Multi-pollutants emissions from the burning of major
 agricultural residues in China and the related
 health-economic effect assessment

4 Chunlin Li¹, Yunjie Hu¹, Fei Zhang¹, Jianmin Chen^{1,2}, Zhen Ma¹, Xingnan Ye¹, Xin

5 Yang^{1,2}, Lin Wang^{1,2}, Xingfu Tang¹, Renhe Zhang², Mu Mu², Guihua Wang²,

6 Haidong Kan³, Xinming Wang⁴, Abdelwahid Mellouk i⁵

¹Shanghai Key Laboratory of Atmospheric Particle Pollution and Prevention (LAP³), Department

8 of Environmental Science & Engineering, Fudan University, Handan Road 220, Shanghai 200433,

9 China

²Institute of Atmospheric Sciences, Fudan University, Handan Road 220, Shanghai 200433, China

³Public Health School, Fudan University, Dongan Road 120, Shanghai 200032, China

⁴State Key Lab of Organ Geochemistry, Guangzhou Institute of Geochemistry, Chinese Academy

13 of Sciences, Kehuajie Road 511, Guangzhou 510640, China

⁵Institut de Combustion, Aérothermique, Réactivité et Environnement, CNRS, 45071 Orléans

15 cedex 02, France

16 Correspondence to: J. M. Chen (jmchen@fudan.edu.cn)

Abstract. Multi-pollutants in smoke particulate matter (SPM) were identified and 17 quantified for biomass burning of five major agricultural residues such as wheat, rice, 18 19 corn, cotton, and soybean straws in China by aerosol chamber system combining with various measurement techniques. The primary emission factors (EFs) for $PM_{1,0}$ 20 and PM_{2.5} are 3.04-12.64 and 3.25-15.16 g kg⁻¹. Organic carbon (OC), elemental 21 carbon (EC), water-soluble inorganics (WSI), water-soluble organic acids (WSOA), 22 water-soluble amine salts (WSA), trace mineral elements (THM), polycyclic 23 aromatic hydrocarbons (PAHs), and phenols in smoke $PM_{1.0}/PM_{2.5}$ are 24 1.34-6.04/1.54-7.42, 0.58-2.08/0.61-2.18, 0.51-3.52/0.52-3.81, 0.13-0.64/0.14-0.77, 25 $(4.39-85.72/4.51-104.79) \times 10^{-3}$, $(11.8-51.1/14.0-131.6) \times 10^{-3}$, $(1.1-4.0/1.8-8.3) \times 10^{-3}$, 26 and $(7.7-23.5/9.7-41.5) \times 10^{-3}$ g kg⁻¹, respectively. BC mainly exist in PM_{1.0}, heavy 27 metal-bearing particles favor to reside in the range of smoke $PM_{1,0-2.5}$, which are also 28

29 confirmed by individual particle analysis.

30 With respect to five scenarios of burning activities, the average emissions and overall propagation of uncertainties at 95% confidence interval (CI) of SPM from 31 agricultural open burning in China in 2012 were estimated for PM_{2.5}, PM_{1.0}, OC, EC, 32 WSI, WSOA, WSA, THM, PAHs, and phenols to be 1005.7 (-24.6%, 33.7%), 901.4 33 (-24.4%, 33.5%), 432.4 (-24.2%, 33.5%), 134.2 (-24.8%, 34.0%), 249.8 (-25.4%, 34 35 34.9%), 25.1 (-33.3%, 41.4%), 5.8 (-30.1%, 38.5%), 8.7 (-26.6%, 35.6%), 0.5 (-26.0%, 34.9%), and 2.7 (-26.1%, 35.1%) Gg, respectively. The emissions were 36 further temporal-spatially characterized using geographic information system (GIS) 37 at different regions in summer and autumn post-harvest periods. It was found less 38 than 25% of the total emissions were released during summer harvest that was 39 mainly contributed by the North Plain and the Central of China, especially Henan, 40 41 Shandong, and Anhui, leading the top three provinces of smoke particle emissions.

Flux concentrations of primarily emitted smoke PM_{2.5} that were calculated using 42 43 box-model method based on five versions of emission inventories all exceed the carcinogenic risk permissible exposure limits (PEL). The health impacts and 44 health-related economic losses from the smoke PM2.5 short-term exposure were 45 assessed. The results show that China suffered from 7836 (95% CI: 3232, 12362) 46 premature mortality and 7267237 (95% CI: 2961487, 1130784) chronic bronchitis in 47 2012, which led to 8822.4 (95% CI: 3574.4, 13034.2) million US\$, or 0.1% of the 48 total GDP losses. We suggest that percentage of open burnt crop straws at 49 post-harvest period should be cut down by over 97% to ensure risk aversion from 50 51 carcinogenicity, especially the North Plain and the Northeast, where the emissions should decease at least by 94% to meet the PEL. Under such emission control, over 52 92% of the mortality and morbidity attributed to agricultural fire smoke PM_{2.5} can be 53 54 avoided in China.

Key words: agricultural straw burning, aerosol chamber, smoke particle, emission
factor, emission inventory, health effect, emission control policy

57 **1 Introduction**

58 Biomass burning (BB) is a significant source of particulate- and gaseous- pollutants (Andreae and Merlet, 2001; Clarke et al., 2007; Ram et al., 2011; Saikawa et al., 59 2009a; Tian et al., 2008). It was estimated that open burning of biomass contributed 60 approximately 40% of the globally averaged annual submicron black carbon (BC) 61 aerosol emissions and 65% of primary OC emissions (Bond et al., 2013). China is 62 the major contributor that bears over 24% of global emissions of carbonaceous 63 aerosols, especially from agricultural field burning, about 0.04~0.5 Tg EC and 64 0.4~2.1 Tg OC are released annually (Bond, 2004; Cao et al., 2006; Qin and Xie, 65 2012; Saikawa et al., 2009), resulting in great radiative forcing, air quality 66 deterioration, visibility reduction, premature mortality, and economic loss regionally 67 and globally (Bølling et al., 2009; Bond et al., 2013; Huang et al., 2014; Janssen et 68 al., 2011; Rosenfeld, 2006; Saikawa et al., 2009; Shindell et al., 2012). 69

BB also represents one of the most uncertainties in the emission, climate effect, 70 and public health assessments, which finally relies on the uncertainties in detailed 71 chemical emissions or related properties and burning activities like strength or 72 73 percentage of biomass fuel burned (Tian et al., 2008; Andreae and Merlet, 2001; Levin et al., 2010). For example, studies have focused on OC and EC emissions due 74 75 to their specific optical properties (Bond et al., 2013; Cao et al., 2006; Qin and Xie, 2012; Ram et al., 2011). OC like sulfate and nitrate can cool the atmosphere by 76 increasing the Earth's reflectivity, however, smoke OC on the other hand has been 77 78 treated as brown carbon to exhibit pronounced light absorption character (Chen et al., 79 2015; Ackerman, 2000; Chakrabarty et al., 2010; Christopher et al., 2000). The coated or internal mixed sulfate or nitrate can act as lens to enhance the light 80 absorption activity of BC (Zhang et al., 2008b), probably also the activity of brown 81 carbon (Chen et al., 2015). However, primary emissions for OC, EC, and alkali 82 components are confused and have a wide range (Sen et al., 2014; Cao et al., 2006; 83 Hayashi et al., 2014), and some study still took OC with negative forcing activity 84 (Saikawa et al., 2009; Shindell et al., 2012). Besides, smoke EC is consisting of soot 85

86 and char, and soot-EC has a higher light-absorption potential compared to char-EC 87 (Arora and Jain, 2015; Reid et al., 2005a). Division and quantification of char- and soot-EC emissions for biomass burning are understudied (Arora and Jain, 2015; Han 88 et al., 2007, 2009). Moreover, other components like organic acids, amines, phenols, 89 and mineral elements that enable CCN activity or endow health hazard of smoke 90 aerosol are also deficient, variable, or outdated, which may hinder our overall 91 92 understanding of biomass burning contributions and also atmospheric process of smoke particles (Li et al., 2015; Akagi et al., 2011; Chan et al., 2005; Dhammapala 93 et al., 2007a; Ge et al., 2011; Reid et al., 2005a, b). 94

Studies using carbon mass-balance (CMB) and pollutant concentration-chamber 95 volume quantification are the two common methods to derive the emission factors 96 for biomass burning aerosols (Akagi et al., 2011; Li et al., 2007; Zhang et al., 2008a). 97 98 Carbonaceous and inorganics components of smoke particles not only vary with biomass issues (fuel types, water content, or burning strength), but also relate to 99 100 burning condition and environment (flaming or smoldering, field burning or laboratory simulation), extent of aging, sampling methods, and measurement 101 technologies (Grieshop et al., 2009; Hayashi et al., 2014; Reid et al., 2005b). 102 Comparing to field observations that are closer to the actual burning (Li et al., 2007; 103 Akagi et al., 2011; Rose et al., 2011; Saffari et al., 2013), laboratory studies have a 104 105 definite advantage over field burning research in emission analysis (Jayarathne et al., 2014; Sun et al., 2016; Zhang et al., 2008a). For example, the environment, amount 106 of fuel, and burning conditions can be precisely controlled, the contamination from 107 ambient atmosphere to the emissions can be excluded, and chemical compositions at 108 different aging extent can be quantified using aerosol chamber system (Li et al., 109 2015, 2016; Aurell et al., 2015; Dhammapala et al., 2007b). 110

