Effect of Large-scale Biomass Burning on Aerosol Optical Properties at the GAW Regional Station Pha Din, Vietnam

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ABSTRACT

In 2014, Pha Din (1466 m a.s.l.) was established as a Global Atmosphere Watch (GAW) regional station for aerosol and trace gas measurements in northwestern Vietnam. This study presents a five-year climatology of aerosol optical properties derived from nephelometer and aethalometer measurements and a comparison with ground-based remote sensing measurements at the nearby AERONET station Son La. The annual variations of the aerosol measurements at Pha Din are clearly dominated by annually recurring periods with high biomass burning activity in northern Southeast Asia (February–May). During these periods, the majority of air masses arriving at Pha Din originate from the southwest (northern Thailand, Laos and Myanmar). Both the meteorological conditions and the aerosol optical properties are very similar during the individual high biomass burning periods (increased temperature: > 20°C; moderate ambient relative humidity: 60–70%; decreased single scattering albedo: 0.8–0.9; increased absorption Ångström exponent: 1.6–2.0; and scattering Ångström exponent significantly larger than 1). Prior to the biomass burning season (October–January), the meteorological conditions at Pha Din are influenced by the SE Asian monsoon, leading to a frequent transport of air masses from SW China with moderate aerosol loadings. The lowest pollution levels are observed from June to September, which represents the wet season.

Keywords: Aerosol optical properties; Biomass burning; Black carbon; Long-term measurements; Global Atmosphere Watch.

INTRODUCTION

Atmospheric aerosols influence the earth’s climate through aerosol-radiation and aerosol-cloud interactions, and they are also known to cause adverse health effects. Light scattering and absorption by atmospheric aerosols influence the solar and terrestrial radiation fluxes, and the change in net radiative flux by anthropogenic aerosol since the preindustrial time contributes to the radiative forcing. Our knowledge of aerosol optical properties such as scattering coefficient, absorption coefficient, single scattering albedo and directional dependence of light scattering, which are needed for radiative transfer calculations for estimating aerosol-radiation interactions (Boucher et al., 2013), still remains incomplete (Jaeglé et al., 2011) due to
the highly complex and variable characteristics of the atmospheric aerosol.

For a reliable and quantitative prediction of the net radiative forcing by aerosols, obtaining a 4-dimensional coverage of aerosol properties in time and space is crucial (Forster et al., 2007). On a local level, a thorough physical and chemical characterization of the near-ground aerosol is done in the existing international networks, for example, GAW (the Global Atmosphere Watch programme of the World Meteorological Organization), ACTRIS (the European Research Infrastructure for the observation of Aerosol, Clouds, and Trace Gases) or the NOAA (National Oceanic and Atmospheric Administration) Federated Aerosol Network, which all aim at providing harmonized data from stations around the globe (Andrews et al., 2011; Collaud Coen et al., 2013). In AERONET (Aerosol Robotic Network), EARLINET (the European Aerosol Research Lidar Network, now incorporated in ACTRIS) and other networks, vertically resolved measurements using lidars and column integrated measurements using sun photometers allow for the retrieval of relevant aerosol optical properties in the air column above ground for further comparison with satellite based aerosol products (Holben et al., 1998; Bösendorfer et al., 2001). Although both in-situ measurements and columnar measurements provide valuable stand-alone information about the composition and dynamics of the troposphere, only interlinking all measurement types at the same site and filling the regional gaps in the global measurement network will provide the full information that is crucially required by climate system models to assess the aerosol climate effects on all spatial scales (Lopatin et al., 2013; Kaskaoutis et al., 2014; Mann et al., 2014; Samset et al., 2014; Sarkar et al., 2018).

Northern Southeast Asia is an area that is well-known for being severely affected by intense seasonal biomass burning (BB) in the form of forest wildfires and agricultural crop burning as well as biofuel usage for cooking (Carmichael et al., 2003; Streets et al., 2003, 2009; Gautam et al., 2013). Myanmar, Thailand and Laos have been identified as source regions for BB in many existing studies (Yen et al., 2013; Krishna Prasad et al., 2015; Duc et al., 2016; Loftus et al., 2016; Pantina et al., 2016; Li et al., 2017; Lin et al., 2017). These show that in the northern part of Southeast Asia (15–25°N, 90–120°E), atmospheric mean flow patterns largely control the transport and dispersion of BB plumes. In the lower mid-troposphere (700–850 hPa), the streamlines are mostly southwesterly/southerly/southeasterly by an upward branch of the local east–west cell/circulation (Lin et al., 2013). In the boundary layer (925–850 hPa), the streamlines are mostly southwesterly/southerly/southeasterly due to the presence of mountain ranges located on the eastern and western coasts of the SE Asian peninsula and of the vast plain in the central peninsula (Chen et al., 2002; Lin et al., 2013). During the rest of the year, the meteorology in the northern SE Asian region is dominated by the NE monsoon (October–January), with frequent air mass transport from SW China (Lin et al., 2017), as well as by the SE monsoon (June–September) associated with frequent rainfall and high relative humidity.

