

1 **Secondary aerosol formation promotes water uptake**
2 **by organic-rich wildfire haze particles in Equatorial**
3 **Asia**

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1 **Abstract**

2 The diameter growth factor (GF) of 100 nm haze particles at 85 % relative humidity and their
3 chemical characteristics were simultaneously monitored at Singapore in October 2015 during a
4 pervasive wildfire haze episode that was caused by peatland burning in Indonesia. Non-
5 refractory submicron particles (NR-PM₁) were dominated by organics (approximating 77.1 % in
6 total mass), whereas sulfate was the most abundant inorganic constituent (11.7 % on average). A
7 statistical analysis of the organic mass spectra showed that most organics (36.0 % of NR-PM₁
8 mass) were highly oxygenated. Diurnal variations of GF , number fractions of highly hygroscopic
9 mode particles, mass fractions of sulfate, and mass fractions of oxygenated organics (OOA)
10 synchronized well, peaking during the day. The mean hygroscopicity parameter (κ) of the haze
11 particles was 0.189 ± 0.087 , and the mean κ values of organics were 0.157 ± 0.108 (κ_{org} , bulk
12 organics) and 0.266 ± 0.184 (κ_{OOA} , OOA), demonstrating the important roles of both sulfate and
13 highly oxygenated organics in the hygroscopic growth of organics-dominated wildfire haze
14 particles. κ_{org} correlated with the water-soluble organic fraction insignificantly, but it positively
15 correlated with f_{44} (fraction of the ion fragment at m/z 44 in total organics) ($R = 0.70$), implying
16 the oxygenation degree of organics could be more critical for the water uptake of organic
17 compounds. These results further suggest the importance of sulfate and secondary organic
18 aerosol formation in promoting the hygroscopic growth of wildfire haze particles. Further
19 detailed size-resolved as well as molecular-level chemical information about organics is
20 necessary for the profound exploration of water uptake by wildfire haze particles in Equatorial
21 Asia.
22

1 **1. Introduction**

2 In the last few decades, wildfire haze has periodically raged throughout equatorial Asian
3 countries (Page et al., 2002; van der Werf et al., 2010; Field et al., 2016; Koplitz et al., 2016),
4 resulting in billions of dollars of economic losses as well as thousands of premature deaths
5 (Johnston et al., 2012; Marlier et al., 2013). The increasing wildfire activity is associated with the
6 recent rapid change in land use for agricultural development, including industrial plantations
7 over peatland (Page et al., 2009; Marlier et al., 2015; Spracklen et al., 2015). Such developments
8 are accompanied by the drainage of water of pristine peat swamp forest, making the tropical
9 peatland susceptible to fire (Langner et al., 2007; Konecny et al., 2016). The occurrence of
10 peatland fires is closely related to El Niño-induced droughts (Page et al., 2002; Field et al., 2016).
11 Enhanced peatland fires have been observed during intense El Niño years, including 1997, 2006,
12 and 2015 (Page et al., 2002; van der Werf et al., 2010; Stockwell et al., 2016). In 1997, the
13 peatland fire was of global concern, as the total carbon emission was estimated to equal 13–40 %
14 of the year’s annual global carbon emission from fossil fuels (Page et al., 2002). The recent
15 equatorial Asian wildfire haze event in 2015 could rival the one in 1997 not only in terms of the
16 hazards to human health but also the significant impacts on global climate (Crippa et al, 2016;
17 Field et al., 2016; Huijnen et al., 2016; Koplitz et al., 2016; Stockwell et al., 2016). During
18 September–October 2015, thick smoke from peatland fires blanketed Equatorial Asia and
19 released huge amounts of organic material and fine particulate matter (particulate matter of
20 aerodynamic diameter below 2.5 μm , $\text{PM}_{2.5}$) (Crippa et al, 2016; Koplitz et al., 2016), which is
21 the leading cause of global air-pollution-related mortality (Kunii et al., 2002; World Health
22 Organization, 2009; Johnston et al., 2012; Marlier et al., 2013; Lelieveld et al., 2015).

23 A previous study on the peatland fire event in 1997 has reported that the wildfire haze
24 particles resulted in dramatic cooling effects on the atmospheric radiative budget, especially over
25 the source region of Indonesia (-150 W m^{-2}) and the tropical Indian Ocean (-10 W m^{-2})
26 (Duncan et al., 2003). In addition to the subsequently affected shallow warm clouds and deep
27 convection processes, the resultant abnormal rainfall in adjacent tropical region and the extra-
28 tropics was also confirmed by both satellite observations and model simulations (Rosenfeld,
29 1999). These studies demonstrate the importance of investigating aerosol-cloud-precipitation

1 interactions of Indonesian wildfire haze particles, including water uptake properties of aerosol
2 particles.

3 Previous studies on water uptake properties of aerosol particles stemmed from Indonesian
4 peatland fires are controversial. On one hand, laboratory studies have demonstrated that fresh
5 Indonesian peat burning particles are weakly hygroscopic and almost inactive as cloud
6 condensation nuclei (CCN) (Chand et al., 2005; Dusek et al., 2005; Chen et al., 2017). On the
7 other hand, a field observation showed that the wildfire haze particles were highly hygroscopic
8 during the 1997 Indonesian peatland fires (Gras et al., 1999). Such discrepancies impede reliable
9 evaluations of the effects on regional and global climate changes driven by Indonesian wildfire
10 haze particles (Lin et al., 2013; Reid et al., 2013). The cause of the discrepancies needs to be
11 understood quantitatively in comparison with the chemical composition of particles.

12 Aerosol particles emitted from wildfires are a mixture of inorganic and organic
13 compounds, which complicates their water uptake properties (Carrico et al., 2008, 2010; Petters
14 et al., 2009; Hallar et al., 2013; Latham et al., 2013). Water uptake properties of inorganic salts,
15 such as ammonium sulfate and ammonium nitrate, are well known, yet the hygroscopic behavior
16 of organic compounds or organic-inorganic mixtures is still difficult to predict due to the
17 complex chemical composition of organics and the associated distinct affinity for water of a
18 specific chemical constituent (Saxena et al., 1995; Gysel et al., 2004; Dinar et al., 2007; Petters
19 and Kreidenweis, 2007; Carrico et al., 2010; Kristensen et al., 2012; Marsh et al., 2017). For
20 instance, experimental and modeling studies have shown that the water uptake by water-soluble
21 matter is governed by the inorganic fraction, whereas the hygroscopic properties of inorganics
22 can be altered substantially by the presence of organics (Saxena et al., 1995; Dick et al., 2000).
23 In general, with the increase in the organic fraction, the water uptake by wildfire particles has
24 presented an overall decreasing trend (Mircea et al., 2005; Carrico et al., 2010), evidencing the
25 high sensitivity of particle water uptake to organic fractions. The high sensitivity to organic
26 fractions has been observed, as inorganic species are much more hygroscopic than most organic
27 compounds. However, the roles of inorganic and organic species in water uptake by Indonesian
28 peatland burning particles have rarely been investigated (Dusek et al., 2005; Chen et al., 2017).

1 Water uptake properties of organic compounds have also been demonstrated to be
2 important especially when the chemical composition of aerosol particles is dominated by
3 organics. Such cases have been frequently observed in particles emitted from wildfires (Petters et
4 al., 2009; Carrico et al., 2010; Cubison et al., 2011; Hallar et al., 2013; Chen et al., 2017). Both
5 theoretical and experimental studies demonstrated that water-soluble organic matter (WSOM)
6 plays the key role in determining the water uptake by organic compounds (Peng et al., 2001;
7 Gysel et al., 2004; Petters and Kreidenweis, 2007; Carrico et al., 2008; Petters et al., 2009;
8 Latham et al., 2013; Chen et al., 2017). For instance, freshly emitted peat burning particles are
9 known to contain a small fraction of WSOM, which explains their limited hygroscopicity (Chen
10 et al., 2017). Chemical aging and oxidation of organic compounds in both gas and particle phases
11 could alter the water uptake properties of aerosol particles in a wildfire plume (Gras et al., 1999;
12 Petters et al., 2009; Rose et al., 2010; Cubison et al., 2011). These chemical processes in the
13 atmosphere enhance the fractions of highly oxygenated organics and polar species, which are
14 typically water soluble (Duplissy et al., 2008, 2011; Jimenez et al., 2009; Chang et al., 2010;
15 Massoli et al., 2010; Cubison et al., 2011; Cerully et al., 2015). Although such chemical
16 transformation and corresponding changes in hygroscopicity have been observed in wildfire
17 particles in both the laboratory and the field (Petters et al., 2009; Massoli et al., 2010; Rose et al.,
18 2010; Cubison et al., 2011; Duplissy et al., 2011), the importance of these processes for water
19 uptake properties has never been investigated for peatland burning particles in the equatorial
20 Asian region.

21 In this work, we investigated the relationships between water uptake properties and the
22 chemical composition of aerosol particles in a tropical peatland fire haze in October 2015 by
23 conducting atmospheric observations in Singapore. We quantified water uptake properties using
24 the Humidified Tandem Differential Mobility Analyzer (HTDMA). In parallel, particle chemical
25 composition was characterized in real-time using the Aerodyne Time of Flight-Aerosol Chemical
26 Speciation Monitor (ToF-ACSM). Furthermore, water-soluble organic carbon (WSOC) and
27 elemental carbon (EC) contents were quantified by an off-line analysis of ambient PM_{2.5} filter
28 samples. The data from these measurements were combined to explore how the water uptake
29 properties of tropical peatland burning particles are regulated.

30

1 **2. Observation**

2 **2.1. Field Campaign**

3 The field observation was conducted at the campus of Nanyang Technological University
4 (NTU), Singapore (1°20'41" N, 103°40'53" E) during October 2015. The campus is located
5 20 km away from the city center, and surrounded by a secondary tropical forest and grassland.
6 The site is located 0.8 km away from a highway, and a petrochemical complex (Jurong Island) is
7 located approximately 8 km south. In September–October 2015, the observation site encountered
8 severe transboundary haze pollution that was caused by recurring Indonesian peatland fires,
9 dominated by the smoldering combustion of underground organic-rich peat soils and mixed
10 surface vegetation burning (Field et al., 2016; Page et al., 2009; Jayaranthe et al., 2017). Particles
11 emitted from the wildfires had experienced for approximately 1–4 days atmospheric aging
12 process before arriving at Singapore (Fig. S1).

13 The observation was performed in an air-conditioned room, with the room temperature
14 maintained at 22 °C. A cyclone (URG-2000-30EN PM_{2.5}, URG) was employed for ambient
15 aerosol sampling at a flow rate of 16.67 L min⁻¹. The inlet was fixed on the rooftop, which was
16 located approximately 10 m above the ground. The sample air was split into several flows for
17 measurements with different instruments after drying by diffusion dryers (the relative humidity,
18 *RH*, of the sample flow was below 30 %). During the observation, particle number size
19 distribution, chemical composition, and hygroscopic growth were monitored.