The activity rates of biomass burning (burning rate of biomass fuels) are also response to the great uncertainties in the emission estimates (Sun et al., 2016; Zhang et al., 2008a). Seldom study ever focused on the burning rates, and the limited data were treated as simplex constant or dynamic values in many studies of emission estimation in a certain year or for annual variations with a long time scales, thus,

significant difference among the results were founded (Qin and Xie, 2011, 2012; 116 Zhang et al., 2011; Zhao et al., 2012). For instance, Cao et al. (2006, 2011) estimated 117 primary smoke carbonaceous materials emissions for 2000 and 2007 in China with 118 same field burning rates, the results were almost the same for the two year with 119 103-104 Gg yr⁻¹ BC and 425.9-433.3 Gg yr⁻¹ OC emitted. He et al. (2011b) found 120 the declining trends in biomass burning emissions in the Pearl River Delta for the 121 122 period 2003-2007 based on constant activity data of burning rates. Lu et al. (2011) developed primary carbonaceous aerosol emissions in China for 1996-2010 with 123 time-dependent activity rates extrapolated from 2008 to 2010 based on national 124 fast-track statistic, rapid increase of OC and EC emissions were reported, and OC 125 increased from 1.5 to 2.3 Tg yr⁻¹, BC increased from 418 to 619 Gg yr⁻¹. Qin and Xie 126 (2012) estimated BC emission from crop straw open burning for 1980-2009 with 127 variable burning rates based on peasants' income development, the increasing trend 128 in BC emission was also confirmed, and BC emission increased from 4.3 to 116.6 129 $Gg vr^{-1}$. 130

As most anthropogenic pollutants are concentrated in submicron particulate 131 matters ($PM_{1,0}$) (Ripoll et al., 2015), more pronounced relationship of ambient $PM_{1,0}$ 132 to haze formation and adverse health effect has been reported (Huang et al., 2003; 133 Roemer et al., 2001; Shi et al., 2014). Nevertheless, associated chemical 134 characterization of $PM_{1,0}$ is still undefined (Li et al., 2015; Safai et al., 2013; Cheng 135 et al., 2006). The study of source-specific $PM_{1,0}$ chemical compositions and 136 emissions are necessary to replenish database for contribution assessment and model 137 application in atmospheric chemistry, climate changes, and public health evaluation. 138

The emission inventories and forecasting in the emissions of atmospheric pollutants have been widely studied, and the incurred mortality, climatic effect, and economic loss have also been estimated (Ostro and Chestnut, 1998; Saikawa et al., 2009; Shindell et al., 2012), based on which the emission control policies were proposed. Shindell et al. (2012) considered ~400 control measures in tropospheric BC and O₃ emissions for the benefit of global or regional human health and food security, and 14 optimal measures targeting CH₄ and BC emissions were identified.

Saikawa et al. (2009) compared different scenarios of OC, EC, and sulfate emissions in China in 2030, concluding that maximum feasible reduction may avoid over 480000 premature deaths in China and decrease the radiative force from -97 to -15 mW m⁻² globally. Wang et al. (2008a) reported field burning restriction may save about 5 billion dollars losses from biological resource and air pollution. However, the generalized strategies in emission reduction were inadequate and lack actual practicality (Streets, 2007; Lin et al., 2010).

In this study, burning experiments with five major agricultural straws were 153 conducted using a combustion stove in combination with an aerosol chamber system. 154 Accurate compositions and emission factors for SPM in PM_{1.0} and PM_{2.5} were 155 characterized and established. Afterwards, up-to-date emissions for agricultural open 156 burning aerosol in 2012 were developed, health and health-related economic impacts 157 158 from smoke PM_{2.5} exposure were also assessed. Finally, emission reduction strategy that was implemented in field burning rate control for the carcinogenic risk concern 159 160 was proposed, which should help establish the policy and provide an idea for the emission control. 161

162 **2 Methodology**

An overview of the research procedures including emission factors acquirement and emission inventory calculation is shown in Fig. 1. Tabulation of emission factors is self-established in our laboratory using a combustion stove to simulate open burning and an aerosol chamber to quantify the emissions. Then, we use a bottom-up approach to calculate the emission inventory of agricultural field burning over China mainland based on crop production data in 2012. Emissions for each species are estimated as:

170
$$E_{k,j} = \sum_{i} A_{k,i} \times EF_{i,j}$$
(1)

where E_j is emission, $A_{k,i}$ is effective biofuel consumption, and $EF_{i,j}$ is emission factor. k, i, and j indicates region, agricultural residue type, and particulate chemical species. 174 State-of-the art chemical transport and box models were commonly applied to reproduce or simulate the ambient aerosol concentrations (Ram et al., 2011; Reddy 175 and Venkataraman, 2000; Saikawa et al., 2009). In this study, spatio-temporal 176 dynamic box model is used to calculate the emission flux concentration. Regional 177 crop straws are premised to be combusted proportionally only in the fire occurrence 178 days. Dismissing interaction of emitted pollutants in space and time, pollutants will 179 180 distribute uniformly in a space covering an area of specific region with mixing height of 0.5 km (atmospheric boundary layer). The flux concentration of 181 agricultural burning smoke can be calculated by Eq. (2): 182

183
$$C_{k,j} = \frac{E_{k,j}}{S_k \times h \times T_k}$$
(2)

in Eq. (2), $C_{k,j}$ is flux concentration of smoke aerosol, S_k is regional area, h is boundary layer height, T_k is agricultural field fire duration time.

186 2.1 Aerosol chamber work and emission factors

187 2.1.1 Crop straws

Five kinds of representative crop residues were used for the burning experiments, i.e., 188 wheat, rice, corn, cotton, and soybean straws. The straws were collected based on 189 regional features of agricultural planting, winter wheat straws were collected from 190 Anhui province, late rice straws from Shanghai, corn straws from Henan province, 191 cotton and soybean residues from Xinjiang. All straws were stored under dark, airy, 192 and cooling condition. Prior to the burning experiment, the dirt and weeds were 193 removed, then straws were dehydrated (at 100 °C for 24 h) to minimize effect of the 194 195 water content on the burning and pollutant emissions, as study found pollutants emissions and combustion efficiencies (CE) are response to water content, increased 196 197 moisture content enhances the emissions but also alter the chemical compositions of 198 smoke aerosols (Reid et al., 2005b; Aurell et al., 2015; Hayashi et al., 2014). Although straws in the field are not well dried and moisture contents vary with 199 weather, ventilation, and storing times, for the convenience of practical application 200

and comparison of burnings and emissions, water contents of the straws were
controlled within 2 wt.%, which has been applied in many studies (Hayashi et al.,
2014; Huo et al., 2016; Li et al., 2015; Oanh et al., 2011; Zhang et al., 2008a, 2011).
The dry straws were then cut to a length of approximately 10 cm and weighted 10.0
g per serving.

206 2.1.2 Burning experiments

The experiments were conducted using an aerosol chamber system (Fig. S1 in 207 supplement information, SI), which was loaded in a temperature-controlled room 208 (18-22 °C, 40%-60% RH). A stainless combustion stove was self-deigned to simulate 209 typical field burning of crop straws by automatic ignition with LPG (Liquid 210 petroleum gas) in particular, albeit on a small scale (ignition time less than 0.1 s). 211 10.0 g conditioned residues were sealed in the 0.227 m³ combustion stove in 212 advance, once ignited, the force-ventilation and HEPA filtrated particle-free air were 213 supplied (300 L min⁻¹). The emissions were immediately injected into a clean, 214 evacuated aerosol chamber. The burning last about 1 min and over 1 m³ particle-free 215 air flushed the stove to ensure all the emissions were transferred into the chamber. 216

The chamber was custom-built to quantify the emissions and characterize the 217 physiochemical properties of smoke aerosols, detailed description of the chamber 218 can be found elsewhere (Zhang et al., 2008a, 2011; Li et al., 2015, 2016). Briefly, the 219 220 chamber has a volume of 4.5 m³ with 0.3 mm Teflon coating on the inner side, a magnetic fan fixed on the bottom to stir the aerosol uniformly, and a hygroclip 221 222 monitor (Rotronic, Model IM-4) equipped inside the chamber to measure the temperature and relative humidity of the aerosol. Before experiment, the chamber 223 was flushed with particle-free air for 6 h, oxidized by high concentration ozone (~3 224 ppm) for 12 h, then flushed and vacuumized, filled with pure dry air to 80 KPa for 225 use. The emissions from straw burning were aspirated into the chamber till room 226 227 pressure, afterwards, size measurement and sampling of smoke aerosols were 228 conducted from the chamber. For each type of straw, four burning experiments were

conducted. The unburned residues were weighted and deducted from 10.0 g aftereach test.