In 2014, the GAW regional station Pha Din (PDI) was established in northwestern Vietnam at a rural background site in a mountainous area at 1463 m a.s.l., providing continuous measurements of ground-based in-situ aerosol parameters (aerosol absorption coefficient and aerosol scattering coefficient) and trace gases (CO2, CH4, CO and O3). The measurements at Pha Din are the first continuous aerosol monitoring activities in northern Vietnam and the entire surrounding region. The goal of this paper is to show the importance of the continuous monitoring at Pha Din for the long-term assessment of BB in the northern SE Asian area. BB and traffic source regions will be identified using air mass backward-trajectories, and the Pha Din data will be compared with other aerosol measurements that have been conducted in northern Vietnam in recent years on a non-continuous basis.

**METHODS**

**Location**

Pha Din, located in the northwest of Vietnam (see Fig. 1), is a mountain pass at the border of the Son La and Dien Bien provinces. The pass is embedded in a mountain chain with hills between 700 and 1600 m a.s.l., stretching from China into Vietnam in a northwest-to-southwest orientation. The Pha Din measuring site (GAW station acronym: PDI) is located on a hilltop close to the pass at 1466 m a.s.l. (21.573°N, 103.516°E) covered with forest. The station itself is above the canopy. It was established in 1962 as a meteorological station of the Viet Nam Meteorological and Hydrological Administration (VNMHA) and started providing official data in 1964. In February 2014, PDI was upgraded with continuous aerosol and trace gas measurements within the framework of the Capacity Building and Twinning for Climate Observing Systems (CATCOS) project (SDC, 2018), and the operation and data management was handed over to VNMHA in 2017 under Decision No. 29/QĐ-BTNMT of the Vietnamese Ministry of Natural Resources and Environment. There are no residents at the station except for the custodians, no relevant residential areas within 10–20 km except for sparse individual farmhouses, and no coal-fired power plants or other industrial activities in the region. The area is under the surveillance of the government; therefore, there is a low probability for extensive growth of residential or industrial activities around the station.

**Inlet System**

The aerosol measurements at Pha Din follow the GAW recommendations for aerosol sampling (WMO/GAW, 2016). The site was audited by the GAW World Calibration Centre for Aerosol Physics (WCCAP) in 2016. Results are available in an openly accessible audit report (WCCAP, 2016). The inlet is a custom made total suspended particulate matter...
Fig. 1. Geographical location of the sampling site (Images: Google Maps, Maphill), in three different zoom levels (A, B, and C).

(TSP) inlet, as recommended for stations that are frequently in clouds. In accordance with GAW guidelines, the sampling air was dried with dried, particle-free dilution air to maintain/achieve a relative humidity (RH) below 40% inside the instruments. During the period of measurement, the RH was 15–40% with a functioning dilution system (see Fig. S1 in the Supplementary Information). Periods with a malfunctioning dilution system were flagged accordingly. The measured ambient aerosol light absorption and scattering data were corrected by the effective dilution factor (set to be around 2) and are reported for standard temperature and pressure (STP) conditions (1013 mbar and 0°C) in this study.

Aerosol Light Absorption and BC Source Apportionment

The aerosol absorption coefficient \( b_{\text{abs}} \) was determined from aethalometer measurements (Model AE31; Magee Scientific Inc.) at 7 wavelengths (370, 470, 520, 590, 660, 880 and 950 nm), with a native time resolution of 5 minutes. The aethalometer raw data processing followed the current GAW/ACTRIS recommendations (Lioussse et al., 1993; Petzold et al., 1997; Weingartner et al., 2003; Collaud Coen et al., 2010; Park et al., 2010; Bond et al., 2013; Drinovec et al., 2015; WMO/GAW, 2016; Zanatta et al., 2016) and includes compensation for non-linear filter loading effects and correction for multi-scattering effects (Weingartner et al., 2003) which are described in detail in the Supplementary Information of this article. The correction for multi-scattering effects is achieved using a specific C-value (Zotter et al., 2017). Lacking simultaneous measurements with an absorption reference method, a constant and wavelength independent C-value of 3.5 was applied in this study, following the GAW recommendations (WMO/GAW, 2016; Zanatta et al., 2016), to infer the absorption coefficients \( b_{\text{abs}} \) from the loading compensated attenuation coefficients measured by the aethalometer.

