and SGP data is provided in Fig. 2 and Table 3. For the FIREX measurements, both analyses (using
- the "default" coefficients and updating the coefficients) suggest good correlation (coefficient of (22)
- 426 determination $(R^2) > 0.9$) between the CLAP and the reference across all three wavelengths. 427 Nevertheless, the corrections using the "default" coefficients result in over-prediction of B_{abs} by
- 427 Revenueless, the concertons using the default coefficients result in over-prediction of D_{abs} by 428 factors of ~2.5. If we update the coefficients in the corrections, there is an obvious improvement
- in the agreement (i.e., slope ≈ 1 ; R² increases). The results are generally similar for SGP, although
- 430 the R² for ambient data is generally lower for ambient data (R² < 0.7). Decreased R² may be due
- 431 to the lower aerosol concentrations measured in ambient air, which could lead to lower signal-to-
- 432 noise in the instruments. Moreover, it is worth mentioning that for both datasets (FIREX and SGP),
- 433 the corrected B_{abs} from different filter-based absorption photometers using the "default"
- 434 approaches does not agree with each other (slopes range from 0.69 to 1.40). However, after435 updating the coefficients, the slopes approach unity (Table S6).



436

Figure 2. Inter-comparison between the CLAP-derived B_{abs} corrected by the B1999 and V2005
 algorithms and the reference B_{abs} at 652, 528, and 467 nm for both FIREX and SGP data. The solid
 lines represent linear regressions, while the dashed line is a 1:1 line.

Table 3 Relationship between the CLAP-derived B_{abs} corrected by the B1999 and V2005 algorithms (including updated coefficients) and the reference B_{abs} at 652, 528, and 467 nm. The relationship is achieved using major axis regression (Ayers, 2001). The value in parentheses represents the coefficient of determination (R^2) of the linear relationship.

		652 nm	528 nm	467 nm
	B1999	y = -39 + 2.69x (0.94)	y = -49 + 2.50x (0.96)	y = -45 + 2.26x (0.97)
EIDEV	V2005	y = -46 + 2.83x (0.96)	y = -57 + 2.75x (0.96)	y = -56 + 2.68x (0.96)
FIREA	B1999 (update coeffs)	y = -8.4 + 1.02x (0.96)	y = -7.7 + 1.01x (0.97)	y = -3.4 + 1.00x (0.96)
	V2005 (update coeffs)	y = -9.4 + 1.03x (0.97)	y = -7.3 + 1.01x (0.97)	y = -3.0 + 1.00x (0.96)
SGP	B1999	y = -2.60 + 3.77x (0.41)	y = -1.90 + 3.20x (0.49)	y = -0.98 + 2.85x (0.55)
	V2005	y = -2.50 + 3.54x (0.41)	y = -2.00 + 3.15x (0.48)	y = -1.10 + 2.96x (0.55)
	B1999 (update coeffs)	y = -0.29 + 1.10x (0.60)	y = -0.29 + 1.08x (0.63)	y = -0.17 + 1.03x (0.65)
	V2005 (update coeffs)	y = -0.57 + 1.24x (0.65)	y = -0.50 + 1.15x (0.67)	y = -0.27 + 1.06x (0.67)

444

In the FIREX data, there is an apparent dependency of the updated coefficients on the wavelength
of light, but more importantly, on the aerosol optical properties, namely SSA and AAE (Tables
S7-S9). However, in the ambient data from SGP, the dependency on optical properties is less

obvious (Tables S10-S11). Nevertheless, all of these coefficients differ from those reported in B1999 and V2005 (again, derived for the PSAP rather than the CLAP), which highlights the potential need to use coefficient values that are appropriate for the instrument being used, its wavelength(s) of light, and optical properties that are representative of the sampled aerosols when applying correction factors to B_{ATN} .

453 3.2. Application of the new algorithms to the FIREX data

The co-location of the CLAP, TAP, AETH, and PAX during FIREX allows us to apply each algorithm (A, B, C) to these data. Similar to Sect. 3.1, we focus our discussion on the CLAP with details on the TAP and AETH presented in the Supplementary Material (Fig. S11-S12). However, we provide the recommended initial guesses in the new algorithms and the comparison of absorption (corrected filter-based B_{abs} versus reference B_{abs}) for all filter-based absorption photometers in Table 4 and Table 5 to help readers quickly retrieve key information of our algorithms.