231 Modified combustion efficiency (MCE) for each burning was monitored with CO and CO₂ measuring to determine the burning phase and to ensure the repeatability. 232 MCE is defined as $\Delta CO_2/(\Delta CO_2 + \Delta CO)$, where ΔCO_2 and ΔCO are the excess molar 233 mixing ratios of CO₂ and CO (Reid et al., 2005b). A gas-chromatograph (GC, model 234 235 930, Shanghai, Hai Xin Gas Chromatograph Co., LTD) equipped with a flame ionization detector, an Ni-H convertor, and a stainless steel column (2 m long) 236 packed with 15% DNP was used to measure CO and CO₂ concentrations in the 237 chamber. And MCE were 0.89-0.96 for all the experiments (see in SI, Table S1), 238 indicating flaming combustion dominated, which were comparable to that in the 239 field burning (Li et al., 2003; Li et al., 2007). 240

241 **2.1.3 Size and morphology of smoke aerosol**

Size distribution (10 nm-10 µm) of smoke particles was measured using a 242 Wide-range Particle Spectrometer (WPS, Model 1000XP, TSI, USA), which has 243 been described by Li et al (2015). Briefly, WPS integrates the function of scan 244 mobility particle sizer (SMPS) and laser particle sizer (LPS), 0.3 L min⁻¹ flow is 245 introduced to SMPS part to classify mobility size from 10 nm to 500 nm in 48 bins, 246 and 0.7 L min⁻¹ flow is introduced to LPS part to measure aerodynamic diameter 247 from 350 nm to 10 µm in 18 bins. Particle density, refractive index, and scanning 248 time were set as 1.0 g cm⁻³, 1.45, and 3 min $loop^{-1}$, respectively, and charge 249 correction mode was on for the measurement. A diffusion dryer tube (45 cm in 250 length) filled with descant-silica gel was set prior to the inlet of WPS. Before 251 experiment, WPS was calibrated with certified polystyrene latex spheres (PSL, 40, 252 80, and 220 nm, Duke Scientific). 253

SPM from the 5 types crop straws burning were sampled onto copper grids coated with carbon film (carbon type-B, 300-mesh copper, Tianld Co., China) using a single-stage cascade impactor with a 0.5 mm diameter jet nozzle at a flow rate of 1.0

L min⁻¹. The sampler has a collection efficiency of 100% at 0.5 μ m aerodynamic diameter. More information about the cascade impactor can be found elsewhere (Fu et al., 2012; Hu et al., 2015). Then, a JEOL-2010F field emission high-resolution transmission electron microscope (FE-HRTEM) coupled with an oxford energy-dispersive X-ray spectrum (EDX) was applied to investigate the morphology, composition, and mixing state of individual particles.

263 2.1.4 Chemical sampling and analysis

PM_{1.0} and PM_{2.5} samples for each burning were collected onto pretreated quartz filter 264 of 90 mm in diameter (Tissuquartz, Pall Corp., USA) from the chamber using a 265 high-volume Particle Sampler (HY-100, Qingdao Hengyuan S.T. Development Co., 266 Ltd) operating at 100 L min⁻¹. Each filter sampling duration time is 5 min, and total 267 44 samples (including 4 blank samples) were gathered. The quartz microfiber filters 268 were prebaked for 8 h at 450 °C to eliminate contamination. Before and after the 269 sampling, the filters were weighted using a balance (Sartorius BP211D) with an 270 accuracy of 10 µg, and the filters were balanced in an electronic desiccator (40 % 271 RH, 22 °C) for 24 h before usage. After weighting, the loaded filters were stored at 272 -20 °C in a refrigerator for further analysis. 273

Water soluble species including general inorganic ions (ions: F^- , $C\Gamma$, NO_2^- , NO_3^- , 274 SO4²⁻, Na⁺, NH4⁺, K⁺, Ca²⁺, Mg²⁺), organic acids (CH₃COOH, HCOOH, C₂H₂O₄, 275 CH₃SO₃H), and seven protonated amines (MeOH⁺, TeOH⁺, MMAH⁺, DMAH⁺, 276 TMAH⁺, MEAH⁺, and DEAH⁺ for short, corresponding to monoethanolaminium, 277 monomethylaminium, 278 triethanolaminium, dimethylaminium, triethylaminium, monoethylaminium, and diethylaminium) were measured from 1/4 of each filter 279 with ion chromatography (IC, Model 850 Professional IC, Metrohm, USA) consists 280 of a separation column (Metrosep A Supp 7 250/4.0 for anion and organic acids, 281 Metrosep C-4 150/4.0 for cation, and Metrosep C4-250/4.0 for water soluble 282 283 aminiums). Sampled filters were ultrasonically extracted with 15.0 mL deionized water (Mili-Q water, 18.2 MΩ cm), extracted solutions were filtrated using 0.2 μm 284

filters before injected into IC for measurement. Detection limits (DLs) for the ions and aminiums were within $0.5 \sim 3.5$ ng mL⁻¹, the correlation coefficients for all calibration curves were better than 0.99, and recovery rates for aminiums were in the range of 93%~106% (see in SI, Table S2). Details for the aminium measurements can be found in the work of Tao et al. (2016).

1/4 of each filter was acid dissolved to measure the selected elements (As, Pb, Cr, 290 291 Cd, Ni, V, Zn, Al), of which As, Zn, Pb, Cr, Cd, and Ni are USEPA priority controlled pollutants (Wu et al., 2011). The smashed filters were digested at 170 °C 292 for 4 h in high-pressure Teflon digestion vessel with 3.0 mL concentrated HNO₃, 1.0 293 mL concentrated HClO₄, and 1.0 mL concentrated HF. Afterwards, the almost dry 294 solution was diluted and characterized using Inductively Coupled Plasma Optical 295 Emission Spectrometer (ICP-OES, Atom Scan 2000, JarroU-Ash, USA). The 296 297 following wavelength lines of the ICP-OES analysis were used: As 189.042, Pb 220.353, Cd 228.802, Cr 205552, Ni 231.604, V 311.071, Zn 206.191, and Al 298 299 394.401. All reagents used were of highest grades, and recovery tests were conducted with standard additions, recoveries of each element were in the range of 300 93%~102% (see in SI, Table S2). 301

Another 1/4 of each filter was ultrasonically double extracted with 15.0 mL 302 HPLC-grade CH₂Cl₂. The extracts were then condensed with rotary evaporator and 303 quantified to 1.0 mL. 16 targeted PAHs (2-ring, naphthalene (Nap); 3-ring, 304 acenaphthylene (Ac), acenaphthene (Ace), fluorene (Fl), phenanthrene (Phe), 305 anthracene (Ant); 4-ring, fluoranthene (Flu), pyrene (Pyr), benzo[a]anthracene (BaA), 306 chrysene (Chr); 5-ring, benzo[b]fluoranthene (BbF), benzo[k]fluoranthene (BkF), 307 benzo[a]pyrene (BaP), dibenzo[a,h]anthracene (DBA); and 6-ring: indeno[1,2,3-cd] 308 pyrene (IP), benzo[ghi]perylene (BghiP)) and 5 selected phenols (phenol, 309 2-methoxyphenol, 4-ethylphenol, 4-ethyl-2-methoxyphenol, 2,6-dimethoxyphenol) 310 311 were measured from the concentrated extracts using an Agilent 6890 Series gas chromatography system coupled with a HP 5973 Mass Selective Detector (GC-MS, 312 Agilent Technologies, Wilmington DE). A DB-5ms (30 m $\times 0.32$ mm $\times 0.25$ mm, 313 Agilent 123-5532) column was installed. The temperature programs were presented 314

as follows: initially at 40 °C, hold for 4 min, to 150 °C at 20 °C min⁻¹, then to 280 °C 315 at 5 °C min⁻¹, hold for 10 min. The interface temperature was kept at 280 °C, the MS 316 was operated in electron impact mode with an ion source temperature of 230 °C, and 317 the high-purity helium (99.999%) carrier gas was maintained at a constant pressure 318 of 16.2 psi with a flow of 2.0 mL min⁻¹. The calibration curves were optimized to be 319 better than 99.9%. Prior to the measurements, PAHs and Phenols recovery studies 320 321 were undertaken, and recoveries were acceptable with rates of 82%~99% (see in SI, Table S2). In addition, Phenanthrene-d10 (Phe-d10) as internal standard surrogate 322 was added into the PAHs mixture, recovery rate of which was 94%. 323