20 **2.2. Particle water uptake measurements**

21 Water uptake properties were measured using the HTDMA system (Chen et al., 2017).
22 Briefly, the sampled particles were desiccated using a diffusion dryer (Model 42000, Brechtel
23 Manufacturing, Inc.), and the resulting dry polydisperse particles were classified by the first
24 differential mobility analyzer (DMA, Model 3081, TSI Inc.). The DMA selects particles of a
25 specific mobility diameter (D_0), which was fixed at 100 nm during the observation. The
26 classified particles were humidified to $RH = 85\%$ using nafion tubings (MD-110-12S-4, Perma
27 Pure) operated under a controlled *RH* condition. The particle residence time in the humidifier
28 was approximately 10 seconds. The variation in *RH* was $\pm 0.5\%$ (peak to peak). The resulting
29 size distribution of humidified particles was measured by the second DMA coupled with a

1 condensation particle counter (CPC, Model 3775, TSI Inc.). The diameter growth factor
2 parameter, g , which is defined as the ratio of the particle diameter after humidification at a
3 conditioned RH ($D_p(RH)$) to the initial dry size (D_0) (i.e., $g = D_p(RH)/D_0$), was calculated from
4 the HTDMA data. Hygroscopicity parameter, κ , was derived from the corresponding g at a given
5 RH and D_0 using the following equation (Petters and Kreidenweis, 2007):

$$6 \quad \kappa = (g^3 - 1) \cdot \left(\frac{1}{RH} \cdot \exp\left(\frac{4\sigma_{s/a} \cdot M_w}{\rho_w \cdot R \cdot T \cdot D_0 \cdot g} \right) - 1 \right), \quad (1)$$

7 where $\sigma_{s/a}$ is the surface tension of the solution/air interface (0.0718 N m^{-1} at $25 \text{ }^\circ\text{C}$), M_w and ρ_w
8 are the molecular weight ($0.018 \text{ kg mol}^{-1}$) and density of water ($1 \times 10^3 \text{ kg m}^{-3}$), respectively; R
9 is the universal gas constant ($8.31 \text{ J K}^{-1} \text{ mol}^{-1}$), and T is the absolute temperature (298 K). The
10 HTDMA calibration results with 150 nm ammonium sulfate particles are shown in Fig. S2,
11 demonstrating the validity of our instrument. Further details about the HTDMA are available in
12 Chen et al. (2017).

13 **2.3. Aerosol chemical analysis**

14 The ToF-ACSM (Aerodyne Inc.) was utilized to measure the chemical composition of
15 non-refractory submicron particles (NR- PM_{10}), including organics (OA), sulfate (SO_4^{2-}), nitrate
16 (NO_3^-), ammonium (NH_4^+), and chloride (Cl^-) (Fröhlich et al., 2013). The ToF-ACSM sampled
17 particles that were desiccated by a nafion tubing. The organic mass spectra measured by the ToF-
18 ACSM were analyzed in detail using the Multilinear Engine (ME-2 solver) software (Canonaco
19 et al., 2013). Four specific types of OA were identified: hydrocarbon-like OA (HOA), peat
20 burning OA (PBOA), non-peat biomass burning OA (briefly BBOA), and oxygenated OA
21 (OOA). The HOA factor was mainly contributed by primary sources, such as the combustion
22 emissions from fossil fuel (e.g., related to traffic, shipping, and industrial use), excluding the
23 influence of biomass burning. The PBOA and BBOA factors were attributed to the long-range
24 transport from Indonesian peatland fires, and the two OA factors were well separated. Details
25 about the ToF-ACSM measurements and data analysis are provided in Budisulistiorini et al.
26 (2018).

1 PM_{2.5} filter samples for chemical analysis were also collected using filter holders (BGI
2 Inc.). The samples were collected for 24 hours using 47 mm (diameter) quartz-fiber filters. The
3 sampling started/ended at 08:00 local time (LT). The collected samples were analyzed for bulk
4 OC, EC, and WSOC. All the quartz-fiber filters were prebaked at 900 °C for 3 hours before
5 sampling, after which they were stored in a refrigerator (−20 °C) until analysis. For each
6 sampling, a back-up quartz-fiber filter was used to account for the potential influence of gas
7 phase organic components on the particulate organics collected on the front quartz-fiber filter
8 (Turpin et al., 1994). The method assumes that all the particulate OC is collected by the front
9 filter, while the gas phase OC is collected equally on both front and back filters. Subtraction of
10 the OC loading on the back filter (i.e., gas phase OC) from that on the front one allows
11 quantification of particulate OC (i.e., corrected OC).

12 Concentrations of OC and EC were determined by thermal-optical reflectance analysis
13 (Chow et al., 1993) using the Sunset Laboratory OC/EC Analyzer, following the IMPROVE-A
14 protocol. WSOC was quantified with the Sievers 800 Total Organic Carbon (TOC) Analyzer
15 following extraction of part of a filter sample (8 mm ϕ) by 10 ml of HPLC-grade water. An
16 orbital shaker was operated for 21 hours for the extraction, and the subsequent solutions were
17 filtered with syringe filters (pore size of 0.2 μ m).

18 **2.4. Particle number size distribution**

19 Particle number size distributions were measured using a NanoScan SMPS Nanoparticle
20 Sizer (NanoScan-SMPS, Model 3910, TSI Inc.) and an Optical Particle Sizer (OPS, Model 3330,
21 TSI Inc.). The detected particle sizes ranged from 11.5–365.2 nm (NanoScan-SMPS) in mobility
22 size and 0.3–10 μ m (OPS) in optical size. Both instruments sampled particles desiccated by a
23 diffusion dryer (Model 42000, Brechtel Manufacturing, Inc.). The time resolution of both
24 instruments was 1 minute.

25

26 **3. HTDMA data analysis**

27 **3.1. Classification of three hygroscopic modes**

1 Figure 1 shows the HTDMA data averaged over the entire observation period. The mean
 2 normalized particle size distribution after humidification at 85 % *RH* has spanned a few different
 3 modes, reflecting mixing states of ambient wildfire haze particles observed at Singapore
 4 (Bougiatioti et al., 2016; Ogawa et al., 2016). For particles in a specific hygroscopic mode, *i*,
 5 with $g_{1,i} < GF < g_{2,i}$, the number fraction of this mode (nf_i) was derived from measured
 6 probability density function of *g* (i.e., $c(g, D_0)$) as $nf_i = \int_{g_{1,i}}^{g_{2,i}} c(g, D_0) dg$. The corresponding
 7 mean *GF* ($g_{mean,i}$) was calculated from $g_{mean,i} = \frac{1}{nf_i} \int_{g_{1,i}}^{g_{2,i}} g c(g, D_0) dg$ (Gysel et al., 2009). The
 8 equivalent values of κ for mode *i* were obtained from $g_{mean,i}$ using Eq. (1).

9 The observed 100 nm dry particles were categorized into the following three groups
 10 based on their hygroscopic properties at *RH* = 85 % (temporal variation of the multimodal size
 11 distribution patterns are shown in Fig. S3 of the supplementary material), facilitating the analysis
 12 of heterogeneity of particle chemical composition:

13 (1) *Nearly non-hygroscopic or weakly hygroscopic particles* ($0 \leq \kappa < 0.1$; $g < 1.15$): Particles are
 14 predominantly composed of black carbon (BC) and non-polar hydrocarbon-like organic
 15 compounds (Peng et al., 2001; Gysel et al., 2007; Kreidenweis et al., 2008).

16 (2) *Moderately hygroscopic particles* ($0.1 \leq \kappa < 0.2$; $1.15 \leq g < 1.27$): They could contain
 17 hygroscopic organics (e.g., carboxylic acids and levoglucosan) and/or mixtures of non-/less and
 18 more hydrophilic compounds (e.g., BC, fatty acids, and/or humic-like substances mixed with
 19 ammonium sulfate or levoglucosan-like species) (Peng et al., 2001; Chan and Chan., 2003; Gysel
 20 et al., 2004, 2007; Chan et al., 2005; Petters and Kreidenweis, 2007);

21 (3) *More hygroscopic particles* ($\kappa \geq 0.2$; $g \geq 1.27$): Aerosol particles contain inorganic salts as
 22 well as some more hygroscopic organic species such as multifunctional organic acids (Peng et al.,
 23 2001; Carrico et al., 2008; Duplissy et al., 2011; Ogawa et al., 2016).

24 In addition, the volume-weighted mean growth factor, *GF*, was also calculated using $c(g,$
 25 $D_0)$ (Gysel et al., 2009):

$$GF = \left(\int_0^\infty g^3 \cdot c(g, D_0) \cdot dg \right)^{1/3}. \quad (2)$$

GF was employed to calculate the mean values of κ , which facilitated the comparison with chemical composition of aerosol particles.

3.2. Effective κ of organic compounds (κ_{org})

Water uptake properties of organic compounds were estimated using the Zdanovskii–Stokes–Robinson (ZSR) mixing rule, employing the observed values of κ and chemical composition as input parameters. The ZSR mixing rule assumes that water uptake by a mixture of materials is additive of the water content retained by each chemical species (Stokes and Robinson, 1966). The rule also assumes that the volume change in the mixing of species within individual particles is almost negligible (Brechtel and Kreidenweis, 2000, Gysel et al., 2007; Petters and Kreidenweis, 2007):

$$\begin{aligned} \kappa &= \sum_i \kappa_i \cdot \varepsilon_i = \kappa_{SNA} \cdot \varepsilon_{SNA} + \kappa_{org} \cdot \varepsilon_{org} + \kappa_{EC} \cdot \varepsilon_{EC} \\ \Leftrightarrow \kappa_{org} &= \frac{\kappa - \kappa_{SNA} \cdot \varepsilon_{SNA} - \kappa_{EC} \cdot \varepsilon_{EC}}{\varepsilon_{org}}, \end{aligned} \quad (3)$$

where κ_i and ε_i stand for the hygroscopicity parameter and the volume fraction of a specific component i in dry particles, respectively. The subscript SNA represents the three major inorganic constituents of sulfate, nitrate, and ammonium; org denotes organic species; EC indicates elemental carbon.

Sulfate, ammonium, and nitrate were considered for inorganics, the majority of which were contributed by sulfate (see Table 2). Other materials, such as sea salt and crustal elements, were demonstrated to be neglected because they are relatively scarce in submicron wildfire haze particles in Southeast Asia (Balasubramanian et al., 2003; Keywood et al., 2003; See et al., 2006; Stockwell et al., 2016). Almost all the sulfate and nitrate were neutralized by ammonia, and these three most abundant inorganic constituents were combined and assumed as ammonium sulfate (i.e., $\varepsilon_{SNA} = \varepsilon_{SO4} + \varepsilon_{NO3} + \varepsilon_{NH4} \approx \varepsilon_{AS}$). Thus, the value of κ_{SNA} was considered to approximate the κ value of ammonium sulfate under the condition when sulfate dominates inorganics (Gunthe et al.,

1 2009; Chang et al., 2010; Ogawa et al., 2016). The elemental carbon (EC) is known as non-
2 hygroscopic (i.e., $\kappa_{EC} \approx 0$).