Figure 3 provides a comparison between the uncorrected B_{ATN} from the CLAP at all three 461 462 wavelengths, as well as photoacoustic Babs interpolated to those wavelengths using AAE. For each wavelength, the slopes are significantly greater than one. Moreover, there is an apparent 463 dependency on SSA and AAE in the agreement between the instruments. This is most obvious in 464 Fig. 3a (652 nm), where data with lower SSA and lower AAE (smaller markers, "brighter" colors) 465 fall below the best-fit line, while data with higher SSA and higher AAE (larger markers, "darker" 466 colors) fall above the best-fit line. This phenomenon is less clear in Fig. 3b-3c, but an apparent 467 468 dependancy on SSA and AAE remains, which highlights the need to include both of these aerosol optical properties (and appropriate interaction terms) when correcting B_{ATN} values. 469





Figure 3. Comparison of the uncorrected CLAP-derived B_{ATN} and the reference B_{abs} at 652, 528,
and 467 nm for the FIREX data. The data points are colored by the corresponding SSA at the given
wavelength. The size of data points reflects their AAE quantified by the two PAX. The solid line
represents the linear regression, while the dashed line is a 1:1 line.

We first apply "Algorithm A" to the CLAP B_{ATN} data in Fig. 3. Using the reference B_{abs} values from the PAX (in addition to B_{scat} values), we are able to derive a set of coefficients that enable

477 the correction of the data (Table 4). Corrected CLAP values are presented in Fig. 4 with the linear

relationships presented in Table 5. The slope for each wavelength is very close to the 1:1 line,

suggesting that our approach works well in correcting these data. Moreover, the heteroscedasticity

that exists in Fig. 3 has been minimized after correction, and there are no apparent trends in how

the data are organized in Fig. 4 due to the aerosol optical properties.

482	Table 4 Coefficient values for Eq. (9) derived using "Algorithm A". We recommend these as the
483	initial guesses when implementing "Algorithm B" ^a .

		G₀	G 1	G ₂	G3	G4	G₅	G ₆	G 7
	652 nm	0.27	-0.16	-0.18	-0.05	0.18	0.08	-0.01	0.03
CLAP (FIREX)	528 nm	0.30	-0.28	-0.18	-0.07	0.25	0.10	0.13	-0.17
	467 nm	0.32	-0.38	-0.20	-0.08	0.33	0.12	0.24	-0.31
	652 nm	0.45	-0.45	0.07	-0.19	0.94	0.10	0.26	-0.35
TAP (FIREX)	528 nm	0.54	-0.51	0.02	-0.26	0.76	0.20	0.38	-0.44
	467 nm	0.62	-0.59	-0.07	-0.32	0.73	0.29	0.53	-0.60
	652 nm	0.37	-0.18	-0.34	-0.11	0.30	0.18	-0.36	0.41
CLAP (SGP)	528 nm	0.40	-0.15	-0.42	-0.14	0.10	0.24	-0.17	0.25
	467 nm	0.43	-0.16	-0.45	-0.16	0.07	0.27	-0.06	0.12
	652 nm	0.24	0.35	-0.16	-0.04	-0.47	0.07	-0.57	0.73
PSAP (SGP)	528 nm	0.30	0.48	-0.26	-0.10	-0.67	0.17	-0.63	0.77
	467 nm	0.35	0.49	-0.34	-0.15	-0.69	0.23	-0.55	0.79
	950 nm	0.47	0.17	0.01	-0.27	-0.4	0.25	-0.12	0.27
	880 nm	0.34	0.13	0.13	-0.17	0	0.10	-0.13	0.12
	660 nm	0.28	0.09	0.11	-0.12	0.15	0.05	-0.12	0.03
AETH (FIREX)	590 nm	0.16	-0.08	0.26	-0.03	0.59	-0.08	-0.02	-0.19
	520 nm	0.16	-0.05	0.14	-0.01	0.54	-0.07	-0.02	-0.21
	470 nm	0.14	-0.05	0.06	0	0.53	-0.05	-0.02	-0.17
	370 nm	0.13	-0.09	0.11	0	0.59	-0.06	-0.01	0.01

^a The coefficients derived from FIREX may be more appropriate for biomass burning aerosols, and
 the coefficients derived from SGP may be more appropriate for rural background environments in
 the absence of marine aerosols and dust (see Fig. S15).

487



488

Figure 4. As in Fig. 3, but the CLAP-based B_{ATN} values have been corrected using our "Algorithm
 A".