Organic carbon (OC) and elemental carbon (EC) were measured with the rest 324 quartz filters using a carbon analyzer (Sunset laboratory Inc., Forest Grove, OR) 325 based on the thermal-optical transmittance (TOT) method with a modified 326 327 NIOSH-5040 (National Institute of Occupational Safety and Health) protocol. Four organic fractions (OC1, OC2, OC3, and OC4 at 150, 250, 450, and 550 °C, 328 329 respectively), PC fraction (a pyrolyzed carbonaceous component determined when transmitted laser returned to its original intensity after the sample was exposed to 330 oxygen), and three EC fractions (EC1, EC2, and EC3 at 550, 700, and 800 °C, 331 respectively) are produced. And OC is technically defined as OC1 + OC2 + OC3 + OC3332 OC4 + PC, while EC is defined as EC1 + EC2 + EC3 - PC (Seinfeld et al., 2012). 333 The instrument detection limits for total OC and EC that deposit on the filter are 0.25 334 and 0.12 μ g C cm⁻². The quality of the data above was guaranteed by standard 335 materials calibration, recovery rate, and operational blank correction. And blank 336 levels were less than 5% of the measured values for all the species. 337

338 **2.1.5 Calculation of emission factors**

The emission quantities derived from the experiment were converted into quantities per unit weight of initial residues as emission factor (EF, unit: g kg-1), which can be calculated from the direct method with effective filter sampling weight, chamber volume, and effective amount of crop straw consumed (Dhammapala et al., 2006, 343 2007a, b; Zhang et al., 2008a), or alternatively from the carbon mass balance method (CMB) via conservation of Carbon in biomass, disregarding the weight of biomass 344 345 that burnt (Dhammapala et al., 2006; Li et al., 2007). EFs determined from these two methods were found to be in good agreement (Dhammapala et al., 2006), 346 nevertheless, CMB method needs more auxiliary information (e.g., concentrations of 347 CO, CO₂, CH₄, non-methane hydrocarbons, and also particulate carbons), which 348 349 may result in data redundancy and uncertainty propagation, hence we applied the direct method to calculate EFs in this work. To be more accurate, influence of wall 350 loss and makeup air dilution on smoke particles sampling from the chamber were 351 considered and corrected, details see in SI. 352

In this study, duration for each test (burning, chamber condition, size measurement, and filter sampling) was controlled within 20 min, therefore, the physicochemical processes of pollutants in the chamber can be negligible, and smoke aerosols we measured were primary emissions.

357 **2.2 Emission inventory calculation**

358 2.2.1 Agricultural field fire survey

Fire sites over China from 2011 to 2013 were statistically analyzed, and the data of 359 mainland agricultural fire sites was derived from the daily report of the Ministry of 360 Environmental Protection of China (MEPC) (website: http://hij.mep.gov.cn/jgjs/). 361 Agricultural fire sites were screened out from MODIS (Moderate Resolution 362 Imaging Spectroradiometer) daily fire products (1 km \times 1 km resolution level 3 363 hotspot) using a high resolution real time land use based on geography information 364 system (GIS). Spatial and temporal distributions of fire sites were displayed in Fig. 365 S2 (SI), over 5000 fire sites were allocated into two prominent burning periods 366 corresponding to summer (May to July) and autumn (September to November) 367 368 harvests, and filed burning last 54 days and 60 days on statistical average during the two harvests. In the North of China, open burning occurred primarily in autumn, 369 while temporal-character of field fires was not obvious in the North Plain and the 370

371 Center of China, where field fires can be observed frequently during the whole372 investigation time.

373 2.2.2 Crop straw production

Crop straw production was generally derived from annul or monthly crop production by multiplying crop-specific ratios of residue-to-production (He et al., 2011b; Cao et al., 2011; Zhao et al., 2012). In this study, crop productions were furtherly classified into summer harvest and autumn harvest productions according to field fire sites analysis and traditional seasonal planting and harvesting. The amount of straw produced was calculated by Eq. (3):

380
$$M_{tki} = P_{tki} \times r_i \times H_{tki} \times D_i$$

(3)

in which M is mass of crop straws produced; P is annual crop-specific amount of crop production; r is the residue-to-production ratio; D is the dry matter content; $H_{t,k,i}$ is production ratio of crop i at region k during summer or autumn harvest period t.

385 Province-level crop production data of wheat, rice, corn, cotton, and soybean were taken directly from the China Yearbook 2013 (National Bureau of Statistics of China, 386 NBSC, 2013). Crop-specific residue-to-production ratios were cited from Chinese 387 Association of Rural Energy Industry (Wang and Zhang, 2008; data available at 388 389 http://www.carei.org.cn/index.php, in Chinese). Dry matter contents of crop straws were referred to He et al. (2011b) and Greenhouse Gas Inventory Reference Manual 390 (IPCC, 2007). The parameters of residue-to-production ratios and dry matter 391 392 contents were summarized in Table S3 (SI). The regional crop production ratios in summer and autumn harvests were listed in Table S4 (SI). 393

394 2.2.3 Field burning rate

Uncertainty of emission estimations mostly relies on intangibility of straw open burning rate (Zhao et al., 2012; He et al., 2011b). However, regional or national percentage of straw open burned was seldom studied, and the limited data were 398 outdated and variable. The available studies indicate national filed burning rate of crop straws range from 15.2% to 27.2% in China (Daize, 2000; Wei et al., 2004; 399 400 Zhang et al., 2008a), and more detailed studies indicate about 31.9% of the crop straw burned in the Pearl River Delta from 2003 to 2007 (He et al., 2011b), while the 401 corresponding figures were almost 100% for the Huabei region in 2003 (Zhao et al., 402 2012). Two versions of province-level field burning rates were commonly used, one 403 404 was from Cao et al. (2005; 2006; 2011) who deduced the rates based on regional economic level, and the proposal of the rates to be proportional to peasants' income 405 was confirmed later, the rates were first used to calculate open burning emission in 406 2000. The other version was reported by Wang and Zhang (2008), they obtained 407 provincial percentage of residue open burnt via filed survey in 2006. Herein, the two 408 versions were both applied directly into the emission estimation of 2012 in this work 409 410 and named as business-as-usual scenarios (BAU, BAU-I from Cao et al. and BAU-II from Wang and Zhang in specific). 411

412 In fact, the burning rates should be dynamic parameters that been influenced by industrial structure, government policy orientation, or public awareness. With crop 413 yields increase and energy consumption structure changes in rural areas, more straws 414 will be discarded and burnt in the field. Nonetheless, rigorous agricultural fire policy 415 may still suppress the condition worsen as it worked during 2008 for Beijing 416 Olympics and 2010 for Shanghai Expo (Huang et al., 2013; Cermak and Knutti, 417 2009; Wang et al., 2010). Qin and Xie (2011; 2012) ever deduced year specific open 418 burning rates in different zone for the period of 1980-2009 according to their 419 respective peasant income changes in a certain year on the basis of peasant income 420 and burning rates in 2006. However, the simple linear relationship should be doubted, 421 as great increase in per capita income after 2006 will surely overestimate the burning 422 423 rates. We supposed that the burning rates were inverse proportional to peasants' 424 agricultural income proportion (AIP), without considering the policy or potential 425 gain or loss related to agricultural residue treatment. Thus the burning rates established in 2000 and 2006 from Cao et al. (2005) and Wang and Zhang (2008) can 426 427 be converted into that of 2012 based on economic data from equation below:

428
$$R_{k,2012} = \frac{I_{k,2012}}{AI_{k,2012}} \times \frac{AI_{k,y}}{I_{k,y}} \times R_{k,y}$$
 (4)

where R is agricultural straw filed burnt rate, $I_{k,y}$ is peasants' annual income, $AI_{k,y}$ is peasants' annual agricultural income. y indicates reference year (2000 for BAU-I, and 2006 for BAU-II). $I_{k,y}$ and $AI_{k,y}$ can be found or calculated from China Yearbook and China Rural Statistic Yearbook (NBSC, 2004-2013).

The versions of converted rates based on primary industry level were called Economic Models I and II (EM -I and EM-II in short) corresponding to BAU-I and BAU-II. Besides, in 2013, the National Development and Reform Commission of China published the Chinese agricultural straw treatment report of 2012 (NDRC, [2014] No.516, data available at http://www.sdpc.gov.cn/, in Chinese) for the first time. The percentages of crop residues discarded in the report were applied in our estimation, which was called NDRC version.

440 **2.2.4 Emission and flux concentration**

441 From above study, emission of SPM pollutants can be calculated by recount of Eq.

(5)

443
$$E_{t,k,j} = \sum_{i} M_{t,k,i} \times R_k \times f_i \times EF_{i,j}$$

where $E_{t,k,j}$ is emission amount of chemical species j at region k during harvest period t; f_i is burning efficiency, the crop specific values were cited as 0.68 for soybean residue and 0.93 for the rest four straws (Zhang et al., 2011; Wang and Zhang, 2008; Zhang et al., 2008a; Koopmans et al., 1997). Thus, flux concentration of corresponded pollutants can be also assessed from box model as mentioned in front.