3 Values of densities are required to compute ε_i from observed mass fractions. The mass
4 fraction is taken as the first-order approximation of the volume fraction, based on the hypothesis
5 that the bulk particle density is similar to the densities of individual compounds when volume
6 additivity is assumed (Kreidenweis et al., 2008; Gunthe et al., 2009; Hallquist et al., 2009). This
7 hypothesis has been demonstrated to be acceptable when particles are composed primarily of
8 organics and sulfate (Cross et al., 2007; King et al., 2007). The densities of ammonium sulfate
9 and EC were assumed to be 1.77 g cm^{-3} and 1.80 g cm^{-3} , respectively (Park et al., 2004; Bond
10 and Bergstrom, 2006). The density of organics is known to vary depending on their elemental
11 composition (Kuwata et al., 2012). The value was assumed as 1.40 g cm^{-3} , which is a typical
12 value for ambient organic aerosols (Hallquist et al., 2009).

13 The EC mass fraction of approximately 10.0 % in submicron wildfire haze particles was
14 utilized according to the time-averaged EC content in ambient $\text{PM}_{2.5}$ filter samples (see Table 2).
15 This assumption was based on the preconditions that the wildfire haze particles were
16 homogeneously mixed in varied sizes and that there was no significant difference between BC
17 and EC. It should be noted that some difference in chemical composition could have existed
18 between particles quantified by the ToF-ACSM (NR-PM_1) and HTDMA (100 nm). Although the
19 data obtained by these two techniques are frequently combined for detailed analysis (e.g.,
20 Whitehead et al., 2016), the difference in particle size range along with the particle mixing state
21 (Whitehead et al., 2014) can be a source of uncertainty for the following discussion in this work.

22 **3.3. κ of oxygenated organic compounds (κ_{OOA})**

23 As described in Sect. 2.3, organics were numerically segregated to HOA, PBOA, BBOA,
24 and OOA. The value of κ_{org} can be calculated by a linear combination of contributions from
25 segregated fractions (Petters and Kreidenweis, 2007; Chang et al., 2010):

$$26 \quad \kappa_{\text{org}} = v_{\text{HOA}} \cdot \kappa_{\text{HOA}} + v_{\text{PBOA}} \cdot \kappa_{\text{PBOA}} + v_{\text{BBOA}} \cdot \kappa_{\text{BBOA}} + v_{\text{OOA}} \cdot \kappa_{\text{OOA}}, \quad (4)$$

27 where v_i stands for the volume fraction of component i in all the organics.

1 Water uptake by a mixed particle is largely driven by the relative abundance of more or
2 less hygroscopic component, and it is more sensitive to uncertainties in the hygroscopicity of
3 more hygroscopic compounds than that of less hygroscopic compounds (Gysel et al., 2007).
4 Hydrocarbon (-like) OA is known to be almost non-hygroscopic, leading to the estimation that
5 the κ value of HOA is 0 (Gysel et al., 2007; Gunthe et al., 2009; Chang, et al., 2010). Water
6 uptake by freshly emitted biomass burning particles is generally limited, especially compared
7 with OOA (Carrico et al., 2010; Chang et al., 2010; Chen et al., 2017). Based on our previously
8 measured κ results of fresh Indonesian peatland burning particles, an overall mean value of κ_{PBOA}
9 = 0.04 was used for the fresh PBOA factor, and $\kappa_{BBOA} = 0.06$ was applied for the non-peat BBOA
10 factor with mixed vegetation burning mainly of acacia and fern species (Chen et al., 2017).
11 Under these assumptions, κ_{OOA} can be calculated by the following equation:

$$12 \quad \kappa_{OOA} = (\kappa_{org} - \kappa_{PBOA} \cdot \nu_{PBOA} - \kappa_{BBOA} \cdot \nu_{BBOA}) / \nu_{OOA} \quad (5)$$

13 The density of OOA/PBOA/BBOA is required to calculate $\nu_{OOA}/\nu_{PBOA}/\nu_{BBOA}$ using the
14 ME-2 resolved corresponding mass concentration combined with the total volume concentration
15 of the bulk OA derived from the ToF-ACSM observed OA mass. The density of oxygenated
16 organics was assumed to be 1.50 g cm^{-3} (as summarized in Table 1), which is a typical value for
17 carboxylic and multifunctional organic acids (Saxena et al., 1995; Peng et al., 2001; Gysel et al.,
18 2004; Carrico et al., 2010; Ogawa et al., 2016). According to the reported average density for
19 common dry smoke particles (i.e., $1.20 - 1.40 \text{ g cm}^{-3}$; Reid et al., 2005), and considering the
20 dominance of non-refractory organic material in fresh Indonesian peatland burning particles
21 (Budisulistiorini et al., 2017), the densities of PBOA and non-peat BBOA were assumed to be
22 1.10 and 1.20 g cm^{-3} , respectively. Detailed information on the parameters utilized for the κ_{org}
23 calculation is provided in Table 1.

24

25 **4. Results**

26 In this section, aerosol number size distribution (Figure 2), chemical composition
27 (Figures 3 and 4), and hygroscopic properties of aerosol particles (Figure 5) are described in
28 addition to diurnal variations in these data (Figure 6).

1 **4.1 Number size distribution of wildfire haze particles**

2 Figure 2a displays the time-averaged particle size distribution within the entire size range
3 of 11.5 nm–10 μm measured by NanoScan-SMPS combined with OPS. The NanoScan-SMPS
4 data were used for the fine particles (11.5–365.2 nm), and the overlapped size range of the OPS
5 (338 nm size bin) was excluded from the analysis. The data in the remaining OPS size range
6 (419 nm–10 μm) were combined with the fine particle data. The temporal average size
7 distribution presented a unimodal structure, in which the number mode diameter was located
8 around 100 nm. Particles in the range of 30–200 nm dominated the total particle number
9 concentration, whereas particles larger than 600 nm accounted for a minor fraction (less than 4.0 %
10 on average; Fig. S4 and Fig. S5). This result suggests that in Singapore, the wildfire haze
11 particles were predominantly contributed by submicron particles, in line with the corresponding
12 chemical characteristics obtained in previous studies (Balasubramanian et al., 2003; See et al.,
13 2006).

14 Figure 2b shows the mean diurnal cycle of particle number size distribution. The growth
15 of ultrafine particles was typically observed in the afternoon. The $dN/d\log D_p$ higher than
16 $1.5 \times 10^4 \text{ cm}^{-3}$ was commonly observed in the 50–200 nm particle size range, while the
17 $dN/d\log D_p$ of super micron particles seldom exceeded $1.0 \times 10^3 \text{ cm}^{-3}$. The particle number
18 concentration was high from 08:00–09:00 LT in the morning. The concentration increased again
19 in the afternoon (about 14:00 LT), which lasted until midnight. The high concentration periods
20 could have been caused by local traffic emissions and by secondary formation processes. Diurnal
21 variations were also observed in the number concentrations of all particles.

22 Figure 2c depicts the mean diurnal variations of the corresponding total particle number
23 and volume concentrations. The total number concentration started to increase after 07:00 LT
24 until around 10:00 LT, and it reached the highest level after 14:00 LT. The particle number
25 concentration was higher than $1.5 \times 10^4 \text{ cm}^{-3}$ before 19:00 LT. After that, the number
26 concentration decreased gradually, reaching $1.2 \times 10^4 \text{ cm}^{-3}$ at midnight. Correspondingly, the
27 aerosol volume concentration was higher than $50.0 \mu\text{m}^3 \text{ cm}^{-3}$ during the day. The volume
28 concentration decreased during the night, although it was still higher than $45.0 \mu\text{m}^3 \text{ cm}^{-3}$. These

1 results demonstrate that the aerosol loading was significantly high during the period of pollution
2 caused by wildfire haze.

3 **4.2 Chemical characteristics of wildfire haze particles**

4 Figures 3a-b show the time series of both the mass concentrations and the corresponding
5 mass fractions of organics, sulfate, nitrate, ammonium, and chloride (expressed as f_{org} , $f_{SO_4^{2-}}$,
6 $f_{NO_3^-}$, $f_{NH_4^+}$, and f_{Cl^-} , respectively) in NR-PM₁ quantified by the ToF-ACSM. The average mass
7 loading of NR-PM₁ was as high as $44.7 \pm 24.5 \mu\text{g m}^{-3}$, confirming the severity of the pervasive
8 wildfire haze. During the observation period, organics were always the most abundant
9 compounds in NR-PM₁ ($34.8 \pm 20.7 \mu\text{g m}^{-3}$). The mass concentration of organics was higher
10 than $50.0 \mu\text{g m}^{-3}$ in many cases, and it occasionally exceeded $100.0 \mu\text{g m}^{-3}$. On average,
11 organics accounted for the highest mass fraction of 77.1 %, followed by sulfate (11.7 %),
12 ammonium (6.4 %), and nitrate (4.2 %). The mass concentration of non-refractory chloride was
13 almost negligible (0.6 % of the total mass). These results demonstrate that the submicron wildfire
14 haze particles were predominantly composed of organics.

