Size-resolved chemical composition, effective density, and optical properties of biomass burning particles

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Abstract. Biomass burning aerosol has important impact on the global radiative budget. A better understanding of the mixing state and chemical composition of biomass burning particles relative to their optical properties is the goal of a number of current studies. In this work, effective density, chemical composition, and optical properties of rice straw burning particles in the size range of 50-400 nm were measured using a suite of comprehensive methods. A Differential Mobility Analyzer (DMA)-Aerosol Particle Mass analyzer (APM)-Condensation Particle Counter (CPC) system offered detailed information on the effective density as well as mixing state of size-resolved particles. The effective density and chemical composition of individual particles were characterized with a DMA in-line with a Single Particle Aerosol Mass Spectrometer (SPAMS), simultaneously. The multiple modes observed in the size-resolved particle effective density distribution indicated size-dependent external mixing of black carbon (BC), organic carbon (OC) and potassium salts in particles. Particles of 50 nm had the smallest effective density (1.16 g/cm³), due to a relative large proportion of aggregate BC. The average effective densities of 100-400 nm particles ranged from 1.35-1.51 g/cm³ with OC and inorganic salts as dominant components. Both density distribution and single-particle mass spectrometry showed more complex mixing states in larger particles. Upon heating, the separation of the effective density distribution modes testified the existence of less volatile BC or soot and potassium salts. Size-resolved optical properties of biomass burning particles were measured by the Cavity Attenuated Phase Shift spectroscopy (CAPS, λ=450 & 530 nm). The single scattering albedo (SSA) showed the lowest value for 50 nm particles (0.741 ± 0.007 & 0.889 ± 0.006) because of larger proportion of BC content. Brown carbon played an important role for the SSA of 100-400 nm particles. The Ångström absorption exponent (AAE) values for all particles were above 1.6, indicating the significant presence of brown carbon. Though freshly emitted, the light absorption enhancement ($E_{abs}$) was observed for particles larger than 200 nm. Concurrent measurements in our work provide a basis for discussing the physicochemical properties of biomass burning aerosol and its effects on global climate and atmospheric environment.

1 Introduction
Biomass burning is a significant source of trace gases and aerosol particles (Andreae and Merlet, 2001). Biomass burning particles affect climate by both absorbing and scattering solar radiation (Chand et al., 2009) and serve as cloud condensation nuclei which would modify cloud microphysical properties (Petters et al., 2009). In addition, biomass burning particles have considerable impacts on air quality, regional visibility, and human health (Naeher et al., 2007; Park et al., 2006). Global annual emissions of black carbon (BC) and organic carbon (OC) aerosols are estimated to be ~8 and 33.9 Tg yr\(^{-1}\) while open burning contributes approximately 42% for BC and 74% for OC (Bond et al., 2004). Along with rapid economic development and increase in agricultural activities, emissions from agricultural residue combustion in China have drawn extensive attention. The total amount of straws from open burning in China is estimated to be ~140 Tg yr\(^{-1}\) (Cao et al., 2008). Controlling biomass burning emissions is thus necessary to improve air quality in China.

Mixing state, composition, and morphology of particles can influence their radiative properties. An enhancement of BC forcing by up to a factor of 2.9 is estimated by models when BC is internally mixed with other components compared with externally mixed scenarios (Jacobson 2001). Quantitative assessment of light absorption properties of biomass burning particles has drawn widely attention as the radiative impacts of these particles are both affected by the strongly absorbing BC and organic matter (Chakrabarty et al., 2010). The co-emission of BC and OC can lead to internally mixed particles, in which the OC coating can enhance particle absorption through lensing effects (Bond et al., 2006; Schnaiter et al., 2005). For internally mixed BC, the assumption of a void-free BC sphere with a material density of 1.8 g/cm\(^{3}\) can lead to overestimations of the shell/core ratio and absorption enhancement by ~13 and ~17%, respectively (Zhang et al., 2016). In addition to absorption enhancement by internal mixing, some organic matter containing specific functional groups (e.g. nitrated/polycyclic aromatics, phenols) can itself absorb radiation in the low-visible and UV wavelengths (Hoffer et al., 2006; Jacobson, 1999) and is referred to as brown carbon (BrC). As biomass burning is a significant source of BrC, the optical properties of biomass burning particles need to be further understood. Field works have been conducted to measure the light absorption enhancement by particle coatings in different areas (Chan et al., 2011; Nakayama et al., 2014). The degree to which particles absorb light depends on their composition, shape, and mixing state. Researches on chemical composition and mixing state of biomass burning particles have been done by our group members previously (Huo et al., 2016; Zhai et al., 2015). However, it remains unclear how mixing states and chemical composition of biomass burning particles influence their morphology and optical properties.

Particles emitted from biomass burning are generally composed of a mixture of spherical and non-spherical particles and chain aggregates (Martins et al., 1998). Scanning electron microscopy (SEM) as well as transmission electron microscopy (TEM) are common techniques widely used to investigate the morphology of biomass burning particles (China et al., 2013; Giordano et al., 2015). However, these methods are unable to provide continuous “on-line” information and suffer from limitations arising from primary particle overlap, screening effects, and cluster anisotropy.
(Wentzel et al., 2003). Effective density is a good predictor for the complex properties of biomass burning particles (Pitz et al., 2008) and is often used to convert particle size distributions into mass loading (Tang and Munkelwitz, 1994). Variations in particle effective density can be used to follow compositional transformations during chemical reactions (Katrib et al., 2005). Online measurements which provide real-time monitoring of particle effective density variation have been developed. McMurry and McMurry (1992) developed a density measurement technique based on the selection of a monodisperse aerosol with a Differential Mobility Analyzer (DMA) followed by classification according to aerodynamic diameter with an impactor. McMurry et al. (2002) reported a technique to determine size-resolved effective density based on using an Aerosol Particle Mass analyzer (APM) to measure the mass of particles that had been classified according to electrical mobility by a DMA. The DMA-APM method has been applied extensively in field studies as well as laboratory experiments (Hu et al., 2012; Barone et al., 2011). However, few measurements of the effective density of biomass burning particles have been done due to the lack of accompanying on-line chemical information.