491**Table 5** Relationship between the filter-based B_{abs} corrected by "Algorithm A" and the reference492 B_{abs} at the operating wavelengths for the filter-based instrument. The relationship is achieved using

493 major axis regression (Ayers, 2001). The value in the parentheses represents the coefficient of 494 determination (\mathbb{R}^2) for the linear relationship.

		652 nm		528	3 nm	467 nm		
	CLAP	y = -7.8 + 1.02x (0.9	8)	y = -6.2 +	1.01x (0.98)	у	y = -3.2 + 1.00x (0.98)	
FIREA	TAP	y = -10 + 1.00x (0.87	7)	y = -13 + ().99x (0.87)	у	r = -16 + 0.99x (0.88)	
COD	CLAP	y = -0.25 + 1.08x (0.6	68)	y = -0.21 + 1.05x (0.67) y =			= -0.04 + 0.99x (0.68)	
30F	PSAP	y = -0.28 + 1.10x (0.4	3)	y = -0.24 +	1.06x (0.55)	y = -0.07 + 1.00x (0.62)		
		950 nm		880 nm	660 nm		590 nm	
EIDEY	AETH	y = -3.19 + 1.01x (0.82)	y = -3.9	2 + 1.02x (0.85) y = -5.97 + 1.03		k (0.88)	y = -5.63 + 1.02x (0.90)	
FIREA		520 nm	470 nm		370 nm		-	
		y = -2.36 + 0.99x (0.90)	y = 2.9	3 + 0.95x (0.88)	y = 18.38 + 0.89	x (0.80)	-	

495

496 We next investigate the repeatability of the coefficient values presented in Table 4 by randomly 497 selecting half of the measurements (N = 1338) from the whole FIREX dataset. By implementing "Algorithm A" to the extracted observations, we obtain new coefficient values for G_0 to G_7 . This 498 is repeated 1000 times to obtain a distribution of coefficient values (Fig. S13). The extraction 499 500 approach mimics the process of obtaining new biomass burning datasets, so that we can estimate the variability of these derived coefficients. From Fig. S13 and Table S12, the derived coefficients 501 502 are mostly insensitive to the different randomly-extracted datasets; most of the quartile deviation (defined as (Q3-Q1)/2, where Q1 and Q3 are the first and third quartile respectively) is within 0.05, 503 except G₄ which has a quartile deviation of ~0.08. Consequently, the coefficient values obtained 504 in "Algorithm A" appear to be reasonable initial guesses to correct filter-based absorption 505 measurements during biomass burning events when the reference B_{abs} is unavailable, such as in 506 "Algorithm B" and "Algorithm C". 507

We next implement "Algorithm B" to the CLAP BATN data from Fig. 3 using the initial guesses of 508 the coefficients derived from "Algorithm A" (Table 4) along with reference B_{scat} values. To get a 509 sense of the variability in the results, we randomly select half of the data and applied the correction; 510 this process is repeated 1000 times. For each iteration, we compare the corrected B_{abs} from the 511 CLAP to the reference B_{abs} from the PAX; the resulting slope, intercept, and R^2 values are 512 summarized as box-and-whisker plots in Fig. 5. For all three wavelengths, the slopes are close to 513 unity, and there is good correlation between the two absorption measurements ($R^2 \approx 0.98$), which 514 indicates that the good performance seen in Fig. 4 is independent of the reference Babs 515 measurements and our algorithm is able to correct "new" BATN. Consequently, when scattering 516 measurements are co-located with filter-based absorption measurements, our new correction 517 algorithm performs well. 518



Figure 5. The box-and-whisker plots for the slope, intercept, and R² of the relationship between the CLAP-derived B_{abs} (corrected by "Algorithm B" in the present work) and PAX-derived B_{abs} for all three wavelengths. For details on how these values were generated, please refer to the text.