15 Table 2 summarizes the mass concentrations of all the analyzed inorganic ionic species in
16 the PM_{2.5} filter samples. The corresponding data for carbonaceous fractions are presented in
17 Fig. 4. Overall, the mass fraction of EC varied from 4.4 % to 15.8 % with a mean value of
18 10.8 %. OC occupied 30.4–50.7 % of the total PM_{2.5} mass concentration, and the mean fraction
19 was 43.0 %. The WSOC fraction was in the range of 17.6–30.2 % with the mean level of 26.7 %.
20 Correspondingly, the water-insoluble OC (WISOC) content was calculated to be 6.1–20.5 %
21 with the mean fraction of 16.3 %. The WSOC/OC ratios were consistently higher than 50.0 %
22 with a mean and maximum values of 63.6 % and 79.9 %, respectively. The mean WSOC/OC
23 value was significantly higher than that for fresh Indonesian peat burning particles emitted from
24 the source region (i.e., 16 %; Jayaranthe et al., 2017), which were demonstrated to be generally
25 water insoluble and thus nearly non-hygroscopic (Chen et al., 2017). This result suggests that the
26 majority of organics in the wildfire haze particles were water soluble, implying the importance of
27 secondary formation as well as the chemical transformation of organic particles during
28 atmospheric transport. Inorganic ions were less abundant and less variable than the organics were.
29 On average, the inorganics accounted for 30.5 % of the PM_{2.5} mass loading with the mean

1 contributions of 0.6 % by Cl^- , 2.6 % by NO_3^- , 17.2 % by SO_4^{2-} , 0.4 % by Na^+ , 8.1 % by NH_4^+ ,
2 0.8 % by K^+ , 0.1 % by Mg^{2+} , and 0.7 % by Ca^{2+} . Sulfate, ammonium, and nitrate were the most
3 abundant inorganic components. More than half of the inorganics was contributed by sulfate.
4 These results show that the wildfire haze particles were dominated by organics especially water-
5 soluble species. Mass concentrations of organics measured by the $\text{PM}_{2.5}$ filter samples and by the
6 ToF-ACSM agreed well when the organics/OC ratio was assumed to be 1.4 (slope = 1.07; $R^2 =$
7 0.91) (Reid et al., 2005; Hallquist et al., 2009; Levin et al., 2010). The total mass concentrations
8 of aerosol particles quantified by the filter samples and the ToF-ACSM also correlated well ($R^2 =$
9 0.96). The mass loading of the $\text{PM}_{2.5}$ filter samples was approximately 30 % higher than that of
10 the ToF-ACSM results, likely because of the difference in particle size range and the lack of EC
11 content for the ToF-ACSM measurements (Budisulistiorini et al., 2018).

12 Figure 3c shows the mean mass spectra of organics averaged over the observation period.
13 Ion signals at m/z 43 (most likely $\text{C}_2\text{H}_3\text{O}^+$) and m/z 44 (CO_2^+) were prominent, accounting for
14 7.5 % and 10.5 % of the total organics mass spectrum. The predominant signal of m/z 44
15 indicates that organic compounds in wildfire haze particles were highly oxygenated. High-
16 molecular weight organic signals with $m/z > 100$ possessed an abundance of 13.3 %. The marker
17 ions for biomass burning particles, such as m/z 60 (mostly $\text{C}_2\text{H}_4\text{O}_2^+$) and m/z 73 (mainly
18 $\text{C}_3\text{H}_5\text{O}_2^+$), which originating in levoglucosan-like species (e.g., levoglucosan, mannosan, and
19 galactosan) were also clearly observed (Cubison et al., 2011).

20 Figures 6d-e show the mean contributions of four organic components classified by the
21 ME-2 method, including HOA (8.8 % of NR- PM_{10} mass), PBOA (10.4 %), BBOA (10.0 %), and
22 OOA (36.0 %). The primary organics originating from biomass burning (i.e., PBOA and BBOA)
23 accounted for 20.4 %. OOA was the dominant type of organics during the wildfire haze episodes.

24 **4.3 Hygroscopic properties of wildfire haze particles**

25 Figure 5 displays the time series of the mean GF data as well as the corresponding values
26 of κ during the entire observation period. The mean values of GF varied between 0.98 and 1.52
27 with the average of 1.25 ± 0.09 . GF values larger than 1.40 were normally observed in the
28 afternoon. The variation in the corresponding κ results ranged from 0.004–0.475, and the
29 average κ value was 0.189 ± 0.087 . Table 3 summarizes the mean κ results for organics (cf.

1 Sect.3.2, with EC considered), which were calculated using the HTDMA and ToF-ACSM
2 measurements during the overlapping observation period from 10–24 October 2015. The
3 HTDMA-derived bulk κ results averaged over the same overlapping period were defined as
4 κ_{HTDMA} . The mean κ_{org} (0.157 ± 0.108) was lower than the mean κ_{OOA} (0.266 ± 0.184), as the
5 whole organic fraction normally contains both non-hygroscopic and hygroscopic organics. The
6 derived κ_{org} results are demonstrated to be comparable to previously reported κ values for bulk
7 organics (Petters and Kreidenweis, 2007; Duplissy et al., 2011; Lathem et al., 2013; Cerully et al.,
8 2015). Moreover, the mean κ_{OOA} value agreed well with mean κ results in previous field and
9 laboratory studies (e.g., $\kappa = 0.25 \pm 0.06$ for biogenic OOA, Chang et al., 2010; $\kappa = 0.2 \pm 0.1$ for
10 aged BBOA, Engelhart et al., 2012). A caveat of this comparison is the representativeness of
11 laboratory experiments for the actual environmental conditions, including types of burnt
12 biomasses, concentrations and types of oxidants, and aging time. As the atmospheric condition of
13 tropical Asia is unique, a systematic study of the chemical aging process of wildfire plume for
14 the region would be required in the future. Note that the mean κ_{OOA} was even higher than the
15 corresponding mean κ_{HTDMA} (0.207 ± 0.093), revealing that the water uptake particularly by some
16 highly oxygenated organics in the wildfire haze particles could be highly significant.

17 **4.4 Diurnal variations of hygroscopic properties and chemical composition**

18 *GF* exhibited a clear pattern of diurnal variation (Fig. 6a). Higher *GF* values were
19 observed during the daytime ($GF = 1.27 \pm 0.05$ for 08:00–20:00 LT). However, the values were
20 lower in the early morning and at night (1.23 ± 0.05 for 20:00–08:00 LT). The corresponding
21 mean bulk κ results averaged over the entire observation period were 0.213 ± 0.051 for the
22 daytime and 0.172 ± 0.043 for nighttime.

23 Similarly, the daytime mean κ_{org} and κ_{OOA} were 0.200 ± 0.104 and 0.323 ± 0.177 ,
24 respectively, whereas the nighttime mean values were 0.103 ± 0.086 (κ_{org}) and 0.186 ± 0.177
25 (κ_{OOA}) (Table 3). These mean κ values were 19.2 % lower (κ_{org}) and 30.8 % higher (κ_{OOA}) than
26 the concurrently measured mean κ_{HTDMA} result of 0.247 ± 0.096 (daytime), whereas they were
27 35.9 % lower (κ_{org}) and 16.3 % higher (κ_{OOA}) than that of 0.160 ± 0.063 (nighttime). A more
28 significant discrepancy between κ_{org} and κ_{HTDMA} was observed at night, whereas a larger
29 difference between κ_{OOA} and κ_{HTDMA} occurred during the day, which was likely due to the greatly

1 inhibited oxidation processes of organic compounds in the evening in contrast to the enhanced
2 situations during the day. This correlation could be visually observed in the corresponding
3 diurnal patterns of both mean GF and OA factors, as the fraction of OOA was demonstrated as a
4 moderately good indicator of the hygroscopicity of organics (Ogawa et al., 2016).

5 The observed variation in GF was predominantly caused by diurnal variations in the
6 probability distribution of g (Fig. 6b). Specifically, the number fractions of the highly
7 hygroscopic mode particles were low in the early morning and evening ($nf_{highly} < 0.3$), and higher
8 during the afternoon (approaching the highest level of 0.6 around 15:00 LT). The value was
9 positively correlated with GF ($R = 0.97$). The number fraction of the weakly hygroscopic
10 particles was opposite that of the highly hygroscopic particles, and it was negatively correlated
11 with the mean GF ($R = -0.95$). There was no clear diurnal variation in the number fraction of the
12 moderately hygroscopic mode particles (stable around 0.2). These results suggest that the water
13 uptake by wildfire haze particles is closely related to the fractions of weakly and more
14 hygroscopic mode particles. The mean g values for each mode were 1.05 ± 0.02 for weakly,
15 1.21 ± 0.01 for moderately, and 1.40 ± 0.05 for more hygroscopic particles. The average values
16 for nf were 0.42 ± 0.18 for weakly, 0.18 ± 0.07 for moderately, and 0.40 ± 0.20 for more
17 hygroscopic particles (Table 4).

18 The diurnal variation in hygroscopic properties coincided with that in chemical
19 compositions (Fig. 6c). The mean GF correlated well with $f_{SO_4^{2-}}$, suggesting the primary role of
20 $f_{SO_4^{2-}}$ in governing the water uptake by wildfire haze particles. The enhancement of $f_{SO_4^{2-}}$
21 accompanied decreased f_{org} . Variations in the chemical composition of organics also correlated
22 well with the water uptake properties. The fraction of signal intensity at m/z 44 (f_{44}), which is
23 considered as a marker ion for the degree of oxidation (Duplissy et al., 2011; Ng et al., 2011;
24 Ogawa et al., 2016), was also high during the day, which was similar to the case of the mean GF
25 and $f_{SO_4^{2-}}$. A similar pattern was also observed in f_{OOA} , whereas that in HOA was the opposite
26 (Fig. 6d). For instance, f_{HOA} was the highest during the morning rush hours, and it subsequently
27 decreased throughout the day. Variations in the fresh PBOA fraction (f_{PBOA}) were similar to those
28 of non-peat BBOA (f_{BBOA}) (Fig. 6e), namely, there was no apparent diurnal pattern during the
29 severe wildfire haze periods. Consequently, the distinctly enhanced mean GF during the day
30 could be attributed to the increase in both $f_{SO_4^{2-}}$ and f_{OOA} . Additionally, the local emission, which

1 primarily contributed to the HOA factor during morning/evening rush hours, could have
2 decreased the mean GF during these periods. The relative importance of both local emissions and
3 long-range transport might be influenced by meteorological conditions such as daily evolution of
4 planetary boundary layer (Krautstrunk et al., 2000; Geiß et al., 2017).

5 The diurnal variations in the organic composition were caused by the enhanced f_{OOA}
6 during the day, which accompanied decreases in f_{HOA} . In general, highly oxygenated organic
7 compounds are highly hygroscopic due to their water solubility, qualitatively explaining the
8 similarities in the diurnal variations among mean GF , f_{OOA} , and f_{44} (Duplissy et al., 2011; Zhao et
9 al., 2015; Ogawa et al., 2016). The relationship between the particle hygroscopicity and the
10 degree of oxidation of organics will be discussed in detail in Sect. 5.2.

11

12 **5. Discussion**

13 **5.1 Chemical composition dependences of water uptake by wildfire haze particles**

14 Figure 7 depicts the relationships between κ and the mass fractions of both inorganic and
15 organic species in NR-PM₁ in the data obtained during the day and at night. κ and $f_{SO_4^{2-}}$ were
16 positively correlated, demonstrating that sulfate was the most important compound in governing
17 water uptake by 100 nm wildfire haze particles due to its high hygroscopicity. Similarly, κ was
18 positively related to $f_{NH_4^+}$, as it is the counter cation of sulfate. In contrast, κ was negatively
19 correlated with f_{org} , as organics are less hygroscopic than inorganic salts are. There was no clear
20 correlation between κ and $f_{NO_3^-}$, implying that the small amount of nitrate had an insignificant
21 contribution to the variability in κ of wildfire haze particles. κ was almost independent of f_{Cl^-} ,
22 partially due to the limited availability of non-refractory chloride. These distinctly different
23 correlations between inorganics with mean κ likely reflect the different formation mechanisms of
24 these species. Sulfate formation occurs as a result of the photochemical process during the day,
25 whereas diurnal variations in gas–particle partitioning are important for the mass concentration
26 of nitrate or chloride (Aan de Brugh et al., 2012; Gong et al., 2013).