The spectral dependence of light absorption can be used to apportion aerosol light absorption and particulate BC mass to biomass burning and traffic emissions, if these are the only sources contributing to light absorbing aerosol (Kirchstetter et al., 2004; Sandradewi et al., 2008). We applied this “aethalometer model” approach to the absorption coefficients \( b_{\text{abs}} \) measured at the wavelengths 470 and 950 nm using the following equations (Sandradewi et al., 2008; Zotter et al., 2017):

\[
b_{\text{abs}, \text{BB}} (950 \text{ nm}) = \frac{b_{\text{abs}} (470 \text{ nm}) - b_{\text{abs}} (950 \text{ nm})}{\left(\frac{470}{950}\right)^{\alpha_{\text{a}}} - \left(\frac{470}{950}\right)^{\alpha_{\text{a}}}}
\]
where \( \alpha \) is the absorption Ångström exponent. BB and TR denote the sources “biomass burning” and “traffic” to be apportioned. The aethalometer model is based on the fact that typical traffic emissions contain BC as the exclusive light absorbing species, whereas biomass burning emissions contain co-emitted brown carbon, which results in a higher \( \alpha \). A recent study (Zotter et al., 2017) referenced the aethalometer model against \(^{14} C\)-based source apportionment and found optimum \( \alpha \)-values of \( \alpha_{\text{TR}} \approx 0.9 \) and \( \alpha_{\text{BB}} \approx 1.68 \) for the above wavelength pair. However, this \( \alpha_{\text{BB}} \) is specific to wintertime Switzerland, where wood burning for domestic heating is the dominant biomass burning source. The spectral dependence of the typical biomass burning aerosol at Pha Din, which originates from wildfires, agricultural crop burning and the use of biofuel for cooking, may be different. Indeed, the \( \alpha \)-values of the whole aerosol were as high as \( \sim 2 \) when wood burning influence dominated, thereby clearly exceeding \( \alpha_{\text{BB}} \approx 1.68 \). Applying \( \alpha_{\text{BB}} \approx 1.68 \) in the aethalometer model would consequently result in negative traffic contribution. Therefore, we chose \( \alpha_{\text{BB}} \approx 1.8 \) instead. The uncertainty of \( \alpha_{\text{BB}} \) will be shown and discussed in the context of Fig. 4 later in this article. The respective contributions of biomass burning and traffic to the equivalent black carbon mass concentrations \( (m_{\text{BC}}) \) were then calculated as:

\[
m_{\text{BC,TR}} = \frac{b_{\text{abs,TR}}(950 \text{ nm})}{\text{MAC}_{\text{TR}}(950 \text{ nm})}
\]

(3)

\[
m_{\text{BC,BB}} = \frac{b_{\text{abs,BB}}(950 \text{ nm})}{\text{MAC}_{\text{BB}}(950 \text{ nm})}
\]

(4)

where \( \text{MAC}_{\text{TR}} (950 \text{ nm}) \) and \( \text{MAC}_{\text{BB}} (950 \text{ nm}) \) are the mass absorption cross sections for traffic and biomass burning, respectively, and were assumed to be 6.6 m\(^2\) g\(^{-1}\). This value translates to a MAC of around 10 m\(^2\) g\(^{-1}\) at 630 nm, which is a typical value for internally mixed BC at rural and remote sites (Zanatta et al., 2016).

**Aerosol Light Scattering**

Aerosol total light scattering coefficients \( (b_{\text{scat}}) \) and hemispheric backwards scattering coefficients were measured using an integrating nephelometer (Aurora 3000, Ecotech Inc.) at 3 wavelengths (450, 525 and 635 nm) with a native time resolution of 1 min. Data were corrected for the truncation error of the angle non-ideality (Müller et al., 2011). Zero-offset correction was performed at a daily interval, and the calibration span value was regularly defined and verified using CO\(_2\) as reference gas.

\[
b_{\text{abs,TR}}(950 \text{ nm}) = b_{\text{abs}}(470 \text{ nm}) - b_{\text{abs}}(950 \text{ nm}) \left(\frac{470}{950}\right)^{-\alpha_{\text{abs}}}
\]

\[
\left(\frac{470}{950}\right)^{-\alpha_{\text{abs}}} - \left(\frac{470}{950}\right)^{-\alpha_{\text{abs}}}
\]

(2)

where \( \alpha \) is the absorption Ångström exponent. BB and TR denote the sources “biomass burning” and “traffic” to be apportioned.

**Aerosol Optical Properties**

Using the aethalometer and nephelometer data, the following aerosol optical properties can be retrieved:

\[
b_{\text{ext}}(\lambda) = b_{\text{scat}}(\lambda) + b_{\text{abs}}(\lambda)
\]

(5)

\[
SSA_{\text{dry}}(\lambda) = \frac{b_{\text{ext}}(\lambda)}{b_{\text{ext}}(\lambda)}
\]

(6)

\[
SSA_{\text{ambient}}(\lambda, \text{RH}) = \frac{b_{\text{ext}}(\lambda) \cdot f(\text{RH}, \lambda)}{b_{\text{scat}}(\lambda) \cdot f(\text{RH}, \lambda) + b_{\text{abs}}(\lambda)}
\]