519

Lastly, we apply "Algorithm C" to the data in Fig. 3. However, we first require a functional 523 relationship between AAE and SSA, because in this scenario, the CLAP BATN values are the only 524 data input to the algorithm (and therefore, SSA is unknown). Liu et al. (2014) proposed that a 525 power function can describe this relationship (AAE = $a + b \times SSA^{c}$); we present these data from 526 FIREX along with power function fits (and associated prediction intervals) in Fig. 6. To define 527 AAE in this figure, we fit a power-law relationship to the three B_{ATN} values from the CLAP; 528 similarly, we define SSA using interpolated B_{scat} from the PAX and B_{ATN} from the CLAP (The 529 rationale for using BATN is that if "Algorithm C" were to be implemented in practice, only BATN 530 would be available). In Fig. 6, the data points are colored by "prediction error", effectively a metric 531 to quantify how well the power function reproduces the individual data points. Although there is 532 a fair amount of error in some of these points, we still obtain an SSA-AAE relationship required 533 to initialize "Algorithm C". 534



Figure 6. AAE plotted against SSA for the FIREX data. In the figures, AAE was computed using a
 power-law fit across all three wavelengths, and SSA was computed using the interpolated B_{scat}

from the two PAX and the reported B_{ATN} from the CLAP. The data points are colored by their prediction error (("true" AAE - "calculated" AAE)/ "calculated" AAE).

540 Even though there is uncertainty in the SSA vs. AAE relationship used in "Algorithm C", after corrections have been applied, the filter-based B_{abs} for the CLAP agrees well with the independent 541 reference B_{abs} ; the slopes for all wavelengths are slightly greater than 1 (1.03-1.05) and the R^2 542 values are all high (0.97-0.98). However, even though the absorption measurements are corrected 543 well, there still remains large uncertainties in values of inferred scattering. Examples of this are 544 provided in Fig. 7, where we compare the SSA inferred from the PAX to the SSA inferred from 545 546 "Algorithm C" as well as B_{scat} for each wavelength. Generally speaking, data that are better represented by the SSA vs. AAE relationship (i.e., smaller prediction error) results in better 547 agreement with the reference for both SSA and B_{scat}, but there is also a clear divergence from the 548 1:1 line in Fig. 7a-c as SSA decreases. Therefore, even though "Algorithm C" performs well at 549 correcting filter-based BATN to agree with the reference Babs, estimates of final SSA values should 550 be considered to be uncertain. 551



Figure 7. Comparison of SSA (a-c) and B_{scat} (d-f) at the three wavelengths for the FIREX data. Vertical axis: values output from "Algorithm C"; horizontal axis: values calculated using the photoacoustic B_{abs} and B_{scat}.

552

To explore the generalization of the new algorithms across different instruments, we next apply our algorithms to the other filter-based absorption photometers operated during the FIREX study (TAP and AETH). Consistent with what we observed for the CLAP results, the corrected TAPand AETH-derived B_{abs} is in good agreement with the photoacoustic B_{abs} (as demonstrated in Table 4 and Table 5, as well as Fig. S11-S12). Moreover, the corrected B_{abs} from the three filter-based instruments agrees with each other for all three wavelengths (Table 6), confirming the consistency of our algorithm.

563 **Table 6** Inter-comparison between different filter-based B_{abs} corrected by "Algorithm A" in the

present work. The value in the parentheses represents the coefficient of determination (R^2) of the linear relationship.

	FIREX: CLAP vs. TAP	FIREX: CLAP vs. AETH	FIREX: TAP vs. AETH	SGP: CLAP vs. PSAP
652 nm	y = 1.84 + 1.02x (0.89)	y = 4.17 + 0.94x (0.87)	y = -0.31 + 0.99x (0.82)	y = -0.04 + 0.99x (0.70)
528 nm	y = 5.75 + 1.02x (0.88)	y = 3.70 + 0.91x (0.85)	y = -6.38 + 0.98x (0.82)	y = -0.11 + 1.02x (0.73)
467 nm	y = 10.57 + 1.01x (0.88)	y = 0.45 + 0.98x (0.83)	y = -13.62 + 1.04x (0.79)	y = -0.11 + 1.02x (0.76)

566

567 3.3. Application of the new algorithms to ambient data

To test our algorithms further, we extended our work to ambient data collected the DOE SGP site during the time period which the PASS-3 was operational. From the SGP data, we derived a different set of coefficients for ambient data using "Algorithm A", which differ from those derived for FIREX (Table 4). The results presented in Fig. 8 and Table 5 suggest that our new algorithm works at least as well as B1999 and V2005 on this dataset (both with updated coefficients). The repeatability of the coefficient values in "Algorithm A" is confirmed for the SGP measurements using the same procedure as described in Sect. 3.2 (see results in Fig. S13 and Table S12).



575

576 **Figure 8.** Inter-comparison between the CLAP-derived B_{abs} corrected by "Algorithm A" in the 577 present work and reference B_{abs} at 652, 528, and 467 nm for the ambient data at the SGP study 578 area. The solid line represents the linear regression, while the dashed line is a 1:1 line.