27 The correlation between κ and f_{org} was relatively scattered. For instance, κ varied from
28 0.10 to 0.40 when f_{org} was 0.7, signifying that factors other than f_{org} could influence the water

1 uptake. The variability in the chemical characteristics of organics might have played a role in the
2 scattering of the data (Fig. 6d-f).

3 **5.2 Relationship between the hygroscopicity and chemical composition of organics**

4 Table 2 summarizes the κ_{org} and κ_{OOA} results averaged over the same PM_{2.5} filter
5 sampling periods. Figure 8 illustrates the relationship between κ_{org} and f_{WSOC} . In general, κ_{org} was
6 insignificantly correlated with f_{WSOC} , especially in the data on October 22. The relatively high
7 value of f_{44} (0.11) on October 22 might have caused the deviation. These results indicate that the
8 oxygenation degree of organics could predominate over the WSOC fraction regarding the
9 hygroscopic growth of organic-rich wildfire haze particles.

10 Figure 9 depicts the relationships between κ_{org} and f_{44} as well as κ_{org} and f_{44}/f_{43} . The daily
11 average data were utilized. Although the data were scattered, a positive correlation between κ_{org}
12 and f_{44} was observed ($R = 0.70$). The signal of m/z 44 (mostly CO₂⁺) is known to originate in
13 highly oxidized organic functional groups, such as dicarboxylic acids and acyl peroxides (Aiken
14 et al., 2008). These highly oxygenated functional groups contribute to promoting hygroscopicity
15 by enhancing water solubility (Topping et al., 2005; Cubison et al., 2006; Hallquist et al., 2009;
16 Duplissy et al., 2011; Psichoudaki and Pandis, 2013; Suda et al., 2014; Riipinen et al., 2015;
17 Ogawa et al., 2016; Petters et al., 2016; Marsh et al., 2017). κ_{org} and f_{44}/f_{43} also presented a
18 similar trend to that of κ_{org} and f_{44} . The correlations shown in Figure 9 clearly demonstrate the
19 important role of oxygenation degree in the water uptake properties of organic compounds in
20 wildfire haze particles.

21 During the day, organic particles tend to be highly oxidized because of the oxidation of
22 primary organic aerosol as well as the formation of secondary organic aerosol from volatile
23 organic compounds (Ng et al., 2010 – see Fig. 6; Zhao et al., 2016). The chemical evolution
24 processes of organic aerosol particles need to be better understood to enhance quantitative
25 predictions of water uptake by wildfire haze particles (Kroll and Seinfeld, 2008; Riipinen et al.,
26 2011; Winkler et al., 2012; Ehn et al., 2014). The evolution process could induce alternations in
27 size dependence of chemical composition, and in the mixing state (Chakrabarty et al., 2006;
28 Zhao et al., 2015). To address these unanswered questions, further knowledge about size- and

1 mixing state-dependent chemical composition as well as the molecular-level chemical
2 characteristics of Indonesian wildfire haze particles are required.

3

4 **6. Conclusions**

5 In September–October 2015, Indonesian wildfire-induced transboundary haze pollution
6 spread throughout Southeast Asia, affecting both environment and climate dramatically and
7 ravaging public health and the economy seriously. As a downwind receptor city, Singapore
8 experienced pervasive wildfire haze events. In this study, we simultaneously measured the
9 hygroscopic growth factor (*GF*) and the chemical composition of ambient wildfire haze particles
10 in Singapore, with the aim of providing comprehensive insights into the linkages between water
11 uptake and particle chemical composition as well as secondary aerosol formation.

12 High aerosol loading of non-refractory submicron particles (NR-PM₁, occasionally
13 exceeding 100.0 μg m⁻³) was frequently observed, stressing the severity of the 2015 wildfire
14 haze pollution. The NR-PM₁ particles are predominantly composed of organics (OA,
15 approximately 77.1 % on average) and sulfate dominates the inorganic constituents (about
16 11.7 %). Chemical analyses of NR-PM₁ indicate the ubiquity and dominance of oxygenated
17 species in organics, in line with the most intense ion signals at *m/z* 44 (mostly CO₂⁺, *f*₄₄ = 10.5 %
18 in total OA mass) and *m/z* 43 (most likely C₂H₃O⁺, *f*₄₃ = 7.5 % on average). Moreover, a major
19 fraction of organics is water soluble, as signified by the high water-soluble organic carbon
20 fraction in ambient PM_{2.5} filter samples (26.7 % of the total PM_{2.5} mass).

21 Wildfire haze particles are generally highly hygroscopic. The hygroscopicity parameter, *κ*,
22 of 100 nm particles varied between 0.004 and 0.475, with a mean *κ* value of 0.189 ± 0.087. The
23 derived mean *κ* results of organics were 0.157 ± 0.108 (*κ*_{org}, bulk organics) and 0.266 ± 0.184
24 (*κ*_{OOA}, oxygenated organic fraction), which are comparable to previously reported values of
25 organic compounds (Petters and Kreidenweis, 2007; Duplissy et al., 2011; Lathem et al., 2013;
26 Cerully et al., 2015). These results highlight the difference in *κ* between wildfire haze particles
27 and fresh Indonesian peat burning particles, which are intrinsically non-hygroscopic due to the
28 rather limited water-soluble organic fraction (Chen et al., 2017). The *GF* data showed a notable

1 diurnal variation that usually peaked during the day. This variation was identical to the diurnal
2 pattern of the number fraction of more hygroscopic mode particles, and it accompanied the
3 opposite fluctuation of the number fraction of weakly hygroscopic mode particles. These results
4 imply the chemical composition dependence of particle hygroscopicity, as supported by the
5 positive correlation of κ with the mass fraction of sulfate but its inverse relation to the mass
6 fraction of organics. In addition to the governing influence of sulfate, κ of the haze particles was
7 promoted by the water uptake of organics. Although κ_{org} was loosely related to the water-soluble
8 organic fraction, a positive correlation between κ_{org} and f_{44} was shown ($R = 0.70$). This denotes
9 that the oxygenation degree of organics may play an important role in water uptake especially by
10 organic-rich haze particles.

11 To the best of our knowledge, this could be the first reported field measurements of water
12 uptake by wildfire haze particles in Equatorial Asia. The results suggest that formation of
13 secondary aerosol particles, including both inorganics and organics, is key in addressing the
14 variability of reported results about the hygroscopic properties of aerosol particles originating
15 from Indonesian peatland fires. Further quantitative studies on size-dependent chemical
16 composition and detailed chemical analyses at molecular levels are needed to deepen our
17 understanding of the water uptake by particles stemmed from Indonesian wildfires.

18

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23

1 **Table 1.** Summary of the hygroscopicity parameters (κ) and material densities of different
 2 chemical constituents utilized in the theoretical κ calculation with chemical data.

Chemical compounds	Hygroscopicity parameter, κ	Density (10^3 kg/m^3)
<i>SNA</i> [†]	0.59	1.77
<i>EC</i>	0	1.80
<i>PBOA</i>	0.04^ζ	1.10
<i>BBOA</i>	0.06^ζ	1.20
<i>OOA</i>	κ_{OOA}^*	1.50
Bulk OA	κ_{org}^*	1.40

3 [†] *SNA* includes all the sulfate, nitrate, and ammonium in submicron wildfire haze particles.

4 ^ζ κ values were assumed according to the mean κ results derived from laboratory studies (Chen et
 5 al., 2017).

6 * κ_{OOA} and κ_{org} were derived from ambient water uptake measurements and chemical data, in
 7 combination of the given parameters in Table 1, using the simplified algorithm introduced in
 8 Sects. 3.3 and 3.2.

1 **Table 2.** Summary of the averaged chemical characteristics of the 24 h $PM_{2.5}$ filter samples
 2 collected during 2015 haze events and the accordingly calculated mean κ results of organics
 3 ($RH = 85\%$).

Sampling date	Cl^- ($\mu g/m^3$)	NO_3^- ($\mu g/m^3$)	SO_4^{2-} ($\mu g/m^3$)	Na^+ ($\mu g/m^3$)	NH_4^+ ($\mu g/m^3$)	K^+ ($\mu g/m^3$)	Mg^{2+} ($\mu g/m^3$)	Ca^{2+} ($\mu g/m^3$)
Oct-14	0.64	2.40	10.21	0.20	4.93	0.62	0.05	0.55
Oct-15	0.38	1.72	10.51	0.18	4.63	0.60	0.03	0.50
Oct-16	0.27	1.62	9.01	0.25	4.03	0.48	0.08	0.65
Oct-19	0.65	2.39	15.90	0.20	7.99	0.58	0.03	0.41
Oct-20	0.24	1.09	10.22	0.31	4.33	0.51	0.05	0.48
Oct-21	0.19	0.83	7.95	0.23	3.15	0.51	0.08	0.52
Oct-22	0.23	0.80	9.71	0.29	3.78	0.38	0.08	0.50
Oct-23	0.56	2.65	15.92	0.18	9.13	0.70	0.005	0.25
Average	0.40	1.69	11.18	0.23	5.25	0.55	0.05	0.48

4 **Table 2. (continued)**

Sampling date	f_{inorg}^* (%)	$f_{SO_4^{2-}}$ (%)	f_{EC} (%)	$WSOC/OC$ (%)	Mean κ	Mean κ_{org}^\dagger ($D_0 = 100\text{ nm}$)	Mean κ_{OOA}^\dagger
Oct-14	42.1	21.9	15.3	79.9	0.114	0.030	0.054
Oct-15	41.1	23.3	11.1	64.8	0.202	0.118	0.185
Oct-16	28.6	15.7	11.7	66.4	0.161	0.133	0.256
Oct-19	26.6	15.0	5.9	60.2	0.265	0.229	0.450
Oct-20	30.3	18.0	11.3	61.5	0.271	0.236	0.443
Oct-21	26.4	15.6	11.0	56.9	0.193	0.148	0.283

Oct-22	42.9	26.4	15.8	59.5	0.265	0.212	0.376
Oct-23	24.6	13.3	4.4	59.6	0.217	0.184	0.337
Average	30.5	17.2	10.8	63.6	0.211	0.161	0.298

1 * The subscript *inorg* stands for all the inorganic species; hence, f_{inorg} is the mass fraction of
2 inorganic particles in the $PM_{2.5}$ filter sample. All the ionic data are provided with the mean mass
3 concentration ($\mu\text{g}/\text{m}^3$).

4 † The mean κ results of organics were calculated assuming 10.0 % elemental carbon (*EC*) in total
5 mass (see Sect.3.2).