Mixing state of individual particle can be very different caused by the chemical composition, aging degree, etc., which greatly influence the morphology and optical property of particles. Thus, distinction among particles might be covered by bulk measurements. Single particle mass spectrometry techniques have been utilized to measure the chemical composition, size, density, and shape of individual particles. Spencer et al. (2007) utilized a DMA-ultrafine aerosol time-of-flight mass spectrometer (UF-ATOFMS) system to detect the effective density and chemical composition simultaneously of ambient aerosol at single-particle level. The comprehensive information about single particles could help to elucidate the morphology, mixing state, and sphericity of biomass burning particles.

The chemical composition, morphology, and optical properties of particles are usually interrelated. However, given that biomass burning particles is a complex mixture of organic and inorganic species, including strongly light-absorbing BC and BrC, size-resolved or even single particle level information on the morphology, chemical composition, and optical properties of biomass burning particles are needed to help get a macroscopic understanding of the relationships. In this study, laboratory experiments were conducted on aerosols from combusting rice straws, a main source of biomass burning particles in Southern China. The size-resolved effective density of biomass burning particles was measured by two different methods. One was based on a DMA-APM-Condensation Particle Counter (CPC) system. For the other method, the mobility size-selected particles by a DMA were transported into a Single Particle Aerosol Mass Spectrometer (SPAMS), where the vacuum aerodynamic diameter and chemical composition of individual particles were measured. Size-resolved optical properties of biomass burning particles were also measured by Cavity Attenuated Phase Shift spectroscopy (CAPS). A thermodenuder (TD) was used to help analyze the mixing state of particles by removing the volatile compounds and leaving behind the less volatile species based on the vaporization temperature of materials. The purpose of our study was to add physicochemical knowledge regarding biomass
burning particles which is an important aerosol source globally.

2 Experiments

2.1 Laboratory-made biomass burning particles

Rice straw, a typical type of crop residue in Southern China, was taken as the representative biomass burning material in our experiment. The self-designed combustion setup was introduced in previous work (Huo et al., 2016). Briefly, the rice straw collected in rural residential area in Shanghai were dehydrated for 24 h at 100°C in an oven prior to combustion. Five replicate tests of straw-burning were conducted for each experiment. For each test, ~50g of dried rice straws were burned in a combustion stove at a flaming condition. The emitted smoke was introduced into a 4.5 m³ (in volume) chamber with a flow rate of 50 L/min. Ambient air was introduced though a high efficiency particulate air filters to maintain the ambient pressure. The particles in chamber were then introduced into the measurement system through a silica gel type diffusion drier (shown in Figure 1).

2.2 Single particle mass spectrometry

A Single Particle Aerosol Mass Spectrometer (SPAMS) (Hexin Analytical Instrument Co., Ltd) was deployed to examine the aerosol chemical composition and aerodynamic diameter at single-particle level. Detailed information on the SPAMS has been described elsewhere (Li et al., 2011). Briefly, particles in the size range of 0.2-2.0 μm are first drawn into the vacuum through an Aerodynamic Focusing Lens. Each particle is accelerated to a size-dependent aerodynamic velocity which is calculated based on two orthogonally oriented continuous lasers (Nd: YAG, 532 nm). The two lasers fixed at a 6 cm distance and the delay of the scatter light is collected by two photomultiplier tubes (PMT). When a particle arrives at the ion source region, a pulsed desorption/ionization laser (Q-switched Nd: YAG, 266 nm) is triggered. Ions are recorded by a bipolar time-of-flight spectrometer, which records both positive and negative mass spectra for each single particle. In this work, the power of desorption/ionization laser was set to ~0.6 mJ per pulse. The aerodynamic diameter measurement is calibrated with curves generated by monodisperse polystyrene latex spheres (Nanosphere Size Standards, Duke Scientific Corp.) with known diameters (0.2-2.0 μm).

All single particle mass spectra acquired were converted to a list of peaks at each m/z by setting a minimum signal threshold of 30 arbitrary units above the baseline with TSI MS-Analyze software. The resulting peak lists together with other SPAMS data were imported into YAADA (version 2.11, www.yaada.org), a software toolkit for single-particle data analysis written in Matlab (version R2011b). In this work, a total of 10220 biomass burning particles were chemically analyzed according to their positive and negative ion spectra, accounting for about 48 % of all sized particles. According to the similarities of the mass-to-charge ratio and peak intensity, the biomass burning particles were classified using an adaptive resonance theory-based clustering method (ART-2a) (Song et al., 1999). Parameters for ART-2a used in this work such as vigilance factor, learning rate, and iterations were 0.85, 0.05, and 20, respectively. The particle clusters resulting from ART-2a were then grouped into 6
particle types based on the mass spectral patterns and chemical similarities. The name of a particle type reflects the dominant chemical species.

2.3 Effective density measurements

2.3.1 Theoretical calculation and methods

Particle density ($\rho_p$) is referenced to the volume equivalent diameter ($d_{ve}$) which is defined as the diameter of a spherical particle with the same volume as the particle under consideration. Particle density can be derived as follows, where $m_p$ is the particle mass:

$$\rho_p = \frac{m_p}{\pi d_{ve}^3}$$  \hspace{1cm} (1)

When particles are not spherical, the “effective density”, not necessarily a true measurement of particle density is derived. Various definitions of effective density are provided in the literature, and a review of these definitions is given by DeCarlo et al. (2004). Different definitions may aim to present different values for a given particle. It is important to understand the derivation, calculation, and measurement for one method of particle effective density.