450 **2.3 Estimate health impacts and health-related economic losses**

451 **2.3.1 Carcinogenic risk of Smoke Particulate Matter (CR**_{SPM})

452 Apart from the enormous climatic effects of smoke particle emissions, new

epidemiological and toxicological evidence have also linked carbonaceous aerosol to cardiovascular and respiratory health effects according to the World Health Organization (Bruce et al., 1987; IPCC, 2007). Here, we present the fuel-specific carcinogenic risk of SPM (CRSPM, unit: per μ g m⁻³) to assess health hazard from agricultural straw burning particles and help source-specific air quality control. The cancer risk attributed to inhalation exposures of smoke PM_{2.5} from crop straw i burning was calculated as:

460
$$CR_i = \sum_j f_j \times UnitRisk_j$$
 (6)

461 where f_j is mass fraction of individual species j in smoke PM_{2.5}, UnitRisk_j is 462 corresponded unit carcinogenic risk value of species j extracted from database 463 provided by the Integrated Risk Information System (IRIS), California 464 Environmental Protection Agency (CEPA), and related documents (Bruce et al., 465 1987; Burkart et al., 2013; Tsai et al., 2001; Wu et al., 2009, 2011).

CR_i is estimated based on dose addition model of selected hazardous air 466 pollutants (HAPs) including USEPA priority pollutants of PAHs and heavy metals. 467 And UnitRisk values of the selected HAPs presented in Table S5 (SI). Synergistic 468 469 interactions among pollutants are dismissed, albeit possible. The cancer risk of chromium is adjusted by multiplying a factor of 0.2, assuming that only 20% Cr 470 measured is in the toxic hexavalent form (Bell and Hipfner, 1997). Benzo[a]pyrene 471 (BaP) is used as an indicator compound of carcinogenicity, legally binding threshold 472 of BaP in most countries ranges from 0.7 to 1.3 ng m⁻³, corresponded carcinogenic 473 risk of BaP is about 1.1×10^{-6} per ng m⁻³ (Bruce et al., 1987; Burkart et al., 2013). 474 Thus, one in million level of carcinogenic potential is frequently used to identify 475 risks of concern in public health and environmental decision making, and 476 permissible exposure limits (PEL, unit: µg m⁻³) of crop straw burning particles can 477 be estimated as: 478

479
$$\text{PEL}_{i} = \frac{10^{-6}}{CR_{i}}$$
 (7)

480 **2.3.2 Human exposure and health impacts**

Robust relationship between surface PM2.5 and health effects has been revealed and 481 confirmed by many studies (Pope et al., 2004; Wong et al., 2008). PM_{2.5}-related 482 health endpoints are composed of a range of elements from sub-clinical effects to the 483 484 onset of diseases and the final death (Davidson et al., 2005). In this study, incidence of commonly studied endpoints like premature mortality, respiratory and 485 cardiovascular hospital admissions, and chronic bronchitis from primary emitted 486 smoke PM_{2.5} short-term exposure were assessed using the Poisson regression model, 487 488 shown as below (Guttikunda and Kopakka, 2014):

489
$$\Delta E = \Delta Pop \times IR \times (1 - \frac{1}{e^{\beta \times \Delta C}})$$
(8)

490 where ΔE represents the number of estimated cases of mortality and morbidity, ΔC 491 is the incremental concentration of particulate matter (PM) or flux concentration; 492 ΔPop is the population exposed to the incremental particulate concentration of ΔC ; 493 IR is short for incidence rate of the mortality and morbidity endpoints, and β is the 494 coefficient of exposure-response function, defined as the change in number case per 495 unit change in concentration per capita.

Concentration-response function and incidence rate of each health endpoint are 496 important in health impacts evaluation and they have variation for different 497 population and regions (Yang et al., 2012; Wong et al., 2008). Here, the variance for 498 sex and ages were neglected. Region-specific exposure-response coefficients for 499 500 individual mortality were summarized from previous studies, as presented in Table S6 (SI). The coefficients for individual respiratory and cardiovascular hospital 501 admission, and chronic bronchitis were cited as 1.2%, 0.7%, and 4.4% (per 10 µg 502 m⁻³, 95% CI) from Aunan and Pan's work (Aunan and Pan, 2004). This is the case 503 504 because seldom studies ever confirmed these topics in China. Region-specific mortality and hospitalization IRs were taken from statistical reports authorized by 505 National Health and Family Planning Commission of the People's Republic of China 506 (NHFPC, 2013), and morbidity of chronic bronchitis were defined as 13.8 % based 507

on the forth national health survey, which was released by the Chinese Ministry ofHealth in 2008 (CMH, 2009).

510 **2.3.3 Economic valuation of the health impacts**

The economic losses of the health impacts associated with smoke $PM_{2.5}$ exposure in 2012 were further evaluated. The amended human capital (AHC) approach was employed to calculate the unit economic cost of premature mortality. The commonly applied AHC method uses per capita GDP to measure the value of a statistical year of life (IBRD and SEPA, 2007) based on Eq. (9). It can be used as a social statement of the value of avoiding premature mortality and estimates human capital (HC) from the perspective of entire society, neglecting individual differences (Hou et al., 2012).

(9)

518
$$HC_{k} = \frac{GDP_{k}}{POP_{k}} \times \sum_{i=1}^{\tau} \frac{(1+\alpha)^{i}}{(1+\gamma)^{i}}$$

 GDP_k and POP_k are gross domestic production and population of target region k 519 that were reported in the statistical yearbook in 2012; α and γ are economic 520 parameters referring to national GDP growth rate and social discount rate, which 521 were 7.7% and 8.0% in 2012 from National Bureau of Statistics of China (NBSC, 522 2013, data available at http://www.stats.gov.cn/tjsj/ndsj/, in Chinese). τ is the 523 524 life-expectancy lost due to aerosol pollution, and 18 years of life was widely applied (Hou et al., 2012). The annual exchange rate of US dollar to RMB was 6.31 in 2012. 525 One can deduce the HC values of the provinces, municipalities, and autonomous 526 regions in the country, and the calculated regional HC values were listed in Table S7 527 (SI). In this paper, the cost of respiratory, cardiovascular hospital admissions, and 528 chronic bronchitis were 632.2, 1223.4, and 948.6 US\$ per case in 2012, which were 529 530 derived from the national health statistical reports (NHFPC, 2013).

The regional and national health-related economic loss from smoke $PM_{2.5}$ exposure can be calculated based on the excess mortality and morbidity multiplied by the corresponding unit economic values.

534 **3 Result**

535 **3.1 Particulate chemical compositions and emission factors**

536 3.1.1 Organic carbon and elemental carbon

An overview of particulate chemical compositions for smoke $PM_{2.5}$ and $PM_{1.0}$ is 537 pie-graphically profiled in Fig. 2, and the corresponded emission factors are given in 538 539 Table 1-2 (detailed EFs for elements, PAHs, and Phenols in Table S8 and S9, SI). From multivariate statistical analysis (P<0.05 at 95% CI), significant differences of 540 chemical compositions and emissions in size range and fuel types can be observed, 541 implying the nonuniform mixing and distribution of particulate pollutants from 542 biomass burning, which is consistent with the conclusion from Lee et al. (2015) and 543 Giordano et al. (2015). EFs of particulate species from this study were compared 544 with that from literature as summarized in Table 3, since EFs in smoke $PM_{1,0}$ were 545 546 seldom reported, only smoke PM_{2.5} or total particulate matter emissions were collected, which were comparable with the results in this work. EFs of smoke PM_{2.5} 547 and PM_{1.0} were in range of $3.25 \sim 15.16$ and $3.04 \sim 13.20$ g kg⁻¹ for the five kinds of 548 crop straws, a high ratio of $PM_{1.0}/PM_{2.5}$ was observed to be over 90 wt.%, which was 549 in line with size distribution analysis of smoke particles given in Fig. S3 (SI). Li et al. 550 (2007) measured the emissions from field burning of crop straws via CMB method, 551 $PM_{2.5}$ EFs for wheat and corn straw were estimated to be 7.6 ± 4.1 and 11.7 ± 1.0 g 552 kg^{-1} (dry basis, MCE > 0.9), which were higher and presented more uncertainties 553 than our result. As study ever found a positive relationship between particulate EFs 554 555 and moisture content of agricultural residue (Hayashi et al., 2014), it was reasonable that combustion of the dehydrated crop straw produced less smoke aerosol in this 556 work. Hayashi et al. (2014) measured particulate EFs to be 2.2 and 15.0 g kg⁻¹ for 557 rice and wheat straw of ~10 wt.% moisture content, while corresponded EFs 558 increased to 9.1 and 19.5 g kg⁻¹ when water content of straw was ~20 wt.%, and the 559 linear equations between smoke EFs and straw moisture content were furtherly 560 proposed. However, the simple linearity and its application scope should be doubted, 561

as Hayashi et al. only considered two water content levels (10 wt. % *vs* 20 wt. %) and disregarded influence of combustion efficiency for the fires. $PM_{2.5}$ EFs given by Dhammapala et al (2006, 2007a, b) were 4.7 ± 0.4 g kg⁻¹ for wheat straw and $12.1 \pm$ 1.4 g kg⁻¹ for herbaceous fuel that were burnt using a chamber under flaming phase, and negative response for particulate EFs to combustion efficiency was observed. After all, smoke EFs vary with fires depend on fuel type and moisture, combustion phase, environmental conditions, and some other variables (Reid et al., 2005b).