6

1 **Table 3.** Derived mean κ values of organics with consideration of 10.0 % *EC* mass fraction in
 2 total NR-PM₁, as well as the mean κ results for HTDMA measurements conducted within the
 3 overlapped ToF-ACSM observation period of 10–24 October 2015 (i.e., κ_{HTDMA}).

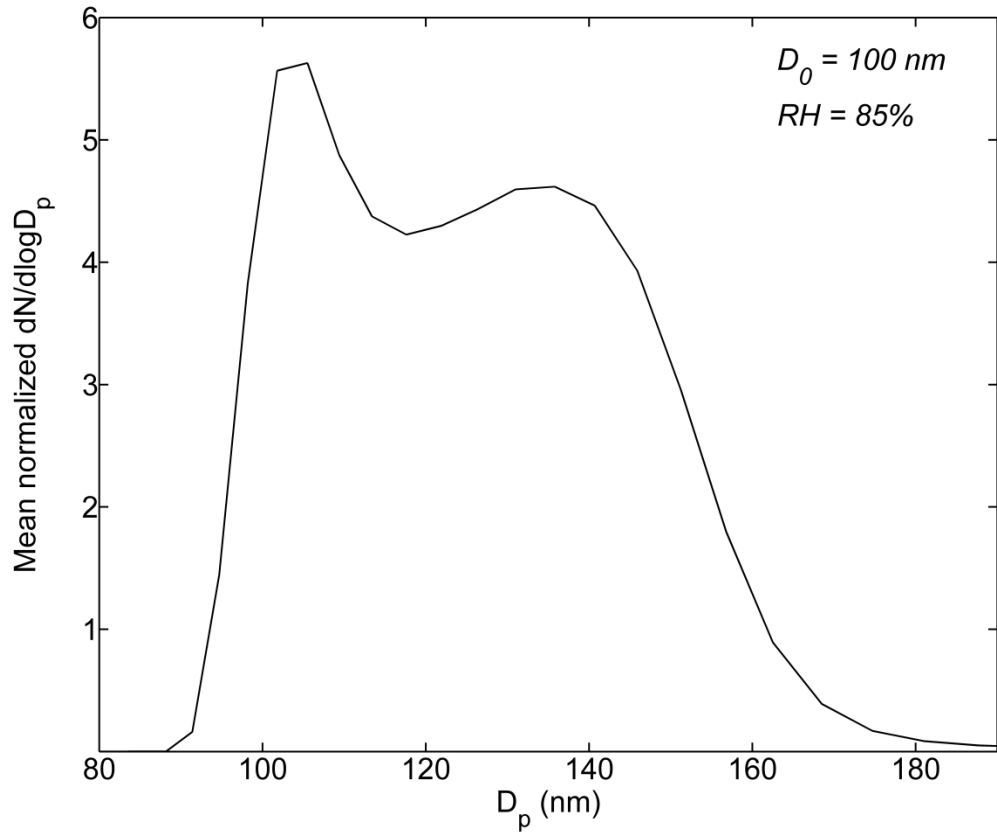
	Mean κ (mean \pm SD)		
	Overall	Daytime	Nighttime
<i>κ_{org}</i>	0.157 \pm 0.108	0.200 \pm 0.104	0.103 \pm 0.086
<i>κ_{OOA}</i>	0.266 \pm 0.184	0.323 \pm 0.177	0.186 \pm 0.177
<i>κ_{HTDMA}</i>	0.207 \pm 0.093	0.247 \pm 0.096	0.160 \pm 0.063

4

1 **Table 4.** The temporally mean number fraction (nf), volume-weighted mean diameter growth
 2 factor (GF), and κ results (mean \pm SD) of 100 nm wildfire haze particles in the three different
 3 hygroscopicity ranges at 85 % RH .

Hygroscopic mode	nf	GF	κ
Weakly ($g < 1.15$)	0.42 ± 0.18	1.05 ± 0.02	0.030 ± 0.013
Moderately ($1.15 \leq g < 1.27$)	0.18 ± 0.07	1.21 ± 0.01	0.151 ± 0.005
More ($1.27 \leq g < 1.85$)	0.40 ± 0.20	1.40 ± 0.05	0.343 ± 0.054
Bulk mean	<i>n/a</i>	1.25 ± 0.09	0.189 ± 0.087

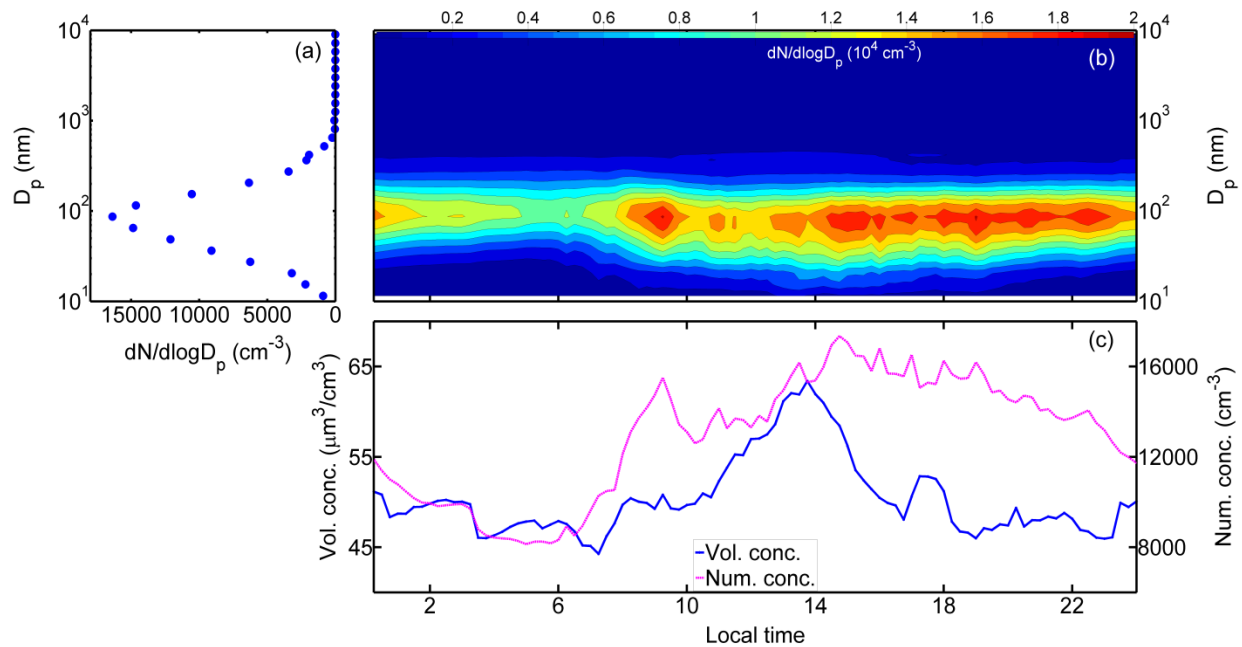
4 *n/a*: not applicable.



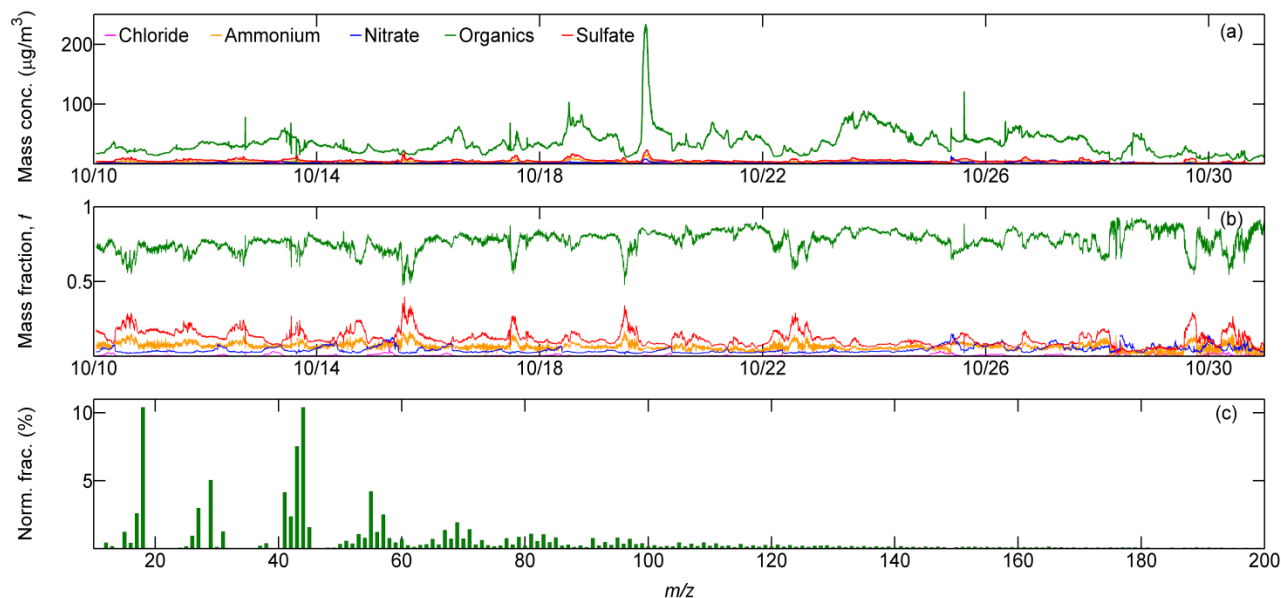
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2 **Fig. 1** Normalized particle number size distribution ($(dN/d\log D_p)/N$) after humidification
 3 averaged over the entire haze observation period.