(7)

\[
\frac{b_{\text{abs}}(\lambda_1)}{b_{\text{abs}}(\lambda_2)} = \left(\frac{\lambda_1}{\lambda_2}\right)^{-\text{AAE}}
\]

(8)

\[
\frac{b_{\text{scat}}(\lambda_1)}{b_{\text{scat}}(\lambda_2)} = \left(\frac{\lambda_1}{\lambda_2}\right)^{-\text{SAE}}
\]

(9)

\[
\frac{b_{\text{ext}}(\lambda_1)}{b_{\text{ext}}(\lambda_2)} = \left(\frac{\lambda_1}{\lambda_2}\right)^{-\text{EAE}}
\]

(10)

where \( b_{\text{ext}}(\lambda) \) is the aerosol extinction coefficient and \( SSA_{\text{dry}}(\lambda) \) is the dry single scattering albedo (at laboratory conditions with RH \( < 40\% \)). \( SSA_{\text{ambient}}(\lambda, \text{RH}) \) is the single scattering albedo at ambient RH, determined using the scattering enhancement factor \( f(\text{RH}, \lambda) \) obtained by external methods (Zieger et al., 2015). \( f(\text{RH}) \) is the key parameter for describing the RH dependence of the particle light scattering coefficient \( b_{\text{ext}} \) and is defined as the ratio of the ambient \( b_{\text{ext}}(\text{RH}_{\text{ambient}}) \) to its dry value \( b_{\text{ext}}(\text{RH}_{\text{dry}}) \). Knowledge of this RH effect is of crucial importance to climate forcing calculations but is also needed for the validation or comparison of remote sensing with in-situ measurements, as it will be performed in the section “Comparison with AERONET data”.

AAE (also called \( \phi \)) is the absorption Ångström exponent; SAE, the scattering Ångström exponent and EAE, the extinction Ångström exponent. Because \( f(\text{RH}) \) is also wavelength dependent (Zieger et al., 2015; Skupin et al., 2016), EAE, SAE and AAE also exhibit a dependence on RH. The reported values will be labelled as “dry” or “ambient” throughout this manuscript, reflecting the different measurement approaches (ground based in-situ sensing with in-situ measurements, as it will be performed in the section “Comparison with AERONET data”).

**Meteorological Measurements**

The PDI station has the function to monitor the meteorological parameters according to the technical requirements of the Vietnamese National Technical Regulation on Meteorological Observations (MONRE, 2012). Wind speed, wind direction, RH and temperature are recorded four times a day, while precipitation is measured at 00:00, 06:00, 12:00 and 18:00 UTC, corresponding to 07:00, 13:00, 19:00 and 01:00 local time.
Additionally, ambient sensors for temperature and RH recording data with 1 min time resolution are mounted at the main inlet for the aerosol and gas phase measurements.

**Carbon Monoxide Measurements**

Carbon monoxide (CO) was measured with a cavity ring-down spectrometer (CRDS) (Picarro Inc., G2401) which allows the simultaneous detection of CO, carbon dioxide and methane. The CRDS has been shown to be a suitable technique providing high-precision CO observations in ambient air (Zellweger et al., 2012; Yver Kwok et al., 2015). The robustness and little required maintenance of these analyzers also allow long-term observations at remote locations (Fang et al., 2016). All data are corrected for dilution and pressure broadening effects due to humidity and thus represent dry air mole fractions. Samples from four calibration tanks are measured every 3 to 6 days. CO concentrations in the calibration tanks were all directly traced back to the GAW Central Calibration Laboratory for CO at NOAA. Data are reported on the WMO CO scale. Samples from an additional target cylinder were measured every 25 hours for quality control.

**AERONET**

For comparison to the ground based in-situ measurements at Pha Din, AERONET sun photometer data (CE318 multi-band sun photometer; Cimel Electronique) collected at the nearby site Son La (see Fig. 1) are used in this study. The Son La AERONET site (21°19'55"N, 103°54'18"E; 675 m a.s.l.) is located approximately 48 km SE of Pha Din (Wang et al., 2015). The site is located on a hill in Son La City (population: 98,751) and surrounded by grass and tree vegetation. In contrast to Pha Din as a remote site, the Son La site is not only influenced by BB emissions (Nguyen et al., 2016; Popovicheva et al., 2016) but is also directly influenced by the anthropogenic emission sources of the city. The AERONET photometers measure the spectral direct-beam solar radiation as well as directional diffuse radiation along the solar almucantar (Holben et al., 1998). In addition to measuring direct solar irradiance, they also measure the sky radiance at four wavelengths (440, 675, 870 and 1020 nm) along the solar almucantar. The solar zenith angles are restricted to > 50° (i.e., in early morning and late afternoon). The sky radiance measurements are used to retrieve additional columnar aerosol properties, including the volume size distribution, phase function, real and imaginary components of the refractive index, particle effective radius, single-scattering albedo and the asymmetry factor, which are typically computed using AERONET inversion algorithms (Dubovik and King, 2000; Dubovik et al., 2006; Holben et al., 2006). The uncertainty in the retrieved single scattering albedo is estimated to be ± 0.03. In addition, for absorption related inversion products, the Level 2.0 direct sun AOD_{500} must be greater than 0.4. AERONET inversion products are quality assured based on literature (Dubovik et al., 2000; Holben et al., 2006). The AERONET data from Son La were downloaded from the AERONET website (AERONET, 2018), as Version 3, Level 2 (cloud screened and quality-assured). For Son La, the available AERONET data consisted of 400 individual measurements, distributed over 124 days in 2014–2017, mostly in March (167 measurements), April (80 measurements) and May (60 measurements).