The carbonaceous materials (Organic matter and EC) are dominated in SPM, 569 accounting for about 73.4 wt.% for $PM_{2.5}$ and 71.3 wt.% for $PM_{1.0}$ on average. 570 Organic matter (OM) was converted from OC by multiplying a factor of 1.3 to 571 account for noncarbon materials like oxygen, hydrogen, and other minor species (Li 572 573 et al., 2007; Li et al., 2015), and Li et al. (2016) ever measured OM/OC ratio as ~1.3 574 for fresh smoke particles via volatility analysis. EFs of EC and OC from this work were consistent with most studies, average OC EFs were 4.21 and 3.58 g kg⁻¹ in 575 smoke $PM_{2.5}$ and $PM_{1.0}$, and the corresponded EC EFs were 1.09 and 1.01 g kg⁻¹, 576 respectively. These values fell within the ranges (0.9~9.3 g kg⁻¹ for OC and 0.2~1.7 577 $g kg^{-1}$ for EC) found in other similar sources (Dhammapala et al., 2007; Hayashi et 578 al., 2014; Li et al., 2007; May et al., 2014). Due to the technical limitation and 579 ambiguous artificial boundary, carbon contents of biomass burning particles have 580 vast variability and uncertainty (Lavanchy et al., 1999; Levin et al., 2010). It was 581 ever reported chamber burn study may overestimate EC EFs due to a misassigned 582 OC-EC split for the heavily mass loaded filter samples (Dhammapala et al., 2007b). 583 Moreover, carbon measurement based on TOT method with NIOSH protocol may 584 overestimate OC fraction by sacrificing EC part compared with that of TOR 585 (Thermal-Optical Reflectance) method with IMPROVE program (Han et al., 2016). 586 Mass ratio of OC/EC is a practical parameter to indicate the primary organic aerosol 587 588 (OA) emission and secondary organic aerosol (SOA) production. The ratio is influenced by burning conditions, source, aging extent, and particle size (Engelhart 589 et al., 2012; Grieshop et al., 2009). Smoke emitted from smoldering fires is 590 OC-dominated while flaming combustion produces more EC, and the discrepancy of 591

592 OC/EC ratio can be an order of magnitude regarding to different combustion phase 593 (Grieshop et al., 2009). SOA production upon photo-oxidation will enlarge OC/EC 594 ratio, and positive relation between oxidation level of OA loading and OC/EC ratio was reported (Grieshop et al., 2009). Here, OC/EC ratio in primary emissions varied 595 from 2.4 to 6.2 under flaming phase, similar to previous studies (Arora and Jain, 596 2015; Dhammapala et al., 2007a, b; Hayashi et al., 2014; Lewis et al., 2009). The 597 598 OC/EC ratios were larger in $PM_{2.5}$ with average value of 3.8, while it was 3.6 in $PM_{1,0}$, indicating more EC resides in $PM_{1,0}$. 599

600

3.1.2 Water soluble organic acids

Smoke particles comprise a considerable amount of water soluble organic acids 601 602 (WSOA), it was 3.35 wt.% in PM_{2.5} and 3.17 wt.% in PM_{1.0} on average, which was in line with previous work that organic acids measured represent less than 5 wt.% of 603 the total smoke aerosol mass load and favor to partition in larger size (Falkovich et 604 al., 2005; Gao et al., 2003). Acetic acid followed by methysulfonic acid contributed 605 606 the most of the measured low molecule weight acids. Oxalic acid is the dominated dicarboxylic acids measured in the ambient environment and biomass burning 607 aerosol (Falkovich et al., 2005; Kundu et al., 2010), and oxalic acid EF was 608 measured to be 2.2 ~ 4.8 and 1.6 ~ 3.6 mg kg⁻¹ for smoke PM_{2.5} and PM_{1.0} in present 609 work. The sums of WSOA EFs ranged from 46.7 to 770.0 mg kg⁻¹. Correlation 610 among the multi-pollutants was analyzed by relevance matrix as shown in Table S10 611 (SI), the strong positive linear relationship (R^2 >0.99, p<0.05 at 95% CI) between 612 WSOA and emissions of OC and PM was observed. Study has confirmed organic 613 acids contribute a significant fraction of both oxygenated volatile organic 614 compounds (OVOCs) in gaseous phase and SOA in particulate phase, the direct 615 emission of particulate organic acids from biomass burning also represents a 616 significant source of precursors for SOA formation, as the low molecular organic 617 acids will evaporate into gas phase or involve in the heterogeneous reaction directly 618 (Takegawa et al., 2007; Veres et al., 2010; Yokelson et al., 2007; Carlton et al., 2006). 619

620 Moreover, as the significant fraction of water soluble organic carbon, organic acids play major response to CCN activity of smoke particles, and organic acids coating or 621 622 mixing can amplify hygroscopic growth of inorganic salts by decreasing the deliquescence RH, enable the particle to be CCN at relative low degree of 623 supersaturation (Falkovich et al., 2005; Ghorai et al., 2014). In the ambient 624 environment, organic acids can enhance atmospheric new particle formation by 625 626 impairing nucleation barrier (Zhang et al., 2004), besides, particulate organic acids can also mobilize the solubility of mineral species, like iron, altering the chemical 627 process of particles (Cwiertny et al., 2008). And prominent optical properties of 628 organic acids like humic/fulvic substance make them as potential contributors to the 629 global warming (Yang et al., 2009; Andreae and Gelencs ér, 2006). 630

631 **3.1.3 Water soluble aminiums**

Interest has been focused on the vital role of amines in particle nucleation-growth 632 process and acidity regulating due to their strong base (Tao et al., 2016; Bzdek et al., 633 2010, 2011). Though ultratrace gaseous amines and particulate aminiums were on 634 the order of pptv or ng m^{-3} , aminium salts exhibit potential climatic and health effect 635 due to their significant different properties in hygroscopicity, optics, and also 636 toxicology (Qiu and Zhang, 2012; Qiu et al., 2011; Samy and Hays, 2013; Zheng et 637 638 al., 2015; Ho et al., 2015; Tao et al., 2016). It ever proposed that biomass burning is an important source for gaseous amines, especially from smoldering burning, and 639 alkyl amides can be served as biomarkers in particular (Ge et al., 2011; Ho et al., 640 641 2015; Lee and Wexler, 2013; Lobert et al., 1990; Simoneit et al., 2003). However, seldom study ever quantitatively explored the particulate water soluble amine salts 642 (WSA) in primary smoke emissions (Schade and Crutzen, 1995; Ge et al., 2011). 643 From this study, WSA contributed about 4.81 wt.‰ of smoke PM_{2.5} and 4.69 wt.‰ 644 of PM_{1.0}, implicating aminium favored to be abundant in fine-mode of smoke 645 particles, especially in PM_{2.5-1.0}. DEAH⁺, TMAH⁺, TEOH⁺ and DMAH⁺ made up 646 over 80 wt. % of the measured WSA. Fuel-dependence of WSA distribution and 647

emissions were evident. EFs of WSA ranged from 4.5 to 104.8 mg kg⁻¹ in smoke 648 PM_{2.5}, the least was from burning of soybean straw and the largest from cotton and 649 rice straws. We used mass ratio of WSA to NH_4^+ to denote the enrichment of 650 aminium in particulate phase. Statistical analysis showed WSA/NH4^+ was 0.16 \pm 651 0.03 and 0.18 \pm 0.06 in smoke PM_{1.0} and PM_{2.5}, respectively, which were almost one 652 order of magnitude larger than that in the ambient aerosol (Liu and Bei, 2016; Tao et 653 654 al., 2016). Tao et al. (2016) ever measured the ratio as a function of particle size during NPF days in Shanghai, and a noticeable enrichment of aminiums for ultrafine 655 particles (<56 nm) was observed with WSA/NH₄⁺ over 0.2, highlighting the 656 competitive role for amines to ammonia in particle nucleation and initial growth of 657 the nuclei, the ratio was then decreased with the increasing particle size, and the final 658 increasing trend was found after ~ 1.0 μ m, and average WSA/NH₄⁺ for ambient bulk 659 660 $PM_{1.0}$ and $PM_{2.5}$ were 3.2% and 3.5%, respectively.