(1) DMA-APM-CPC system

The effective density of a particle can be calculated by combining mobility and mass measurements under the assumption that the particle is spherical, thus its physical diameter equals to the electrical mobility diameter ($d_{m}$) measured by a DMA. The effective density ($\rho_{eff}$) can be calculated by the following equation:

$$\rho_{eff}^1 = \frac{m_p}{\pi d_{m}^3}$$  \hspace{1cm} (2)

where $m_p$ stands for particle mass obtained by an APM. In our work, we selected biomass burning particles with mobility diameters of 50 nm, 100 nm, 200 nm, and 400 nm and determined their effective density using the DMA-APM-CPC system.

(2) DMA-SPAMS system

Another approach of deriving effective density is through a combination of mobility and aerodynamic measurements. Simultaneously measuring the particle electrical mobility diameter ($d_{m}$) by DMA and the vacuum aerodynamic diameter ($d_{va}$) by SPAMS allows for the determination of particle effective density ($\rho_{eff}^2$) by the following equation:

$$\rho_{eff}^2 = \frac{d_{va}}{d_{m}} \rho_0$$  \hspace{1cm} (3)

where $\rho_0$ is the standard density (1.0 g/cm³). In this study, since particles smaller than 200 nm may not scatter sufficient light to be detected by SPAMS and the number concentration of biomass burning particles above 400 nm was low (shown in Figure S1), we selected 200 nm and 400 nm particles by DMA and then introduced them into SPAMS.

(3) Shape factor calculation

The shape of particles can influence the optical properties and can reflect the mixing state of particles to some degree. It is possible to extract the shape information
based on the measurements above.

The relationship between the volume equivalent diameter ($d_{ve}$) and mobility diameter ($d_m$) is shown in the following equation:

$$\frac{d_m}{C_c(d_m)} = \frac{d_{ve}X}{C_c(d_{ve})}$$  \hspace{1cm} (4)

where $\chi$ is the shape factor, the ratio of the resistance force on the nonspherical particle to the resistance force on its volume equivalent sphere (Hinds, 1999). The $\chi$ value equals 1 for spherical particles and is greater than 1 for nonspherical/irregular particles.

$C_c$ is the Cunningham Slip Correction Factor parameterized as:

$$C_c(d) = 1 + \frac{2\lambda}{d} [\alpha + \beta \exp (-\gamma \frac{d}{\lambda})]$$  \hspace{1cm} (5)

where $d$ is the particle diameter ($d_m$ or $d_{ve}$) and $\lambda$ is the mean free path of gas molecules. The empirical constants $\alpha$, $\beta$, and $\gamma$ are 1.142, 0.558, and 0.999, respectively (Allen and Raabe, 1985).

The vacuum aerodynamic diameter ($d_{va}$) is related to the volume equivalent diameter ($d_{ve}$) by:

$$d_{va} = \frac{\rho_p d_{ve}}{\rho_o \chi}$$  \hspace{1cm} (6)

As the measurements of mobility and aerodynamic diameters are readily available, we assumed the error was in the particle mass measurement if the measured $\rho^\prime_{eff}$ was used to replace $\rho^\prime_{eff}$ in Equation (2) (Decarlo et al., 2004). With assumed particle density ($\rho_p$) and known particle mass ($m_p$) measured by an APM, a calculated $d_{ve}$ could be obtained using Equation (1). Here we assumed $\rho_p$ equals to 1 which would be canceled out later. Using the same $d_{ve}$ and for any shape factor ($\chi$), a calculated $d_m$ and $d_{va}$ was obtained by Equation (4) and (6), respectively. Thus, $\rho^\prime_{eff}$ could be obtained by the calculated $d_m$ and $d_{va}$ and an estimated $m_p$ was calculated by replacing $\rho^\prime_{eff}$ by $\rho^\prime_{eff}$ in Equation (2). We then calculated the ratio of the estimated $m_p$ to the exact $m_p$ as a function of $d_m$ and $\chi$ (shown in Figure S5, discussed in Section 3.1.5).

**2.3.2 Instruments**

The size distribution of biomass burning particles was detected by a Scanning Mobility Particle Sizer (SMPS) consisting of a Differential Mobility Analyzer (DMA, Model 3080, TSI Inc.) and a Condensation Particle Counter (CPC, Model 3775, TSI Inc.). An Aerosol Particle Mass analyzer (APM, Model 3601, Kanomax Inc.) was used to classify aerosol particles according to their mass-to-charge ratio. The detailed information of the APM classification principle was previously reviewed by Tajima et al. (2011). Briefly, particles were size-selected by DMA after being charged with a Kr85 neutralizer. Particles with a known size were then introduced into APM. When the radial electrical and centrifugal forces were in balance, particles passed through the rotating cylinders to CPC. Mass distribution was obtained by voltage scanning and
particle counting.

2.4 Optical measurements
Cavity Attenuated Phase Shift (CAPS) spectroscopy (Shoreline Science Research Inc.) was used to determine the particle extinction and scattering coefficient. Detailed information on the CAPS is available in Onasch et al. (2015). Briefly, a square-wave modulated light-emitting diode (LED) is transmitted through an optical cavity cell. A sample cell incorporating two high reflectivity mirrors (R~0.9999) with a vacuum photodiode detector (Hamamatsu R645) centers at the wavelength of the LED. The particle extinction coefficient \(b_{\text{ext}}(\lambda)\) can be obtained from the changes in the phase shift of the distorted waveform of the LED. An integrating nephelometer using a 10 cm diameter integrating sphere is operated to measure the scattering coefficient \(b_{\text{scat}}(\lambda)\). Particles are illuminated by the collimated light beam which has measured the extinction. The scattered light of particles is collected at all angels by the integrating sphere. A PMT (H7828-01, Hamamatsu) with a high voltage power supply and an amplifier records the scattered light. In this work, we used two CAPSs with the LED light sources at wavelength of 450 nm and 530 nm to detect the optical properties of biomass burning particles, respectively.