579 On the SGP data, we see similar performance to the FIREX data when we apply "Algorithm B", 580 where we again sampled half of the CLAP data, used the initial guesses derived in "Algorithm A", 581 and repeated this process 1000 times. Although the slopes tend to be larger than 1 (i.e., the 582 corrected CLAP B_{abs} remains high relative to the PASS B_{abs}), the results still represent an 583 improvement over B1999 and V2005 using their recommended coefficients for their correction 584 equations.

Implementing "Algorithm C" is challenging for ambient data, because there is no distinct power
function relationship in AAE vs. SSA (Fig. 9); this is consistent with other field studies reporting
both SSA and AAE (e.g., Backman et al. (2014) and Lim et al. (2018)). Our approach described

here is only appropriate for ambient aerosols that follow a power function, such as sites impacted by biomass burning. Nevertheless, we did apply this to a subset of the SGP data where the AAE-SSA prediction error is within 30% (N = 86), and for this subset of data, "Algorithm C" works fairly well (slopes ≈ 0.95 ; see Fig. S14). Therefore, while "Algorithm C" may have utility for ambient data, we advise caution when using this algorithm since the aerosols influencing the site may not be represented by a clear AAE-SSA power function (e.g., when biomass burning and coarse aerosols are equally prevalent at a long-term monitoring site).



595

Figure 9. AAE plotted against SSA for the SGP ambient data. The power law fit (AAE = $a + b \times SSA^c$) is performed on SSA (SSA = $B_{scat}/(B_{scat} + B_{ATN})$) and AAE computed by three-wavelength B_{ATN} .

These new algorithms are also applicable to the PSAP deployed at the SGP site. The results of the correction for the PSAP are presented in Table 5 and Fig. S11, and the recommended initial guesses when implementing "Algorithm B" to PSAP-B_{ATN} at ambient environments are given in Table 4. As expected, there is good agreement between corrected PSAP- and CLAP-B_{abs} (Table 6).

602 To investigate if our algorithms are suitable to correct Babs obtained from different ambient environments, the aerosol properties from the SGP site are compared to those from the other 603 NOAA/ESRL observational sites. We use the similar "AAE-SAE space" as that in Cappa et al. 604 605 (2016) and Schmeisser et al. (2017) to infer dominant aerosol types (e.g., BC, BrC, dust, mixed dust/BC/BrC, see Fig. S15). Our results of AAE and SAE overlap with most of the values from 606 the NOAA/ESRL sites, except when marine aerosols or dust contribute to the local aerosol 607 608 emissions. Though none of the NOAA/ESRL sites fall into the clusters of BC or BC/BrC, some of our data can represent the optical properties of aerosols from these clusters. Therefore, we highlight 609 that our algorithm developed by the SGP data may have the potential to be generalized to a variety 610 611 of environmental conditions, but we would need to validate this using observations from more studies. 612

613 3.4. Impact of the implemented correction algorithm on aerosol optical properties

In addition to the direct comparisons of B_{abs} between the filter-based and photoacoustic measurements, we compare derived optical properties (AAE and SSA) from different instruments to assess the algorithms' performance on derived aerosol optical properties. For example, we have discussed the discrepancy of SSA between the filter-based and photoacoustic measurements when implementing "Algorithm C" in Sect. 3.2. In this section we will more broadly discuss the impact

of different correction algorithms on AAE and SSA.

620 In Fig. 10, we present the frequency distribution of AAE for both FIREX and SGP data generated 621 from different campaign/instrument pairs using different correction approaches. For the FIREX data (Fig. 10a-b), most corrections (with the exception of the "default" B1999) are consistent with 622 623 the photoacoustic data, while for the SGP data (Fig. 10c-d), most corrections (with the exception of "default" V2005) are consistent with the photoacoustic data. However, updating the coefficients 624 for B1999 and V2005 improves the agreement with the photoacoustic data. The 50% difference 625 that exists between the B1999 and V2005 algorithms in all panels in Fig. 10 are consistent with 626 previous studies. For example, both Backman et al. (2014) and Davies et al. (2019) found that the 627 V2005-derived AAE is greater than B1999-derived AAE by 33% to 50% for ambient aerosols. 628 Therefore, we highlight that the default coefficients in B1999 and V2005 may have some 629 limitations when deriving AAE using the corrected B_{abs}; instead, updating the coefficients or using 630 the new algorithm proposed in this work may yield more robust AAE results. 631



632

Figure 10. The frequency distribution of AAE calculated for different instrument/correction
 combinations of multi-wavelength B_{abs}.