661 **3.1.4 PAHs and Phenols**

662 Atmospheric PAHs are primarily the byproduct of incomplete combustion of biomass and fossil fuels (Simcik et al., 1999; Galarneau, 2008). Due to their high 663 degree of bioaccumulation and carcinogenic or mutagenic effect, the sources and 664 environmental fate of the ubiquitous PAHs have been the subjects of extensive 665 studies (Santodonato, 1997; Kim et al., 2013). PAHs can involve in photochemical 666 reaction to form SOA, the process is influenced by gas-to-particle partition and 667 meteorological conditions. Moreover, oxidation may increase the toxicity of PAHs 668 669 (Arey and Atkinson, 2003; Wang et al., 2011). Biomass burning is one of the main sources of gaseous and particulate PAHs, which even contributes to about half of 670 anthropogenic PAHs emissions in China (Xu et al., 2006; Zhang et al., 2011). 671 Burning conditions can significantly influence the emission of PAHs, under the 672 flaming phase in this study, PAHs contributed 0.46 wt.‰ of smoke PM_{2.5} and 0.28 673 wt.% of PM_{1.0}, over 60% of the total PAHs were associated to respiratory submicron 674 particles. The sum of EFs of 16 PAHs in smoke PM2.5 ranged from 1.81 to 8.30 mg 675

 kg^{-1} , which were consistent with the values from literature (Dhammapala et al., 676 2007a, b; Lee et al., 2005; Zhang et al., 2011). Hays et al. (2005) estimated total EFs 677 of 16 PAHs to be 3.3 mg Kg⁻¹ in wheat straw burning PM_{2.5}. Korenaga et al. (2001) 678 measured PAHs EFs from rice straw burning to be 1.9 mg Kg⁻¹ in particulate phase, 679 while the value from Jenkins et al. (1996) was 16 mg Kg⁻¹. Dhammapala et al. 680 (2007b) found negative linear response for biomass burning source PAHs emissions 681 to burning efficiency, and under flaming combustion, particulate total 16 PAHs EFs 682 were $2 \sim 4 \text{ mg Kg}^{-1}$. Zhang et al. (2011) simulated burning of rice, corn, and wheat 683 straws, the corresponded PAHs EFs were measured as 1.6, 0.9, and 0.7 mg Kg⁻¹ in 684 fine smoke particles, respectively. Great uncertainties for PAHs EFs were evident 685 that relied on burning phase, fuel types, moisture content, and also measurement 686 techniques. Dhammapala et al. (2007a) also found laboratory simulation might 687 688 overestimate the emission factors of PAHs compared with field burnings. EFs for individual PAHs were included in Table S8 and S9 (SI). The distribution of 689 690 particulate PAHs emission factors was presented in Fig. 3a. Of the particle bound PAHs, 3~4-rings components were the primary ones, including Pyr, Ant, Ace, Flu, 691 Phe, and Chr. Concentration ratios of selected PAHs, namely diagnostic ratios, were 692 usually used to trace the source and make apportionment of specific pollutions 693 (Yunker et al., 2002; Simcik et al., 1999). In this work, average Ant/(Ant+Phe), 694 Flu/(Flu+Pyr), BaA/(BaA+Chr), and IP/(IP+BghiP) ratios of 5 types agricultural 695 residue burning smokes were 0.72, 0.36, 0.47, and 0.58, respectively. There was no 696 significant difference (P<0.05 at 95% CI) of the ratios in $PM_{1.0}$ and $PM_{2.5}$. According 697 to previous work, Ant/(Ant+Phe) above 0.1 and BaA/(BaA+Chr) above 0.35 indicate 698 the dominance of combustion and pyrolytic sources, Flu/(Flu+Pyr) and IP/(IP+BghiP) 699 ratios greater than 0.50 suggest coal or biomass burnings dominate (Simcik et al., 700 1999; Yunker et al., 2002). However, validation of source apportionment using 701 702 specific diagnostic ratios should have its constraints, because of variations in source 703 strengths and atmospheric processing of PAHs (Arey and Atkinson, 2003; Galarneau, 2008). 704

From Table S10 (SI), PAHs in smoke particles were highly correlated with EC and

OC contents. PAHs primarily originate from pyrolysis of organic materials during 706 combustion, and formation mechanisms of PAHs and soot are closely intertwined in 707 708 flames. High-molecular-weight PAHs (>500 atomic mass unit) act as precursors of soot particles (Lima et al., 2005; Richter et al., 2000). Thus, PAHs with 3, 4, and 5 709 rings accumulate and dominate in the emissions of biomass burning, as larger 710 molecular weight PAHs tend to incorporate into soot particles. PAHs 711 712 expulsion-accumulation in OC and EC fractions were analyzed by linear fitting of PAHs mass fractions and EC mass fractions in carbonaceous materials (EC+OC) in 713 Fig. 3b. The partitions can be parameterized as Eq. (10): 714

715
$$f_{PAHs} = \frac{m_{PAHs}}{m_{OC} + m_{EC}} = \beta_{EC} \times \frac{m_{EC}}{m_{OC} + m_{EC}} + \beta_{OC} \times \frac{m_{OC}}{m_{OC} + m_{EC}} = \beta_{EC} \times f_{EC} + \beta_{OC} \times f_{OC}$$
(10)

where f_{EC} and f_{OC} are the mass fraction of OC and EC in carbonaceous materials (EC+OC). β_{EC} and β_{OC} are expulsion-accumulation coefficients of PAHs in OC and BC. The coefficient of β_{EC} is 1.1×10^{-3} in smoke PM_{1.0} and 1.9×10^{-3} in PM_{2.5}, the corresponded β_{OC} is 0.3×10^{-3} and 0.5×10^{-3} .

720 Phenols are the most common SOA precursor/product and organic pollutants in the atmosphere (Berndt and Böge, 2006; Schauer et al., 2001). Hydroxyl functional 721 group and aromatic benzene ring make phenols a paradigm in heterogeneous 722 reaction upon photo oxidation research and aqueous phase reaction research. Phenols 723 724 are also ROS (reactive oxidized species) precursors that present health hazard (Bruce et al., 1987). Phenol and substituted phenols are thermal products of lignin pyrolysis 725 during biomass burning (Dhammapala et al., 2007a), and the most abundant 726 methoxyphenols can be markers of biomass burning sources (Urban et al., 2016). 727 The five measured phenols contributed 3.0 wt.% and 2.5 wt.% of $PM_{2.5}$ and $PM_{1.0}$. 728 2, 6-dimethoxyphenol was the major one of the measured phenols. Mass fraction of 729 phenols was about 7~9 times of PAHs in smoke aerosols. EFs for the sum phenols 730 were 9.7 ~ 41.5 and 7.7 and 23.5 mg Kg⁻¹ for smoke PM_{2.5} and PM_{1.0}, respectively. 731 Dhammapala et al. (2007a) estimated particulate methoxyphenols emissions to be 35 732 \pm 24 mg Kg⁻¹ for wheat straw burning, while Hays et al. (2005) measured the same 733 compounds to be 6.8 mg Kg⁻¹. Carbonaceous materials like PAHs and Phenols or 734

aromatic and phenolic deviates are the main chromophores in the atmosphere, and the considerable fractions of PAHs and Phenols justify biomass burning as a significant source of brown carbon (Laskin et al., 2015), study has proved ~ 50% of the light absorption in the solvent-extractable fraction of smoke aerosol can be attributed to these strong BrC chromophores (Lin et al., 2016).