2.5 Thermodenuder
A thermodenuder (TD, Model 3065, TSI Inc.) was utilized to separate volatile and less volatile species of biomass burning particles at specific temperatures. The TD consists of a 40 cm long desorber section and a 70 cm long adsorption tube. The sample can be heated up to 400 °C in the desorber section while we selected 150 °C and 300 °C in this work. The adsorption tube is surrounded by an annular bed of activated carbon which adsorbs the evaporated gas-phase compounds, leaving behind the less volatile fractions. With a flowrate of 0.6 L/min, the residence time of particles in the TD heating section was approximately 9 s in this work.

The particle number fractions after heating do not necessarily represent the actual number fractions before heating as some of the particles can evaporate completely. Besides, particle loss could be produced both in the TD heating and adsorption section due to thermophoretic forces and diffusion, respectively (Philippin et al., 2004). On account of the quantitative measurements of optical properties, particle loss could lead to the underestimate of \(b_{\text{ext}}\) and \(b_{\text{scat}}\).

Sodium chloride (NaCl) aerosol produced by a single-jet atomizer (Model 9302, TSI Inc.) was used to determine the transport efficiency (\(\eta\)) in TD. The transport efficiencies of NaCl of different electric mobility diameters selected by DMA (\(d_{\text{m}}\): 50, 100, 200, and 400 nm) at a range of temperatures (\(T_i\): 20, 150, and 300 °C) are shown in Figure S2. In TD, \(\eta\) decreased with increasing \(T_i\) and decreasing \(d_{\text{m}}\), which was consistent with the result in Philippin et al. (2004). The measured \(\eta\) were used to correct the particle number concentration in the calculation of optical properties.

3 Result and discussion
3.1 Size-resolved effective density
3.1.1 Effective density from DMA-APM-CPC measurements (\(\rho_{\text{eff}}^I\))
The effective density of particles, measured using the DMA-APM-CPC system \( \rho_{\text{eff}} \), provided useful information on the mixing state of particles. A Gaussian model was applied to determine the effective densities of the biomass burning particles selected by DMA (shown in Figure 2). The density distribution of 50 nm \( (d_{50}) \) particles showed a single peak profile with a peak value of 1.17 g/cm\(^3\) (Table S1). Two possible factors could be inferred from this feature: a nearly-monodisperse aerosol effective density distribution or a juncture of two models with very close peak values. Biomass burning particles contain highly agglomerated structures like soot (Martins et al., 1998). Although the material density of black carbon (BC) is \( \approx 1.8 \) g/cm\(^3\) (Malm et al., 2005), fresh BC particles with an aggregate structure can have an effective density less than 1.0 g/cm\(^3\). The density of organic matter varies in the range of 1.2-2.0 g/cm\(^3\) depending on sources (Hand et al., 2010; Turpin and Lim, 2001). Since particles of 50 nm have the possibility of containing organic matter rather than BC alone, the apparent single-peak density distribution of these particles was more likely due to the combination of two models representing BC and organic particles respectively (as the dash lines shown in Figure 2). The thermal desorption method can help to explain the mixing state of 50 nm particles which will be discussed in Section 3.1.3.

The density distribution of 100 nm particles exhibited a peak at 1.45 g/cm\(^3\), which suggests that these particles were dominated by organic matter. However, less-massive composition with effective density of 0.9-1.1 g/cm\(^3\) was also obtained for 100 nm particles. This range is identical with the density of fresh BC with aggregate structure. The bimodal distribution of the density profile of 100 nm particles suggests that BC was partly externally mixed with other components in ultrafine particles from biomass burning emissions. Similar result has been found by Lack et al. (2012) and Adachi et al. (2011). The external mixing of BC and organic particulate matter was evident in the density distribution of 200 nm particles as well (Figure 2). For 400 nm particles, besides a dominant density mode at 1.34 g/cm\(^3\), a relative weak mode with effective density of 1.92 g/cm\(^3\) was observed. Previous studies have shown that potassium chloride crystals, which have a material density of \( \approx 1.99 \) g/cm\(^3\) (Lide, 2008), were observed in the TEM of fresh biomass burning particles (Li et al., 2015). Thus, we estimate that the mode at 1.92 g/cm\(^3\) was associated with KCl, and possibly KSO\(_4\) and KNO\(_3\), and that these crystalline species were more likely externally mixed with organic matter in biomass burning particles.

Though freshly emitted, biomass burning particles can be coated by secondary species, such as ammonium nitrate and ammonium sulfate, pronouncedly in a very short period (Leskimen et al., 2007). The bulk densities of ammonium nitrate and ammonium sulfate are \( \approx 1.75 \) g/cm\(^3\). The differences in the peak values of the dominant mode observed for 50-400 nm particles are associated with the composition and morphology of particles. Different proportions of the same material can lead to differences in particle effective density. The dominant modes for biomass burning particles in the size range of 50-400 nm (Figure 2) could be a mixture of similar composition (organic matter and secondary species) but different proportions. Detailed information and discussion about the particle composition can be found in...
Section 3.2.