635 Similar to Fig. 10, we also investigate the distribution of SSA computed by using corrected B_{abs} 636 along with B_{scat} . We provide the results at 652 nm as an example in the main text (Fig. 11); figures 637 for 528 nm and 467 nm can be found in the Supplementary Material (Fig. S16 and S17). For both 638 FIREX and SGP data, the SSA obtained using the new algorithm agree very well with the B1999 639 and V2005 but only when their coefficients have been updated. Calculations of SSA using B1999 640 and V2005 with their recommended coefficients suggest that these values may be biased low, 641 which follows the over-estimation of corrected B_{abs} demonstrated in Fig. 2.



642

Figure 11. The frequency distribution of SSA (652 nm) calculated for different
 instrument/correction combinations of B_{abs} and B_{scat}.

Moreover, we plot similar figures as Fig. 10-11 using all algorithms (A, B, and C). As shown in Fig. S18, the results using "Algorithm B" agrees very well with those using "Algorithm A", but the use of "Algorithm C" results in some obvious discrepancies compared to the photoacoustic reference, again highlighting the potential for large uncertainty using this algorithm.

In Fig. 12, we directly compare the distributions of both AAE and SSA at 652 nm for all of the filter-based absorption photometers considered here, using our "Algorithm A" to correct the B_{ATN} data. For both datasets, after the corrections have been applied, there are only marginal differences of the AAE (Fig. 10a and 10b) derived by different instruments. Similarly, there is good agreement among the SSA values when using corrected- B_{abs} from different instruments (Fig. 10c and 10d). Overall, the derived properties using the new correction are consistent across all instruments,

655 suggesting its generalizability.





Figure 12 The probability density of AAE and SSA (652 nm) derived by different filter-based photometers B_{abs} (corrected by "Algorithm A" in the present work). Note that the number of total observations vary across instruments.

660 3.5. Uncertainty of the new algorithms

In this section, we estimate the uncertainty of the new algorithms due to both measurement uncertainties of the instruments and the uncertainties of parameter computation. We then simulate the propagated uncertainty in the corrected filter-based B_{abs} reported in this paper.

664 Measurement uncertainties of the instruments considered here have been reported in previous work (e.g., (Anderson et al., 1996; Nakayama et al., 2015; Ogren et al., 2017; Sherman et al., 2015)) and 665 are summarized in Table 1. The typical sources of measurement uncertainty of the aerosol 666 instruments include: 1) instrument noise (often associated with the averaging time); 2) calibration 667 668 uncertainties (such as the accuracy of the operating wavelengths and the properties of the calibration materials); 3) standard temperature and pressure (STP) correction uncertainties 669 (Sherman et al., 2015); and 4) flow rate uncertainties. Additional uncertainties that are specific to 670 filter-based absorption photometers include spot size and filter medium corrections (Bond et al., 671 1999; Ogren et al., 2017). Regardless, these values all tend to be $\leq 30\%$, which is consistent with 672 other commonly-used aerosol instrumentation. 673

Because correction algorithms for filter-based absorption instruments also require aerosol optical 674 properties, the algorithms' performance will be affected by these values as well. For example, 675 uncertainties in SSA are directly related to uncertainties associated with Babs and Bscat, which are 676 both included in our simulations. However, capturing uncertainties in AAE is more complex, as 677 AAE can be computed by either " 2λ fit" (a linear fit using B_{abs} at two wavelengths) or " 3λ fit" 678 (same as the power fit used in the present work). Davies et al. (2019) used the 3λ fit to calculate 679 AAE and compared this to calculations using 662 nm and 785 nm (i.e., AAE_{662/785}), finding that 680 the 3λ results was about 50% greater. Moreover, similar differences (-35% to 85%) can exist 681 comparing two different 2λ combinations (AAE_{440/870} and AAE_{675/870}), depending on the 682 contribution of brown carbon to absorption at 440 nm (Wang et al., 2016). However, based on Fig. 683 S19 and S20, we demonstrate small ($< \sim 10\%$) differences in the calculated values of AAE using 684

685 our Algorithm A using different 2λ combinations for linear fits and the 3λ power-law fit, when 686 considering both FIREX and SGP data. Consequently, we do not include AAE calculation 687 uncertainty in our simulation.