**Levoglucosan as Biomass Burning Marker**

Vegetative biomass burning emits gases and particles into the atmosphere and has been described as an important source of organic aerosols (Simoneit and Elías, 2001). Levoglucosan and mannosan, produced by pyrolysis of cellulose and hemicellulose, respectively, are well-established tracers for BB (Simoneit et al., 1999; Simoneit, 2002). During a field campaign with an intensive BB period from 22 March to 12 April 2015, PM_{2.5} (particulate matter with an aerodynamic cut-off diameter ≤ 2.5 µm) was collected on quartz fiber filters (47 mm diameter, 2500QAT-UP; PALL Life Sciences, Inc., USA) by MiniVol™ TAS samplers (Airmetrics, USA) with a 24-h sampling protocol. The quartz fiber filters were extracted ultrasonically with deionized distilled water and filtered through Teflon syringe filters (0.45 µm pore size; Millipore Corp., Billerica, USA). The extract solutions were then analyzed by high performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD) for the quantitative determination of various carbohydrates, including levoglucosan and mannosan. The technical details of the analysis method have been reported previously (Engling et al., 2006; Zhang et al., 2013).

**FLEXTRA Backward Trajectories**

The FLEXTRA backward-trajectory model (Stohl et al., 1995; Stohl and Seibert, 1998) was used to trace the origin of the air masses arriving at Pha Din. Kinematic backward trajectories were calculated on operational 3-hour ECMWF analysis/forecast fields (1° × 1° horizontal resolution and 137 vertical levels). For the whole observation period, trajectories were initialized every 4 hours at different heights above the site and were followed 10 days backward in time. For the analysis presented below, an initial height for the trajectories of 420 m above ground was selected, which roughly represents the middle between the real station altitude and the altitude of the smoothed model topography.

**AVHRR Land Fire Hotspots**

The monthly sum of satellite detected land fire hotspots for Vietnam and the surrounding countries were downloaded from the website of the ASEAN Specialised Meteorological Centre (ASMC, 2018). The hotspots published by ASMC are observed using the NOAA satellite Advanced Very High Resolution Radiometer (AVHRR) sensor. Hotspots may be undetected due to cloudy conditions or partial satellite pass. Hotspot counts from 2016 onwards are based on the NOAA-19 satellite, while for the period of 2006–2015, they are based on the NOAA-18 satellite. Depending on the size and proximity of fires, they may or may not appear as separate hotspots. If the total size of a detected fire is smaller than a pixel, the fire will appear as a single hotspot. As for active fires larger than a pixel, they will appear as multiple hotspots.
RESULTS AND DISCUSSION

Annual Variation of Aerosol Optical Properties and their Relation to Biomass Burning

Fig. 2 shows the time series for the aerosol absorption coefficient ($b_{abs}$), the total aerosol scattering coefficient ($b_{scat}$), the dry absorption Ångström exponent (AAE$_{dry}$), carbon monoxide and the monthly land fire hotspot sum for Vietnam and the surrounding countries, for the entire period of the continuous measurements (February 2014–June 2018). High $b_{abs}$ values clearly correlate with high carbon monoxide (CO) mixing ratio and high number of land fire hotspots for Vietnam and the surrounding countries, with distinct maxima between February and May. BB is known to be a major source of CO (van der Werf et al., 2006; Duncan et al., 2007; Schultz et al., 2008). Due to its atmospheric lifetime on the order of weeks to months (Holloway et al., 2000), CO is a suitable tracer for BB. In regions such as Southeast Asia, the systematic variation in BB intensity throughout the year results in pronounced seasonal cycles of CO emissions (Arellano et al., 2006), which are also reflected in the atmospheric CO levels at Pha Din. The periods with the highest BB influence at Pha Din are consistently characterized by no rainfall, reduced RH and increased temperature, as well as AAE$_{dry}$ values higher than 1.4, peaking up to 1.9–2 (see also Figs. S2–S10 in the

Fig. 2. Time series 2014-2018 of (A) the monthly sum of satellite detected land fire hotspots for Vietnam, Laos, Thailand, Cambodia and Myanmar, (B) the carbon monoxide mixing ratio (daily average values), (C) the aerosol absorption coefficient (daily average values), (D) the absorption Angstrom exponent (daily average values, derived from the aerosol absorption coefficients at 470 nm and 950 nm) and (E) the aerosol total scattering coefficient (daily average values).
Supplementary Information). In contrast, the lower values in the summer months largely coincide with the SE Asian summer monsoon with more rain and generally more humid conditions. During the time period covered so far by the measurements at Pha Din, a decrease in peak intensity for all parameters is observed between 2014 and 2018.