3.1.2 Effective density from DMA-SPAMS measurements ($\rho_{\text{eff}}^\text{II}$)

The vacuum aerodynamic size distributions of 200 nm and 400 nm electrical mobility selected biomass burning particles are shown in Figure 3. The dominant mode for the 200 nm mobility selected particles was 280 nm in vacuum aerodynamic diameter with an effective density ($\rho_{\text{eff}}^\text{II}$) of 1.40 g/cm$^3$ and a second mode at 360 nm ($d_{\text{a}},$) with an effective density of 1.80 g/cm$^3$. This is quite consistent with the result from the DMA-APM-CPC method. The less intense mode at 520 nm ($d_{\text{a}},$) should be due to doubly charged particles (Spencer et al., 2007). For 400 nm mobility selected particles, the dominant mode in aerodynamic diameter was 540 nm with an effective density of 1.35 g/cm$^3$. Since the less massive modes at 660 nm and 840 nm were not in the deviation range of doubly charged particles, these two modes were singly charged particles with effective density of 1.65 and 2.10 g/cm$^3$, respectively. The single-particle level chemical composition of biomass burning particles will be discussed below.

3.1.3 Thermal denuded particle effective density

The average density distributions of 50-400 nm ($d_{\text{a}},$) biomass burning particles after heating at 150 °C and 300 °C, respectively, are shown in Figure 2. After heating by TD, the bi-model density distributions of biomass burning particles became more pronounced. At 150 °C, the effective density mode with peak at ~1.0 g/cm$^3$ protruded for the whole size range of 50-400 nm particles. The separation of the peaks helped to testify that the less volatile BC or soot with effective density of ~1.0 g/cm$^3$ was partly externally mixed with other compositions. The dominant density peak values for 50, 100, 200, and 400 nm particles at 150 °C were 1.64-1.80 g/cm$^3$. Li et al. (2016) reported that the density of organic matter vaporized at 150 °C was 0.61-0.90 g/cm$^3$. The increase of the dominant density peak value (1.34-1.45 g/cm$^3$ for unheated vs. 1.64-1.80 g/cm$^3$ for 150 °C heated) could be due to the volatilization of organics with small effective density. The dominant density peak values of 50-400 nm particles at 300 °C were 1.75-2.04 g/cm$^3$. The volatilization temperatures of ammonium nitrate and ammonium sulfate were reported to be ~48-89 °C and ~178-205 °C, respectively (Johnson et al., 2004a; Johnson et al., 2004b). Thus, the fractions of ammonium nitrate and ammonium sulfate should be small at 300 °C. The increase of dominant density peak value for 50-400 nm biomass burning particles upon heating could be due to the vaporization of volatile organics with small effective density and secondary inorganic species such as NH$_4$NO$_3$ and (NH$_4$)$_2$SO$_4$ with density of ~1.75 g/cm$^3$. Upon heating, the density mode of KCl and partly K$_2$SO$_4$ at ~2.0 g/cm$^3$ was ambiguous as the dominant mode shifted right and overlapped the KCl mode (dash lines shown in Figure 2). However, at 300 °C, the dominant mode of 400 nm particles was at 2.05 g/cm$^3$ which fitted the density of potassium salts, indicating the main material of 400 nm heated (~800 nm unheated, detected by a tandem DMAs, discussed in Section 3.3.3) biomass burning particles should be potassium salts.

With heating by TD, the aerodynamic size distributions of 200 nm and 400 nm
electrical mobility size-selected biomass burning particles at 300 °C are shown in Figure S3. The increase of $\rho_{\text{eff}}^I$ upon heating was consistent with that of $\rho_{\text{eff}}^I$.

3.1.4 Comparison of $\rho_{\text{eff}}^I$ and $\rho_{\text{eff}}^I$

Figure S4 summarizes that the average effective densities ($\rho_{\text{eff}}^I$ & $\rho_{\text{eff}}^I$) of biomass burning particles that were size-selected at 6 different mobility diameters. Note that the density distributions of the 300 nm and 350 nm ($d_m$) particles are not contained in Figure 2 since they were similar to those of the 200 nm and 400 nm ($d_m$) particles. The 50 nm biomass burning particles had the lowest effective density of 1.15 ± 0.23 g/cm³ which could be due to the aggregate structure of black carbon. Compared with the 50 nm ($d_m$) particles, the effective density of 100 nm particles was larger (1.45 ± 0.15 g/cm³). Since the sampling limitation of SPAMS was 200 nm, $\rho_{\text{eff}}^I$ was derived only for particles in the size range of 200-400 nm ($d_m$). Overall, these two methods had consistent results although $\rho_{\text{eff}}^I$ were generally smaller than $\rho_{\text{eff}}^I$.

3.1.5 Shape factor

The shape of particles has been suggested to play an important role in their optical properties (Zhang et al., 2008) and mixing state (China et al., 2013). Shape factor was introduced to account for the ratio of the drag forces on a particle due to nonspherical/irregular shape. Shape factor, which can be extracted based on the measurement of particle density and mass has been introduced in Section 2.3.1.

We calculated the ratio of the estimated $m_p$ to the exact $m_p$ as a function of $d_m$ (shown in Figure S5). For nonspherical particles ($\chi > 1$), the estimated mass was larger than the actual mass. We calculated the estimated mass using the exact $\rho_{\text{eff}}^I$ measured by the DMA-SPAMS to replace the $\rho_{\text{eff}}^I$ in Equation (2) as well. The ratios of the estimated mass by this mean to the exact mass for 200, 300, 350, and 400 nm mobility selected particles were 1.4, 1.3, 1.3, and 1.2 respectively (red dots in Figure S5). Thus, we could estimate the $\chi$ of the particle measured using the DMA-SPAMS in the size range of 200-400 nm. Totally, the $\chi$ of 200-400 nm biomass burning particle in this work exceeded 1.2 (~1.2-2.2). The $\chi$ decreased with the increase of $d_m$ while the effective density showed the same trend. The more regular shape and lower effective density of 400 nm particles compared with that of 200 nm particles could be due to the particle chemical composition and particle voids (discussed in Section 3.2).