In our simulations, the propagated uncertainty of corrected B_{abs} is estimated by implementing the 688 new algorithm to datasets in which filter-based BATN, reference Babs, and Bscat are subject to 689 measurement uncertainties. The full procedure is outlined in the Supplementary Material, but we 690 691 provide a brief overview of our Monte Carlo approach here. First, we create a synthetic dataset (n 692 = 500 records) that defines B_{abs} at 652 nm and AAE that is intended to represent biomass burning. Values of BATN and SSA are then computed using the relationships presented in Fig. 3 and Fig. 6, 693 respectively. Respective uncertainties associated with each of these values are applied following 694 695 Table 1, assuming that these follow a normal distribution. We then applied "Algorithm B" to the BATN dataset, repeated 1000 times, to quantify overall uncertainty associated with our correction 696 697 algorithm.

Figure 13 provides a graphical summary of our uncertainty simulation results, which was derived 698 by fitting linear equations to the "true" Babs value (that we defined) and the "corrected" Babs values 699 700 (outputs of each iteration). Considering the slopes (Fig. 12a), our algorithm can generally reproduce the "true" value within 10% at 652 nm and 528 nm, but the performance is slightly 701 degraded at 467 nm. The median intercept for our simulations is close to zero, but the interquartile 702 range increases with decreasing wavelength (Fig. 12b), suggesting that the uncertainty may 703 increase at shorter wavelengths. The coefficients of determination (Fig. 12c) range from 0.47 (652 704 705 nm) to 0.68 (467 nm), showing that the algorithm may be less precise if large measurement uncertainties exist. Even though these sources of uncertainty exist when implementing our 706 correction algorithms and propagate through to the corrected values, we argue that our new 707 algorithm will "standardize" uncertainties across corrected Babs values from filter-based absorption 708 photometers. Moreover, the new algorithms perform, at least, better than the previous algorithms 709 with "default" coefficients, or as well as the previous algorithms with updated coefficients. 710



711

Figure 13. The box-and-whisker plots (slope, intercept, and R²) for the Monte Carlo simulation of
 the relationship between the CLAP-derived B_{abs} (corrected by "Algorithm B" in the present work)

714 and "true" B_{abs} for all three wavelengths.

715 4. Conclusions

Filter-based absorption instruments are widely used at global observational sites due to their 716 717 relatively low cost, fast response, and easy operation. Despite the existence of different correction algorithms to correct the filter-based Babs measurements, these are not "standardized" as 718 differences in corrected Babs values exist across different instrument/correction combinations, even 719 when the instruments are co-located. This study provides a systematic evaluation of the previous 720 721 correction algorithms (B1999 and V2005 corrections) on the CLAP and similar instruments (TAP 722 and PSAP) using both laboratory-generated biomass burning emissions and ambient aerosols. We also developed new correction algorithms that are applicable to any filter-based absorption 723 photometer (e.g., PSAP, CLAP, TAP, AETH), which will have utility for any historic or future 724 725 filter-based absorption measurements and which have the potential to standardize absorption coefficients across all filter-based instruments. This latter point is demonstrated in Table 6 and Fig. 726 727 12 in that there is good agreement across all filter-based absorption photometers when applying our corrections to both biomass burning and ambient data. In practice, we anticipate that our 728 Algorithm B will be most common, because at long-term monitoring sites, filter-based absorption 729 photometers are typically co-located with a nephelometer. 730

Using the existing corrections on our CLAP measurements, we find that the corrected Babs 731 overestimate photoacoustic B_{abs} by factors of ~2.6 (biomass burning aerosols) and ~3.2 (ambient 732 aerosols). Similar overestimations of absorption by filter-based instruments are seen in the results 733 of TAP from the FIREX study and PSAP deployed at the SGP. Comparing between B1999 and 734 V2005, B_{abs} corrected by the two corrections differ by -6% to 18%. These discrepancies in our 735 results are consistent with those reported for the inter-comparisons between filter-based and 736 photoacoustic absorption instruments (e.g., (Arnott et al., 2003; Davies et al., 2019; Li et al., 2019; 737 Müller et al., 2011a)). 738