Fig. 3 shows a classification of aerosol types based on AAE and SAE (Cazorla et al., 2013). AAE_{dry} reaches 1.6–2.0 during BB periods, a typical value for BB dominated aerosol in northern Indochina (Wang et al., 2015). SAE_{dry} is > 1.5, pointing to a dominant contribution of accumulation mode particles to light scattering, as is expected for BB aerosols, and does not indicate a significant influence of coarse mode dust. According to the classification used in Fig. 3, the aerosol detected during high BB episodes is thus characterized as “OC dominated”, while for the rest of the year, the major fraction of the data is classified as “EC/OC mixture”.

The aethalometer model (see “Methods”) allows for splitting the eBC mass concentration into its traffic and BB fractions as shown in Fig. 4 (daily and monthly mass concentrations and monthly fractions). The aethalometer model clearly attributes a major part of the eBC mass concentration to BB during the time periods with maximal aerosol absorption coefficients (~80%; see top panel in Fig. 4), while the contributions of eBC(TR) and eBC(BB) are similar during the rest of the year, with higher absolute values during the dry season and lower values during the wet season. Traffic is even a slightly more dominant source than BB during the wet season.

Origin of Air Masses at Pha Din

The origin of the air masses arriving at Pha Din was assessed with the help of FLEXTRA backward trajectories (hereafter called “back-trajectories”). Based on the well-known seasonal meteorology patterns in northern SE Asia (see “Introduction”), the trajectory analysis was performed for the three dominating meteorological regimes, namely (a) the SE monsoon (“wet season”, June–September), (b) the NE monsoon (October–January) and (c) the winter/spring SE Asian tropical high (February–May). For these three regimes, the aerosol optical properties as well as meteorological parameters were apportioned to specific grid cells on a world map, following existing approaches described in the literature (Zieger et al., 2015; Franke et al., 2017). The 3-day FLEXTRA back-trajectories were mapped onto a grid with a horizontal resolution of 0.5° × 0.5°. For each trajectory in an individual grid cell, the aerosol/meteorological parameter measured at Pha Din at the time of the trajectory arrival is assigned to the cell, and the average of all values assigned to a grid cell is formed. This method provides qualitative information on how much of a given quantity is (on average) observed at the receptor station after the air mass has crossed a certain grid cell. The method has clear limitations (Franke et al., 2017) and does not replace an in-depth particle dispersion analysis, but it represents a simple way to qualitatively identify source regions for a parameter measured at the receptor site.

In the main BB season (Fig. 5), the trajectory density plot (Fig. 5(A)) shows two equally dominant branches, one with back-trajectories covering the area SW of Pha Din (Laos, northern Thailand, Cambodia and Myanmar) and a second one covering the NE sector (northern Vietnam and southwestern China). The average trajectory altitude above model ground (Fig. 5(B)) is lower than 300–600 meters for a large part of the area, indicating that the BB influenced

![Fig. 3. Absorption Angstrom exponent (AAE, 470/950 nm) versus scattering Angstrom exponent (SAE, 450/635 nm), color coded by the aerosol absorption coefficient at 880 nm. Dry daily average values are shown (February 2014–June 2018). The classification by aerosol type is based on Cazorla et al. (2013).]
Fig. 4. (A) Equivalent black carbon (eBC) mass concentrations (daily/hourly values) from biomass burning (BB) and traffic (TR) at Pha Din, using the aethalometer based approach by Zotter et al. (2017), (B) Same with monthly average values and (C) Corresponding BB and TR fractions (monthly average values).

air masses arriving at Pha Din had sufficient contact with the BB aerosols emitted on the ground. The source region for eBC(BB) mass (Fig. 5(C)) is clearly located in the SW sector and fully matches the previously identified BB source regions (Laos, northern Thailand, Cambodia and Myanmar; see “Introduction”). The identified source region is furthermore associated with increased temperature (> 20°C) and rather low relative humidity (< 60%; Fig. 5(E)). The source region for eBC(TR) mass (Fig. 5(C)) is more diffuse, with some isolated hotspots in the SE sector along the coastal areas of Thailand and Vietnam. During a field campaign at Pha Din from late March to early April in 2015, molecular BB markers (levoglucosan and mannosan) were measured on additionally collected PM2.5 filters (see “Methods”) during a BB intensive period with air masses advected from the SW. Levoglucosan and mannosan concentrations averaged at 473 ± 635 (2–2285) ng m$^{-3}$ and 58 ± 33 (16–137) ng m$^{-3}$, respectively, indicating a significant impact of BB on the ambient aerosol. Fig. 6 shows that the apportioned eBC(BB) mass concentrations correlate very well with the levoglucosan/mannosan measurements. The distinctly high levoglucosan concentrations (exceeding those reported at many other sites across the globe; Simoneit, 2002) along with the eBC(BB) pattern clearly demonstrate the strong biomass-burning impact on air quality in this region. The fact that the aethalometer model apportions the peak concentrations to BB further confirms the validity of the apportionment approach.