3.2 Size-resolved chemical composition

Based on the mass spectra of single particles, the biomass burning particles were classified into 6 particle types: 1) BB-CN: biomass burning (BB) particles with a strong CN (m/z -26 [CN⁺]) peak; 2) BB-EC: BB particles with strong elemental carbon clusters ($C_n^{+/−}$); 3) BB-Nitrate: BB particles with strong nitrate (m/z -46[NO₂⁻]).
-62[NO$_3$] signals; 4) BB-Sulfate: BB particles with strong sulfate (m/z -97[HSO$_4$] signals; 5) BB-KCl: BB particles with strong potassium chloride (m/z 113[K$_2$Cl]) signals; and 6) BB-OC: BB particles with strong organic carbon peaks (e.g., m/z 27[C$_2$H$_5^+$], 37[C$_2$H$_7^+$], 43[C$_3$H$_7^+$], 51[C$_4$H$_9^+$], et al.). The naming of the chemical classes is based on some of the dominant chemical species in an attempt to keep the names short. The mass spectra for each particle type are presented in Figure S6. The percentages of 6 particle types in different modes of aerodynamic size distribution for 200 nm and 400 nm mobility selected particles are shown in Figure 3. For 200 nm mobility selected particles, the dominant particle types were BB-EC and BB-CN. The percentages of particle types within the two aerodynamic modes differ slightly. Compared with the first mode, the second mode contains more BB-CN (24.4% vs. 29.6%), more BB-KCl (1.0% vs. 4.3%) and less BB-EC (32.2% vs. 22.9%). We supposed that the density of each particle type largely depended on the dominant species. The exact effective density of each particle type could not be obtained directly while the relative value compared with other particle types could be inferred from the material density of dominant species. For example, the BB-KCl type may have larger effective density compared with others since the dominant composition of BB-KCl type has material density of ~1.99 g/cm$^3$ (Lide, 2008). The increased BB-KCl type and the decrease of BB-EC (~1.0 g/cm$^3$) resulted in a larger effective density in the second mode than the first mode.

The fractional distributions of the 6 particle types for 200 nm and 400 nm mobility selected particles were apparently different (Figure 3). For 400 nm mobility selected particles, the proportions of BB-Nitrate, BB-Sulfate and BB-KCl types were larger than those of 200 nm mobility selected particles. The dominant chemical species for BB-Nitrate and BB-Sulfate particle types could be NH$_2$NO$_3$ and (NH$_4$)$_2$SO$_4$ with material density of ~1.75 g/cm$^3$ (Lide, 2008). Compared with other types, BB-Nitrate, BB-Sulfate and BB-KCl were particle types with larger density. However, the effective density for 400 nm mobility selected particles was smaller than that of 200 nm. In addition to the compositional differences, particle morphology could be another reason responsible for the observed differences in the effective densities between these two sizes. Indeed, it has been found that the morphology like void ratio, particle shape factor, and fractal dimension of particles all greatly affect particle effective density (DeCarlo et al., 2004). Though the shape factor discussed in Section 3.1 had shown that the 400 nm ($d_m$) particles had a more spherical morphology, their lower average effective density compared to smaller particles could be due to the voids in particles. Amorphous species such as NH$_2$NO$_3$ could lead to the low effective density of particles. Thus, we supposed the lower effective density of 400 nm particles compared with 200 nm particles was caused by the large proportion of NH$_2$NO$_3$ and (NH$_4$)$_2$SO$_4$ with fluffy material properties.

For 400 nm mobility selected particles, there was no big difference of particle types between the dominant and the secondary modes. The difference between effective densities of these two modes could be due to the different proportions of particle types. However, the proportion of BB-KCl in the third mode at 840 nm with effective density of 2.10 g/cm$^3$ greatly increased compared with the first two modes.
(8.8%, 9.2% vs. 32.7%). The increased BB-KCl indicated that the KCl crystals were external mixed and tended to be mixed with larger size particles which were consistent with the DMA-APM-CPC result.

Upon heating by TD, the proportions of BB-CN and BB-KCl increased, indicating that these types of particles were composed of less volatile species (shown in Figure S3) (Zhai et al., 2015). At 300°C, the fractions of BB-Nitrate and BB-Sulfate decreased, consistent with the volatilization temperature ranges of ammonium nitrate (~48-89°C) and ammonium sulfate (~178-205°C) (Johnson et al., 2004a; Johnson et al., 2004b). The high effective density (>2.0) of biomass burning particles at 300°C could be due to the vaporization of volatile organics with small density and the secondary species (NH₄NO₃ and (NH₄)₂SO₄ with density of ~1.75 g/cm³).

3.3 Size-resolved optical properties

3.3.1 Single scattering albedo (SSA)

The single scattering albedo (SSA), was calculated using the following equation:

$$SSA(\lambda) = \frac{b_{\text{scat}}(\lambda)}{b_{\text{abs}}(\lambda) + b_{\text{scat}}(\lambda)}$$

where $b_{\text{scat}}$ is the particle light scattering coefficient, $b_{\text{abs}}$ is the light absorption coefficient, and $\lambda$ is wavelength. The light scattering and extinction coefficients ($b_{\text{ext}} = b_{\text{abs}} + b_{\text{scat}}$) for biomass burning particles in this work were measured at 530 nm and 450 nm wavelengths using CAPSs.

The size-resolved SSAs for biomass burning particles are shown in Figure 4. Totally, the SSAs for biomass burning particles in the mobility size range of 50-400 nm varied narrowly. However, it’s worth noting that our measurement of optical properties was based on bulk measurement by CAPSs rather than single particle like the chemical information obtained by SPAMS. Thus, the mixing states of size-resolved biomass burning particles which can strongly affect the balance of radiative forcing by BC and organic matter might be offset in our measurement.