Overall, our new developed algorithms (A, B, and C) perform well on correcting B_{abs} for different 739 filter-based absorption photometers (CLAP, TAP, PSAP, and AETH) from both biomass burning 740 and ambient measurements. Our work suggests that if the filter-based instrument is co-operated 741 with a reference absorption instrument and a NEPH at field for a period, researchers can compute 742 site-specific initial guesses (same as "Algorithm A" in the present work). Otherwise, either 743 "Algorithm B" or "Algorithm C" proposed in this paper can be used to correct the filter-based 744 measurements. In "Algorithm B" when a filter-based absorption photometer is co-located with a 745 NEPH but without a reference instrument, the set of coefficients yield in this work (Table 4) can 746 be used as initial guesses to implement the algorithm. In "Algorithm C" when a filter-based 747 absorption photometer is operated by itself, a "representative" relationship between AAE and SSA 748 can be used to estimate SSA from AAE at each step in the iterative process, but we advise caution 749 if this relationship is not monotonic (e.g., as in the ambient data from SGP and from Backman et 750 al. (2014) and Lim et al. (2018)). The only scenario not included in the present work is that the 751 filter-based absorption photometer is co-located with a reference absorption instrument, but no 752 instrument for scattering. However, under this scenario, one could simply use the photoacoustic 753 Babs data because no filter-induced biases exist for those instruments. 754

In terms of the aerosol optical properties (AAE and SSA) computed by different corrections, the
new algorithm suggests no bias of AAE and SSA when compared to that derived by updatedB1999and updated-V2005 for both aerosol datasets.

758 However, the new algorithm is not without limitations. First, we used the photoacoustic B_{abs} as the 759 reference to develop the algorithm and the initial guess of the coefficients; meanwhile, some studies argue that photoacoustic absorption is not a "ground truth" (e.g., (Lack et al., 2006; Lewis 760 761 et al., 2008)). Thus, we simulate the propagated uncertainty of our algorithms considering the measurement uncertainties due to the photoacoustic Babs (as well as BATN and Bscat) and find that 762 the corrected B_{abs} can be biased by -17% to 5%, depending on the operated wavelength. Although 763 potential bias due to the precision of photoacoustic Babs cannot be excluded, using our new 764 algorithm to correct the filter-based B_{abs} will at least eliminate correction-related biases among 765 different filter-based instruments. Second, we only tested the algorithms with data from biomass 766 burning and ambient measurements. It is unclear how the algorithms will work for other absorbing 767 aerosols (e.g., dominated by fossil fuel emissions or mineral dust). Further evaluation of the 768 performance of the new algorithm on other aerosol sources may help to address this issue. 769 Regardless, we argue that our approach can standardize reported absorption coefficients at long-770 term monitoring sites, which has the potential to yield a better data set with which to evaluate 771 chemistry-climate models. 772

Code and data availability. The code for the algorithm was initially written in Igor Pro 773 774 (WaveMetrics Inc.) and has been developed to be accessible through either Python (Version 3.7.5) Pro. The implementation of algorithm 775 or Igor our is available at https://doi.org/10.5281/zenodo.3742342. The FIREX aerosol products are available 776 at https://esrl.noaa.gov/csd/project/firex. The SGP aerosol products 777 are available at https://www.archive.arm.gov/discovery/ (February 2013–July 2013, 36° 36' 18.0" N, 97° 29' 6.0" 778 W: Southern Great Plains Central Facility, data set accessed 01/16/19). 779

Author contributions. HL developed the model, conducted the evaluations, and drafted the manuscript. AAM and GRM edited the manuscript and obtained funding.

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

Acknowledgements. The authors would like to thank Allison Aiken (Los Alamos National Laboratory) for useful discussions regarding the SGP PASS-3 data. The authors greatly acknowledge Vanessa Selimovic, and Robert Yokelson from University of Montana for lending us the PAX-405; Patrick Sheridan, John Ogren, and Derek Hageman from NOAA for lending us the CLAP; and Anthony Prenni from the National Park Service for lending us the AETH during the FIREX campaign. We also thank Nick Good (Colorado State University), Jim Roberts (NOAA) and Carsten Warneke (NOAA) for their on-site support at FSL.

Financial support. The authors and FIREX data were supported by NOAA Climate Program
Office Grant NA16OAR4310109. The ambient data at SGP site in Lamont (OK, USA) were
obtained from the atmospheric radiation measurement (ARM) user facility, a U.S. Department of
Energy (DOE) Office of Science user facility managed by the Office of Biological and
Environmental Research.

Review statement. The authors thank Associate Editor Paolo Laj for handling this submission and
 the anonymous reviewers for their valuable comments.

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