For the warm and humid summer wet season (Fig. S11, Supplementary Information), no spatially relevant source regions are identified, with the exception of small hotspots along the Chinese sea (eBC(BB) mass) and SW Chinese mainland (eBC(TR) mass). During the NE monsoon in the colder and humid winter months (Fig. S12, Supplementary Information), a relatively large source region for both eBC(BB) and eBC(TR) mass is found over mainland SW China. For this period of the year, a previous study (Co et al., 2014) reported that BB plumes from the NE were also reaching Tam Dao, a rural mountain site in northern Vietnam 200 km ESE of Pha Din.

Comparison with AERONET Data

The cloud-free conditions at Pha Din during BB plumes from the SW also coincide with a generally low RH (< 50%). This provides an excellent opportunity to intercompare ground-based in-situ measurements with column-based in-situ measurements under dry conditions. The semi-continuous AERONET measurements performed at Son La, approximately 50 km from Pha Din at 675 m a.s.l. (see Fig. 1), allow for an intercomparison with the continuously measured in-situ aerosol optical properties. Fig. 7(A) shows the in-situ EAE$_{dry}$, in-situ AAE$_{dry}$, AERONET EAE$_{ambient}$ and AERONET AAE$_{ambient}$ for an episode in March 2015 with high BB and available AERONET data.
Fig 5. Source regions for BB and traffic eBC mass concentrations during the winter tropical SE Asian high pressure system, derived from 3-day FLEXTRA back-trajectories. The 3-day FLEXTRA back-trajectories were mapped into a grid with a horizontal resolution of 0.5° × 0.5° (trajectory density shown in panel A). For each trajectory in an individual grid cell the aerosol/meteorological parameters measured at Pha Din at the time of the trajectory arrival is assigned to the cell, and the average of all values assigned to a grid cell is formed.
Despite the difference in location and methodology (ground-based in situ measurements vs. column-integrated measurements), the EAE values (top panel) do not significantly differ, and the time series follow the same general pattern. In contrast, the AAE comparison shows lower values for the AERONET measurements at Son La. To assess this difference in a more quantitative way, all available data between 2014 and 2017 were taken into account (see Fig. S10(a)–10(b)) in the Supplementary Information. On average, the Son La AERONET AAE_{ambient} values are 20% lower compared to the Pha Din in-situ values and show a large lower interquartile range caused by occasionally very low AERONET AAE_{ambient} values. Besides the general methodological differences in the sampled volume (in-situ vs. column), it is also likely that the lower AAE_{ambient} values at Son La are caused by a stronger influence from traffic sources within the PBL above the Son La site (i.e., a lower ratio of brown carbon to EC in the traffic emissions compared to BB emissions). Fig. 7(B) compares the SSA retrieved by the two methods, again for the same period in 2015. Since the ambient relative humidity was low during the peak of the BB event (40% at Pha Din and 50% in Son La), the in-situ values compare well with the AERONET SSA values. Before and after this event, the ambient RH is considerably higher, and the AERONET SSA is correspondingly higher as a result of the scattering enhancement by water uptake (see Eq. (7)).

Scattering enhancement by water uptake, commonly described by a wavelength dependent scattering enhancement factor, $f(RH)$ (Zieger et al., 2015), plays an important role when retrieving scattering coefficients, extinction coefficients and SSA values for the Pha Din in-situ measurements, which are measured under dry conditions according to the GAW recommendations (see “Methods”). Fig. 8 shows the correlation between the in-situ and AERONET SSA for all available data between 2014 and 2017 for two wavelengths, additionally color coded by the ambient relative humidity. The plot first of all shows that with the consistently dry conditions during the BB events (low SSA values), the SSA values are closer to the 1:1 line compared to the other time periods with higher SSA and higher ambient RH. The agreement is slightly better for the red wavelength, while the offset seems higher for the blue wavelength. The good agreement between in-situ and remote sensing measurements during the dry conditions supports the general comparability and good quality of these data. It can therefore be assumed that discrepancies during higher RH are truly caused by light scattering enhancement effects. To obtain an idea of the quantitative range of the expected scattering enhancement for the aerosol at Pha Din, “climatology-based $f(RH, \lambda, \text{clim})$” values for SSA were determined according to Eq. (7) by selecting an $f(RH, \lambda)$ so that the in-situ SSA matches the AERONET SSA. This approach is based on the assumption that the aerosol absorption coefficient is only weakly dependent on RH (Nessler et al., 2005). Fig. 9 shows the determined $f(RH, \lambda, \text{clim})$ plotted against RH. The upper panels of Fig. 9 are color coded with $b_{\text{abs}}$ and the lower panels, with $b_{\text{scat}}$. The color coding shows—in accordance with the trajectory plots in Figs. 5(C) and 5(F)—, that the aerosol concentration and