The SSA (530 nm) for 50 nm particles was the lowest (0.889±0.006) as the percentage of strong light-absorbing black carbon for particles in this size range was larger (shown in Figure 3, discussed in Section 3.2). For 100-400 nm biomass burning particles, the SSAs were relatively steady (0.897±0.006 - 0.900±0.006).

The size-resolved SSAs at 450 nm ($\lambda$) for biomass burning particles were generally lower than those at 530 nm ($\lambda$). Previous studies have shown that biomass burning was an important source of brown carbon (BrC) which is light-absorbing in the UV-vis range (Lack and Cappa, 2010). For 50 nm ($d_m$) particles, the SSA ($\lambda$=450 nm) was also the lowest, due to the dominance of the strong light-absorbing BC in these particles. However, unlike the trend of size-resolved SSAs ($\lambda$=530 nm), the SSA ($\lambda$=450 nm) of 100-400 nm particles decreased as the size increased. It has been shown that brown carbon arising from biomass burning is primarily composed of extremely low volatility organic compounds (Saleh et al., 2014). The CN in biomass burning particles is representative for some extremely low-volatile nitrogen-containing organics (Zhai et al., 2015). As shown in Figure 3, compared with 400 nm particles, the proportion of organic matter (BB-CN, BB-OC) was larger for 200 nm particles. The nitrogen-containing species might indicate the existence of...
light-absorbing organics. The lower SSA ($\lambda=450$ nm) for 200 nm particles might indicate higher proportion of BrC. We assumed the lower SSA ($\lambda=450$ nm) for 100 nm performed in a similar way with larger proportion of BrC.

3.3.2 Ångström absorption exponent (AAE)

To investigate the wavelength dependence of the absorption coefficients, we determined the Ångström absorption exponent (AAE) based on absorption measurements at two different wavelengths ($\lambda_1$ & $\lambda_2$) using the following equation:

$$\text{AAE} (\lambda_1 / \lambda_2) = -\frac{\ln b_{abs} (\lambda_1)}{\ln b_{abs} (\lambda_2)}$$

The AAE in this work was calculated from the light absorption coefficients at wavelengths of 450 nm and 530 nm measured by the CAPSs. The uncertainties in the calculated AAE values can be caused by the uncertainties in the calibration factors of CAPSs. The size-resolved AAEs for biomass burning particles are shown in Figure 4. Black carbon is highly absorbing in the visible spectrum with little variation with wavelength and shows an AAE of $\sim$1.0 (Bergstrom et al., 2002). As brown carbon species absorb light in the UV-vis range, BrC-containing particles usually exhibit an AAE above 1 (Martinsson et al., 2015). Lack and Cappa (2010) used modeling to calculate AAE values and suggested that particles with AAE exceeding 1.6 should be classified as BrC. In our study, the AAE values of particles in the size range of 50-400 nm were higher than 1.6, indicating that they were BrC containing particles from biomass burning. Among all sizes, the AAE of 50 nm biomass burning particles was the lowest ($\sim$5.8) while that of 100 nm particles was the highest ($\sim$6.3). The main light-absorbing functional groups in the UV-vis range is conjugated double bond (Laskin et al., 2015). BB-CN and BB-OC particle types identified by mass spectra in our work tended to contain more large molecules of BrC with light-absorbing functional groups. We noticed that the proportion of BB-OC type species was larger in 200 nm particles (Figure 3) and with higher AAE value, compared with 400 nm particles. Thus, we suppose the highest AAE value observed for 100 nm particles might be the result of the highest BrC proportion.

The SSA and AAE values of total biomass burning particles are shown in Table S2. The decrease of SSA values upon heating was due to the vaporization of secondary inorganic species like NH$_4$NO$_3$ and less absorbing organics. The AAE values for all particles at 150 °C and 300 °C were $\sim$19% and $\sim$64%, respectively, lower than those at room temperature (20 °C). The large decrease of AAE at 300 °C could be due to the vaporization of light-absorbing organics in the temperature range of 150-300 °C. However, the AAE value at 300 °C was still above the standard of BrC (1.6), indicating the presence of extremely low-volatile light-absorbing organics in biomass burning particles.

3.3.3 Absorption enhancement ($E_{abs}$)

The impact of other particle components on BC absorption, either internally or externally mixed of BC with organic aerosol and inorganic salts, has drawn significant attention. The light absorption by an absorbing core can be enhanced when coated with a purely scattering shell which acts as a lens. Absorption enhancement has been observed in laboratory for BC particles coated with various materials (Schnaiter et al., 2005; Zhang et al., 2008), and in field observation (Schwarz et al.,
2008; Spackman et al., 2010). In this study, we measured the absorption enhancement of freshly emitted straw combustion particles.

The light absorption enhancement ($E_{abs}$) due to coating was estimated by the ratio of $b_{abs} (\lambda)$ for particles that did and did not pass through the TD:

$$E_{abs} (\lambda, T) = \frac{b_{abs} (\lambda, T)_{TD}}{b_{abs} (\lambda, T)_{TD}} 
$$

where $T$ is the TD temperature (150 or 300 °C), $T_0$ is the room temperature (20 °C in this work).

As heating by the TD, the particles might shrink to smaller sizes. A tandem DMA (TDMA) was utilized to detect the size change of particles. Here, we used the ratio of the particle diameter after heating ($d_{a2}$) to the diameter before heating ($d_{a1}$) as the shrink factor ($d_{a2}/d_{a1}$) of particles (shown in Figure S7). The absorption coefficient of particles was calibrated by the shrink factor and transport efficiency as mentioned in Section 2.5.