4



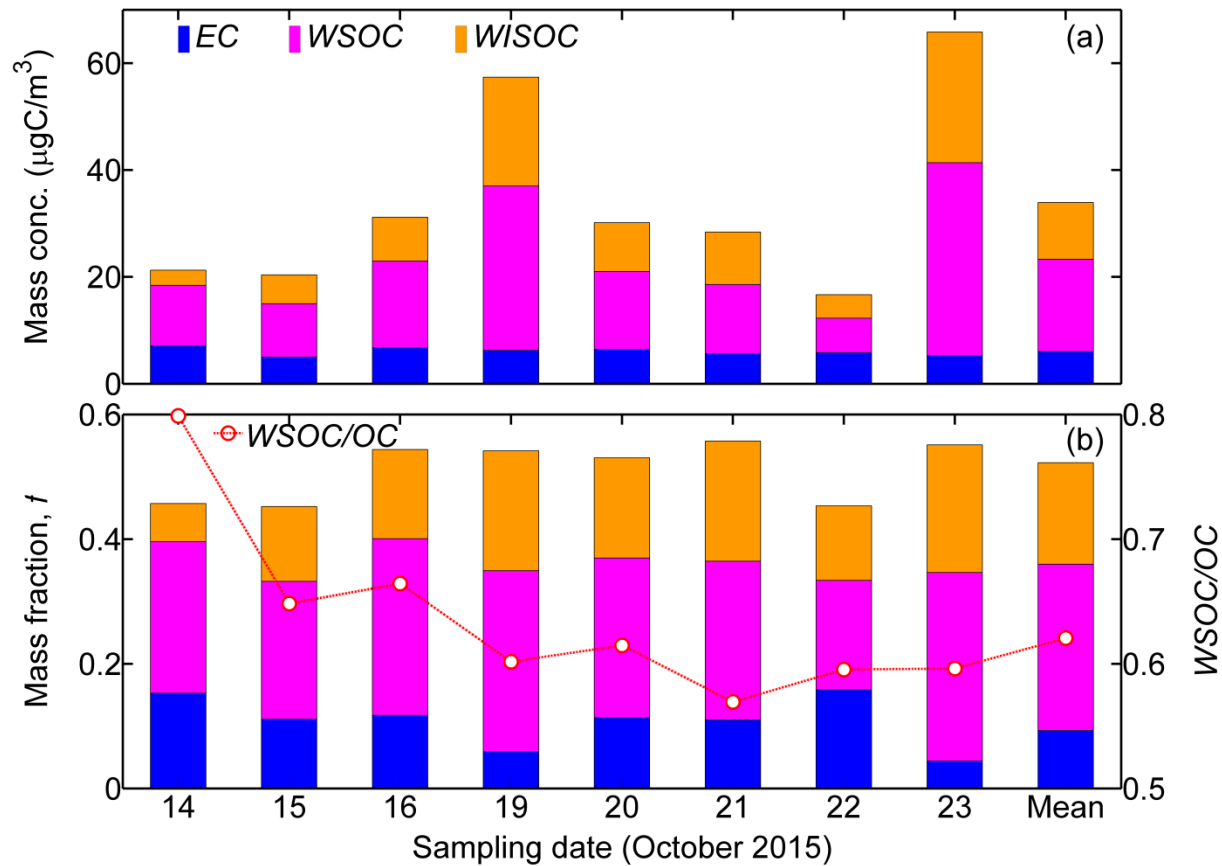
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 2 **Fig. 2** (a) The mean particle number size distribution ($dN/d\log D_p$, cm^{-3}) during the ambient
 3 wildfire haze observations as well as the mean diurnal variations of (b) particle size distribution,
 4 (c) number concentration (Num. conc., cm^{-3} , denoted by the magenta solid line), and volume
 5 concentration (Vol. conc., $\mu\text{m}^3/\text{cm}^3$, as the blue line displayed) measured with NanoScan-SMPS
 6 (11.5–365.2 nm) and OPS (419 nm–10 μm) (local time, LT).
 7 High aerosol loading was commonly observed during the transboundary wildfire haze.
 8 Submicron particles in the size range of 30–200 nm dominated the total number concentration of
 9 wildfire haze particles.



1
 2 **Fig. 3** Time series of (a) mass concentration (Mass conc., $\mu\text{g}/\text{m}^3$) and (b) corresponding mass
 3 fraction, f , of the five specific chemical species in NR-PM₁ measured by ToF-ACSM (abscissa
 4 shows the observation dates in October 2015 in the date format of month/day). (c) Temporally
 5 averaged OA mass spectra for submicron wildfire haze particles, displayed with the normalized
 6 ion fraction (Norm. frac., %) of each ion fragment.

7 Wildfire haze particles were predominantly composed of organics. Ion signals (m/z) from
 8 oxygenated organics (e.g., m/z 43, 44) were prominent, while intensities of ions for hydrocarbon-
 9 like organic compounds (e.g., m/z 41, 55, 57) and biomass burning tracers (e.g., m/z 60, 73) were
 10 relatively less intense. See the text for details.

11

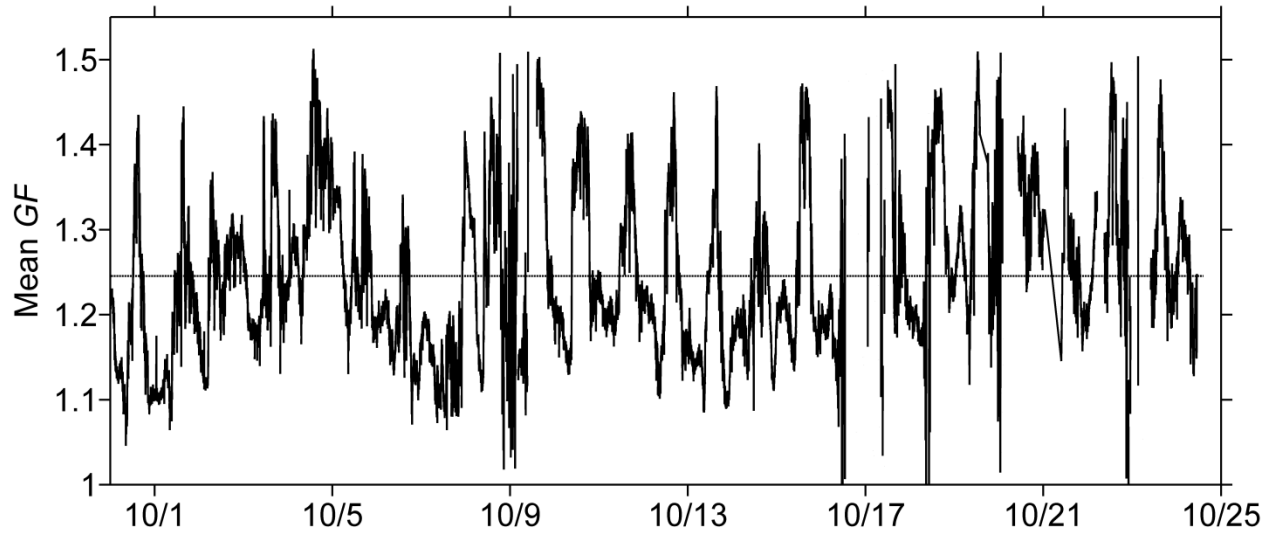


1

2 **Fig. 4** (a) Mass concentration (Mass conc., $\mu\text{gC}/\text{m}^3$) and (b) corresponding mass fraction, f , of
 3 the carbon contents including *EC*, *WSOC*, and water-insoluble *OC* (*WISOC*) in $PM_{2.5}$ filter
 4 samples. The *WSOC/OC* ratio is also displayed by the scattered dots in panel (b). All the
 5 corresponding temporal mean results are shown as “Mean”.

6 The *WSOC* fraction was exclusively higher than that of *WISOC* or *EC*, highlighting the
 7 dominance of *WSOC* in the carbon content of $PM_{2.5}$ filter samples. In general, the *EC* fraction
 8 fluctuated around 10.0 % of the total $PM_{2.5}$ mass.

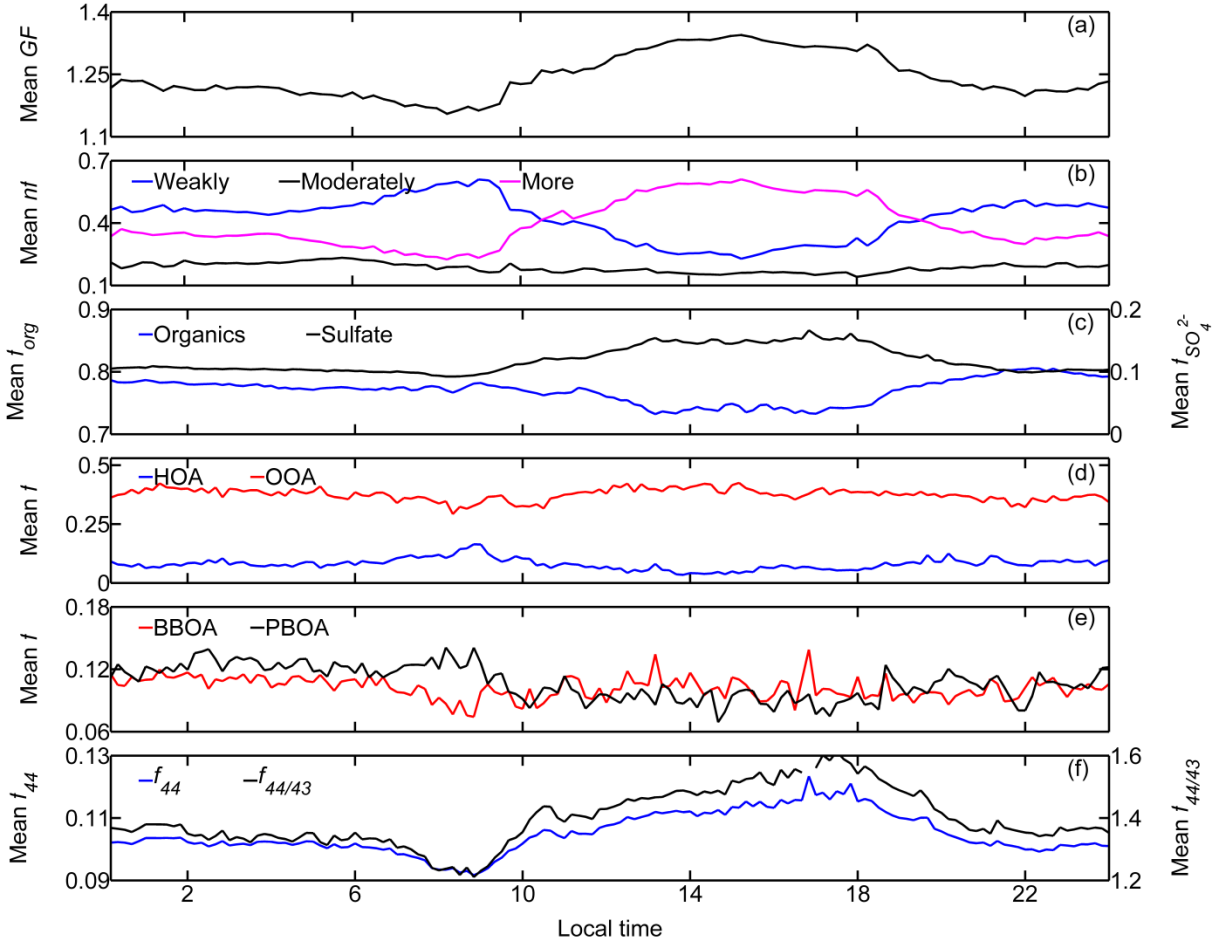
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2

3 **Fig. 5** Time series of the volume-weighted mean particle diameter growth factor (GF) derived
4 from HTDMA measurements (date format: month/day, 2015). The black dashed line stands for
5 the temporal mean GF averaged over the entire observation period.