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Fig. 6. (A) Timeline of biomass burning marker mass concentrations (levoglucosan and mannosan) during an intensive measurement campaign at Pha Din during spring, 2015. Daily values are shown (markers represent start time) and (B) BB and traffic eBC mass concentrations, apportioned by the aethalometer model.
Fig 7. (A) Extinction Angstrom exponent (EAE) and absorption Angstrom exponent (AAE) from ground based in-situ measurements at Pha Din and column-based measurements in Son La (48 km distance from Pha Din) during a high biomass burning episode in March 2015, (B) Single scattering albedo (SSA), calculated for the three nephelometer wavelengths by interpolation of the 7-wavelength aethalometer data, (C) eBC mass concentration, apportioned into BB and traffic contributions by the aethalometer model (using AAE_{BB} = 1.8 in the model) and (D) ambient and laboratory relative humidity.

RH exhibit substantial co-variance at Pha Din: there is a general trend of increasing median $b_{abs}$ and $b_{scat}$ with decreasing RH. Whereas during high $b_{abs}$ events with low relative humidity, $f(RH, \lambda)_{clim}$ is close to unity, $f(RH, \lambda)_{clim}$ increases to ~1.5 (interquartile range: 1.2–2) during the other time periods with high ambient RH and low ambient concentrations of $b_{abs}$ (i.e., the wet season from June to September). Thus, the dry in-situ measured $b_{abs}$ or SSA values are lower than the corresponding values under ambient conditions. The climatology-based $f(RH, \lambda)_{clim}$ values for Pha Din compare well with the few values available in the literature for this region, e.g., an $f(RH, \lambda)$ value of 1.2 at a 450 nm wavelength for 85% RH for near-source BB aerosol in northern Thailand using wet nephelometry (Pantina et al., 2016). Finally, the climatology-based $f(RH, \lambda)_{clim}$ values in Fig. 9 also allow for a simple estimate of the effect of RH on SAE. Based on the $f(RH, \lambda)_{clim}$ values at 450 nm (median values of 1.2 and 1.4 for 30% and 70% RH, respectively) and at 675 nm (median values of 1.1 and 1.6 for 30% and 70% RH, respectively), SAE decreases by approx. 20% at 70% RH compared to the dry value.

CONCLUSIONS

This study presents for the first time results from multi-annual continuous ground-based in-situ aerosol monitoring at a remote site in northern SE Asia. A clear correlation between the time series of the absorption coefficient values and satellite detected land fire hotspots exists, with decreasing peak intensities in the last five years. Also, the ground-based in-situ aerosol data measured at Pha Din agree well with the semi-continuous AERONET data from
Fig. 8. Comparison between the single scattering albedo (SSA) retrieved from AERONET measurements at Son La (48 km distance from Pha Din) and the SSA (dry sampling conditions, see WMO/GAW, 2017) retrieved from the ground-based in-situ measurements at Pha Din. Data points represent all hourly values from 2014–2017 with existing simultaneous in-situ and AERONET data. The comparison was made for two wavelengths (panel A: blue, 440/450 nm, panel B: red, 675/635 nm). Data points are color coded with the ambient relative humidity.

Fig. 9. Climatology-based scattering enhancement factor f(RH) plotted against relative humidity. The comparison was made for two wavelengths (left panel: 450 nm, right panel: 675 nm). Data points are color coded with the absorption coefficient (panels A and B) and the scattering coefficient (panels C and D). The plots are retrieved from all hourly values during 2014–2017 with existing simultaneous in-situ data (Pha Din) and AERONET data (Son La).
the Son La site, which is 50 km from Pha Din. Due to the consistently low ambient relative humidity during the peak BB periods (< 50% RH), RH dependent parameters, such as $b_{\text{scat}}$ or SSA, retrieved from dry ground-based in situ measurements at Pha Din can also be considered to be reasonably representative for ambient conditions during the dry season. Finally, the findings based on the continuous monitoring help to tie together past data obtained during temporally restricted aerosol measurements in the region. Thus, the aerosol monitoring at Pha Din offers an important contribution to a quantitative assessment of the biomass burning impact on regional air quality in northern SE Asia.

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SUPPLEMENTARY MATERIAL

Supplementary data associated with this article can be found in the online version at http://www.aaqr.org.

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