The size-resolved $E_{abs}$ observed at wavelengths of 530 nm and 450 nm are shown in Figure 5. Though freshly emitted, absorption enhancements ($E_{abs}$) of biomass burning particles in the size range of 50-400 nm were observed ($E_{abs} > 1$). Totally, the $E_{abs}$ increased with increasing particle diameters with the largest $E_{abs}$ ($\lambda=530$ nm) of 1.197 ± 0.082 and the $E_{abs}$ ($\lambda=450$ nm) of 1.460 ± 0.101 for 400 nm particles. This could be explained by the thicker coating (both primary and secondary organic and inorganic species) for larger particles. The $E_{abs}$ ($\lambda=450$ nm) were overall larger than the $E_{abs}$ ($\lambda=530$ nm). You et al. (2016) reported that the $E_{abs}$ of BC internally mixed with humic acid (HA/BC) ranged from 2 to 3 and was strongly wavelength dependent.

Removal of the HA absorption contribution revealed the independence of wavelength. Thus, the larger $E_{abs}$ ($\lambda=450$ nm) in this work could be due to the absorption of light-absorbing organics.

4 Conclusions

As a major primary source of aerosols, biomass burning emissions significantly impact the global radiative budget. It is important to understand the physical and chemical properties of biomass burning particles, as well as their links to optical properties. In this work, rice straw was combusted as a representative material of biomass burning in Southern China. A series of comprehensive methods was used to detect the size-resolved chemical composition, effective density, and optical properties of the particles emitted from the burns.

Two methods were utilized to measure the effective density of the biomass burning particles. The DMA-APM-CPC system, which has been widely used in chamber and field work, offered size-resolved information on the particle effective density. The DMA-SPAMS system provided physical property and chemical composition at single-particle level. The 50 nm ($d_{a1}$) biomass burning particles had the lowest effective density of $1.15 \pm 0.23$ g/cm$^3$, which was due to the large portion of aggregate black carbon. The apparent single-peak density distribution of 50 nm particles was due to the combination of two modes (BC and organic matter, respectively). The independent modes at 0.9-1.1 g/cm$^3$ shown in the density distribution of 100 nm and 200 nm particles and ~1.92 g/cm$^3$ mode shown in that of
400 nm particles indicated that BC and crystalline species such as KCl in fresh biomass burning particles tended to be externally mixed with organic carbon. With heating by TD, the separation of the effective density distribution modes testified the presence of BC, potassium salts and less volatile OC in the biomass burning particles.

The effective density measured by DMA-SPAMS system was consistent with the result by DMA-APM-CPC method. The dominant modes in the effective density distributions of 200 nm and 400 nm mobility selected particles were 1.40 g/cm$^3$ and 1.35 g/cm$^3$, respectively. The crystalline KCl with an effective density of 2.10 g/cm$^3$ (with BB-KCl type accounting for 32.7%) was observed in the density distribution for 400 nm particles measured by DMA-SPAMS. The proportions of BB-Nitrate, BB-Sulfate, and BB-KCl types in 400 nm mobility selected particles were larger than those in 200 nm mobility selected particles. Compared with 200 nm particles, 400 nm particles showed more spherical morphology but lower effective density, which could be due to the larger proportion of amorphous NH$_4$NO$_3$.

The size-resolved extinction and scattering coefficients were measured by CAPSs at wavelengths of 450 nm and 530 nm. The SSA ($\lambda$=530 nm) for 50 nm particles was the lowest (0.889±0.006) because of the presence of a larger percentage of the strongly light-absorbing black carbon particles in this size mode. The size-resolved SSAs ($\lambda$=450 nm) for biomass burning particles were generally lower than the SSAs ($\lambda$=530 nm). The AAE values in the size range of 50-400 nm particles were all above 1.6, the acceptable standard of brown carbon. The AAE value was the lowest for 50 nm particles (~5.8) while was the highest for 100 nm particles (~6.3).

Compared with 400 nm particles, the proportions of BB-OC and BB-CN, the extremely low-volatile organic compounds, were larger for 200 nm particles which might indicate a higher possibility for the existence of light-absorbing organics. The $E_{abs}$ was observed in freshly emitted biomass burning particles. The $E_{abs}$ increased with larger diameter due to increasing coating thickness. The wavelength-dependent $E_{abs}$ of particles were likely due to the absorption of light-absorbing organics. Our work emphasizes on the complex mixing states of aerosols from primary source. Further research on how particle morphology affects the optical properties of biomass burning particles is needed.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (91544224, 21507010), the Ministry of Science & Technology of China (2012YQ220113-4), the Science & Technology Commission of Shanghai Municipality (14DZ1202900), and the Changjiang Scholars program of the Chinese Ministry of Education.

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Figure 1. Schematic of the instrumental setup. The CAPS, DMA, CPC, APM and SPAMS represent Cavity Attenuated Phase Shift spectroscopy, Differential Mobility Analyzer, Condensation Particle Counter, Aerosol Particle Mass analyzer and Single Particle Aerosol Mass Spectrometer, respectively.
Figure 2. Average density distributions of 50, 100, 200, and 400 nm particles selected by DMA at 20 °C (room temperature), 150 °C, and 300 °C. Gaussian model was applied in fitting each density scan (red and green lines). Black dashes were the assumption Gaussian models application.
Figure 3. Vacuum aerodynamic size distributions detected by the SPAMS of 200 nm (a) and 400 nm (b) electrical mobility size-selected biomass burning particles and pie charts for the particle types in different aerodynamic modes at 20 °C (room temperature).
Figure 4. (a)-(b): Size-resolved single scattering albedo (SSA) at wavelengths of 530 nm and 450 nm. (c): Ångström absorption exponent (AAE) of biomass burning particles at room temperature (20°C).
Figure 5. The size-resolved absorption enhancement ($E_{abs}$) at wavelengths of 450 nm and 530 nm.