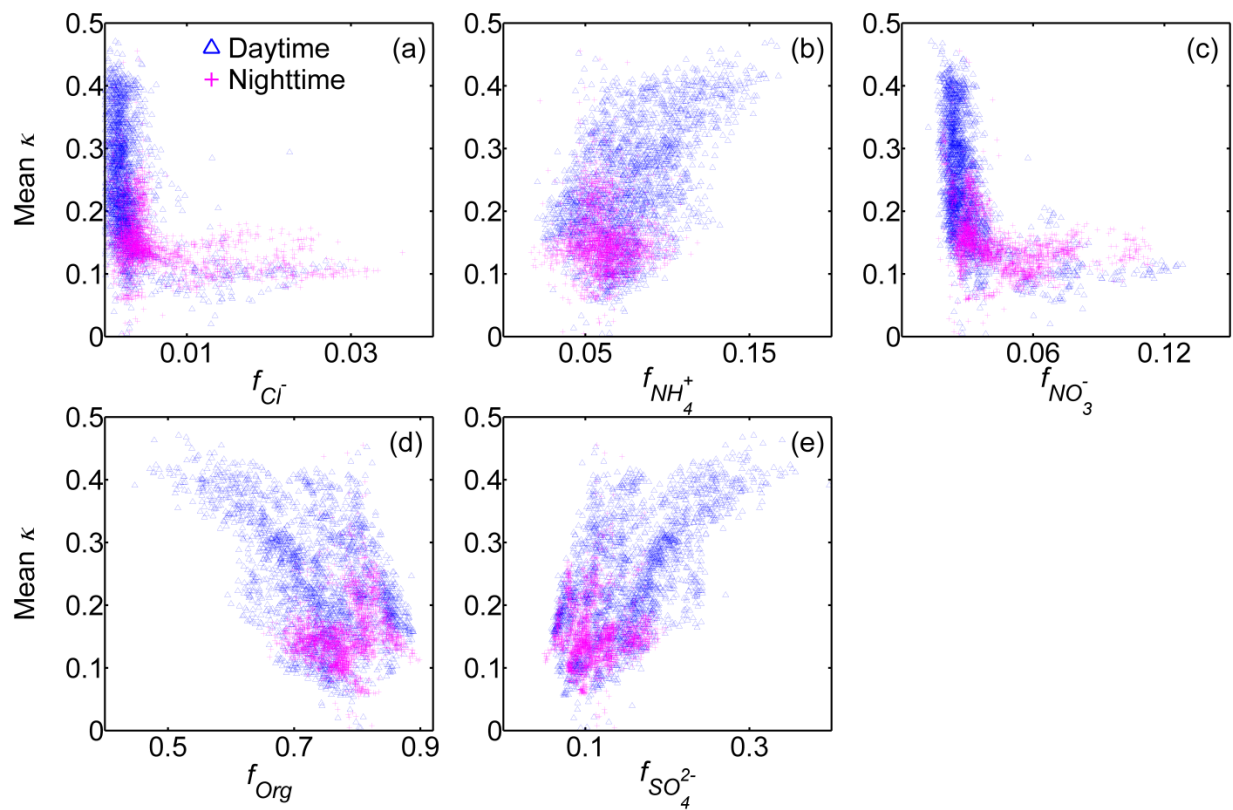
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1
 2 **Fig. 6** Temporal averaged (10–24 October 2015) diurnal variations of (a) the mean GF , (b)
 3 number fraction (nf) of the three hygroscopic modes, (c) mass fraction of the two main
 4 components in NR-PM₁, i.e., organics (f_{org}) and sulfate ($f_{SO_4^{2-}}$), (d-e) mass fraction of the ME2-
 5 resolved four OA factors in NR-PM₁ (i.e., f_{HOA} , f_{OOA} , f_{BBOA} , and f_{PBOA}), and (f) mean f_{44} and $f_{44/43}$
 6 of organics in NR-PM₁ (local time, LT).

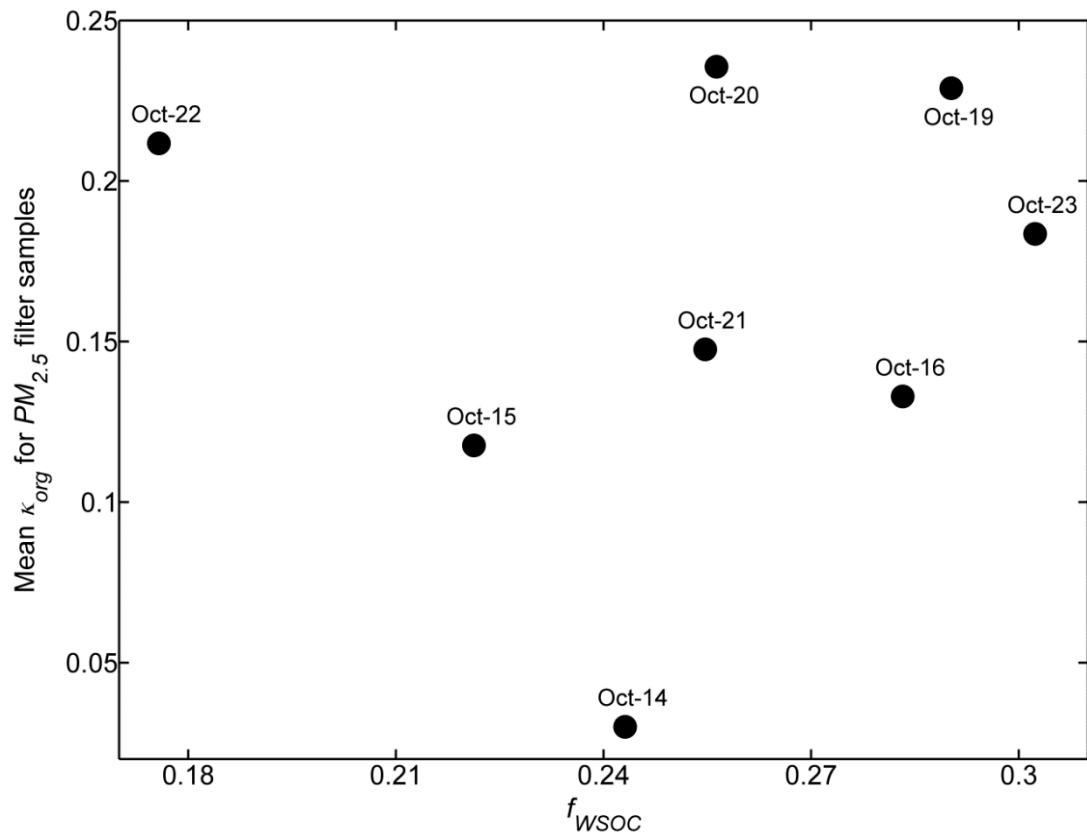
7 Pronounced diurnal patterns were observed for the mean GF , number fractions of both weakly
 8 and more hygroscopic modes, f_{44} , $f_{44/43}$, and mass fractions of organics and sulfate as well as
 9 HOA and OOA; nf_{more} , $f_{SO_4^{2-}}$, f_{44} , $f_{44/43}$, and f_{OOA} showed variations similar to that of the mean
 10 GF , whereas contrasting diurnal patterns were found for nf_{weakly} , f_{org} , and f_{HOA} .

11



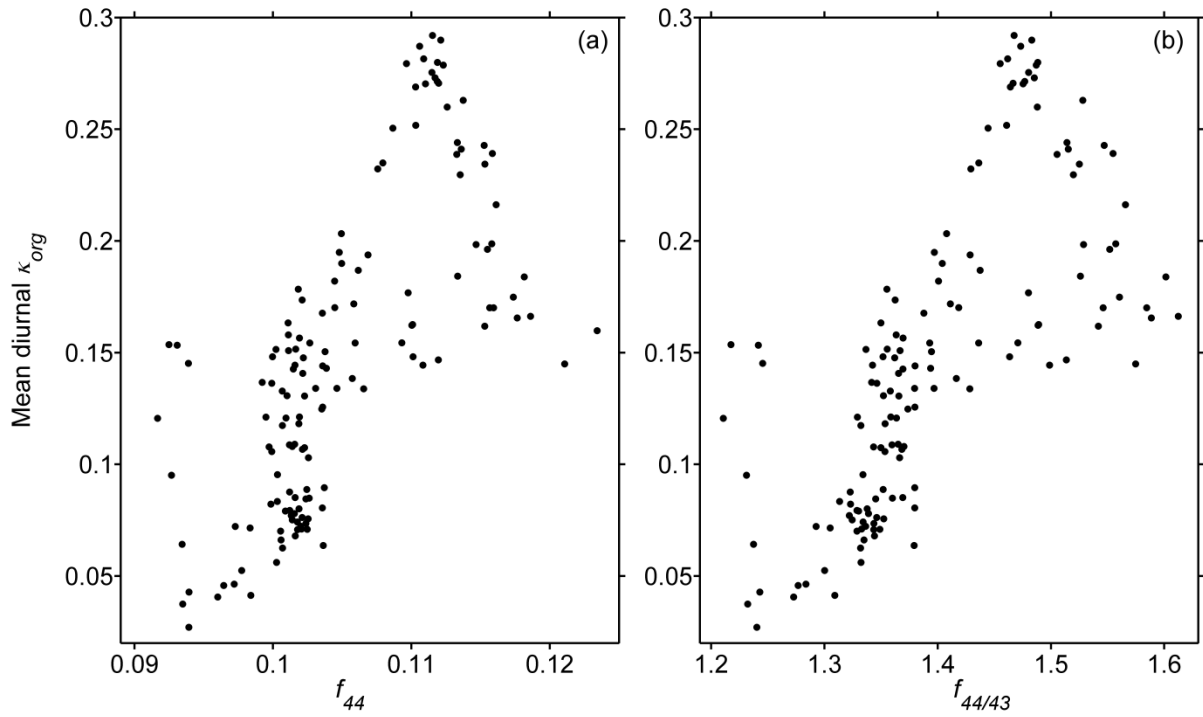
1
 2 **Fig. 7** Relationships between the mean κ results (100 nm) and the mass fractions of the five non-
 3 refractory chemical compositions of submicron wildfire haze particles.

4



1

2 **Fig. 8** Correlation between the mean κ of organics (κ_{org} , with 10.0 % *EC* mass fraction
 3 considered in the κ calculation) and the mean *WSOC* fraction (f_{WSOC}) of $PM_{2.5}$ filter samples.



1

2 **Fig. 9** Relationships between the mean diurnal κ_{org} results vs. (a) f_{44} and (b) $f_{44/43}$ in NR-PM₁
 3 haze particles. Mean diurnal here represents that the diurnal results for each day were further
 4 averaged over the overlapping observation period, i.e., with both HTDMA and ToF-ACSM
 5 measurements taken from 10–24 October 2015.