740 **3.1.5 Inorganic components**

From Fig. 2, smoke particles consisted of approximately 24 wt.% water soluble 741 inorganics (WSI), and the inorganic salts resided more in submicron particles. Great 742 amount of inorganics enable smoke particles to be efficient CCN, and the distinct 743 optical scattering characters of the inorganic fractions may neutralize the warming 744 745 effect of brown carbon for smoke aerosol, otherwise, inorganics coating or mixing will enhance light absorbing of BC. K⁺, NH₄⁺, Cl, and SO₄²⁻ were the principle 746 inorganic ions. Particulate enriched K⁺ together with levoglucose are treated as tracer 747 of pyrogenic source (Andreae et al., 1998). And specific mass ratio of K⁺/OC or 748 749 K^+/EC will help make source apportionment of particulate pollutants with PMF (Positive Matrix Factorization) and PFA (Principle Balance Analysis) models (Lee et 750 al., 2015). K⁺/OC in smoke particles ranged from 0.11 to 0.25 with average value of 751 0.17 in $PM_{1.0}$ and 0.14 in $PM_{2.5}$, which were similar to those reported for the 752 Savannah burning and agricultural waste burning emissions in India and China 753 (Echalar et al., 1995; Ram and Sarin, 2011; Li et al., 2015). However, OC represents 754 large uncertainty arise from degree of oxidization and burning condition, K⁺/EC is 755 756 more practical parameter to distinguish the pyrogenic pollutants in ambient study. To smoke particle emitted from flaming fires, K^+/EC was 0.58 \pm 0.24 in PM_{1.0} and 0.53 757 ± 0.18 in PM_{2.5}. Cl⁻ was the main anion to balance the charge of WSI in smoke 758 particles. Mean charge ratio of Cl^{\cdot} : K⁺ was 1.46 and 1.49 in PM_{1.0} and PM_{2.5}, 759 implicating surplus chloride will associate with other cations. With atmospheric 760 aging, the Cl/K ratio will decrease as chloride being replaced by secondary sulfate 761 and nitrate (Li et al., 2015; Li et al., 2003). Equivalent charge ratio of primary 762

cations $(NH_4^+ + K^+)$ to primary anions $(SO_4^{2-} + CI)$ was 1.05 in PM_{1.0} and 1.01 in 763 PM_{2.5} on average, and charge ratios of total cations to anions (R_{C/A}) was 1.09 and 764 1.07 in PM_{1.0} and PM_{2.5}. R_{C/A} was used to indicate the neutralizing level of 765 particulate matters in many studies. $R_{C/A} \ge 1$ indicates most of the acids can be 766 neutralized, while R_{C/A}<1 means atmospheric ammonia is deficient and the aerosol is 767 acidic (Adams et al., 1999; He et al., 2011a; Kong et al., 2014). In ambient 768 769 environment, acidic aerosol was prevailing urban pollutants in many cities from field investigation (He et al., 2011a; Kong et al., 2014). Acidic aerosols can increase the 770 risks to human health and affect the atmospheric chemistry by activating hazardous 771 materials and promoting the solubility of particulate iron and phosphorus (Amdur 772 and Chen, 1989; Meskhidze, 2005). The emission and transport of biomass burning 773 particles may neutralize the acidity of ambient particles. However, only limited WSI 774 775 were brought into in the analytical system, it is not really to tell the acidity or base of smoke particles, considering the existence of massive organic acids and ammniums. 776

777 Trace mineral elements attracted great attention for the role as catalyst in atmospheric heterogeneous reaction and health cares (Davidson et al., 2005; 778 Dentener et al., 1996). Wet/dry deposition of particles during long range transport 779 will affect the ecological balance by releasing mineral elements (Jickells et al., 2005). 780 Dust storm, weathering, and industrial process are the main sources of particulate 781 metals, and incineration can also produce a lot of mineral elements (Moreno et al., 782 2013). However, the emissions of trace metals from biomass burning are highly 783 uncertain (Li et al., 2007; Zhang et al., 2012), the great influence from local soil 784 785 environment and soil heavy metal pollution will certainly affect the metal content in biomass fuel and smoke particle. In this study, THM resided more in PM_{2.5} than in 786 PM_{1.0}. Smoke PM_{2.5} consisted of 6.7 wt.‰ THM on average, PM_{1.0} comprised 4.1 787 wt.‰ THM. Average EFs of THM in $PM_{2.5}$ and $PM_{1.0}$ were 0.056 g kg⁻¹ and 0.028 g 788 kg⁻¹ in this work, of which Al contributed over 90 wt.‰, in line with result from 789 domestic burning of wood and field investigation of crop straw burning (Li et al., 790 2007; Zhang et al., 2012). Smoke particles from wheat, rice, and corn straws 791 contained more mineral elements than that from cotton and soybean residues 792

combustion. Regardless the difference in biomass fuels, the result may imply that
soil heavy metal pollution is heaver in the East China than that in Xinjiang in the
West North of China (Wei and Yang, 2010).

796 **3.2 Size, morphology, and mixing state of smoke particles**

797 Fresh smoke particles exhibited unimodal size distribution within 500 nm (Fig. S3, SI), and previous chamber study has also confirmed that agricultural fire produces 798 large amount of ultrafine particles, implying the great potential role to act as CCN 799 and more profound threat to human health (Araujo et al., 2008; Delfino et al., 2005; 800 Zhang et al., 2011). However, the role of particles in the atmospheric process and 801 health hazard depends not only size, but also morphology and chemical mixing 802 803 states (Dusek et al., 2006; Kennedy, 2007; Mikhailov et al., 2006; Schlesinger, 1985). From TEM images in Fig. 4, agricultural straw burning aerosols comprised a broad 804 class of morphological and chemically heterogeneous particles. Non-uniformly 805 internal mixing of the agglomerates was noticeable, including the major 806 807 carbonaceous particles and a considerable amount of inorganic salt particles, which was consistent with previous particulate chemical analysis. KCl particles containing 808 minor sulfate or nitrate were the primary inorganic particles, which presented crystal 809 or amorphous state from X-ray diffraction analysis (Fig. 4 a, b, c). And 810 811 potassium-bearing particles have been used as a tracer of ambient biomass burning pollutants. Fly ash particles were arresting due to visible morphology difference and 812 mineral chemical composition (Fig. 4 d, e, f). Fly ash particles were more compact 813 814 and rich in mineral elements like Ca, Si, Fe, Al, Mn, and Cr. Besides, these particles had larger size, statistical average diameter of fly ash particles obtained from bulk 815 analysis was $2.2 \pm 1.6 \mu m$. The result also proved heavy metals resided more in 816 PM_{2.5} than PM_{1.0}. Fly ashes are by products of incineration process (Buha et al., 817 2014), including coagulation of fuel issue debris, condensation of evaporated 818 mineral metal from biomass fuels or adhered dirt at different burning phase. These 819 820 fly ashes coated by or agglomerated with carbonaceous materials were like mash of

821 mineral without clear lattice. Tar ball as a specific form of brown carbon and soot 822 were representative particles of biomass burning aerosol (Wilson et al., 2013; 823 Chakrabarty et al., 2010; Tóth et al., 2013). From Fig. 4 g, chain-like soot particles were coagulated with tar ball. Soot particles were agglomerates of small roughly 824 spherical elementary carbonaceous particles, these chemical consistent particles were 825 within 20~30 nm, and high-resolution TEM showed the soot spheres consisted of 826 827 concentrically wrapped graphitic layers, while monomeric tar balls possessed disordered microstructure. Tar balls and soot corresponded to different stages in the 828 aging of organic particles; tar balls abundant in fresh or slightly aged biomass smoke 829 are formed by gas-to-particle conversion of high-molecular weight organic species 830 or from aged primary tar droplets upon biomass burning. Soot represents further 831 aged carbon-bearing particles, formed from the pyrolysis of lignin, cellulose, or tar 832 833 balls (Pósfai, 2004; Tóth et al., 2013). The botryoid aggregates in Fig. 4 g can be viewed as transformation of tar ball to soot. Tar ball and soot were also internal 834 mixed with inorganic salt including sulfate and nitrate (Fig. 4 g, h, i), which made 835 the physiochemical properties of BC even complicated, as study has confirmed 836 inorganic sulfate mixing will enhance light absorption and hygroscopicity of BC 837 (Zhang et al., 2008b). Dark-ring like shell of tar ball (Fig. 4 g, h) and spot-like 838 particles adhered to the surface of tar ball (Fig. 4 i) were K-rich materials. And size 839 of soot particles was mainly within 200 nm, while tar ball and other carbonaceous 840 841 particles can be over one micrometer.

842 **3.3 Open burning emissions**

843 **3.3.1 Crop straw production**

The agricultural straw productions were calculated and geographically displayed in Fig. 5 a-c. Totally 647.3 Tg agricultural straws were produced in 2012 and dispersed mainly in the North and Northeast of China. The distributions of the straws clearly correspond to the distinct planting regions that are divided by Qinling Mountain-Huaihe River line and the Yangtze River. Rice is primarily planted in the 849 south of Qinling Mountain-Huaihe River line, only 10 % rice (single cropping rice dominate) is planted in Heilongjiang, Jilin, and Liaoning province, while wheat and 850 851 corn are grew mostly in the north of the Yangtze River. Over 90 % of the wheat planted in China is winter wheat that gets ripe in summer, and more than 80 % rice 852 including middle and late rice grows mature in autumn. Summer harvest contributed 853 about 25 % of the agricultural straw production, which solely consists of rice and 854 855 wheat straws in this period and distributes uniformly in the central and east of China. 493.9 Tg crop straws were produced mainly from corn and rice harvesting in autumn. 856 Soybean and cotton straws account for about 8.6 % of autumn straw production that 857 were primarily produced in Heilongjiang and Xinjiang province. 858

859 3.3.2 Open burning rate

The five scenarios of field burning rates and regional AIP $\left(\frac{I_{k,y}}{AI_{k,y}}\right)$ in the year of 860 2000, 2006, and 2012 were listed in Table 4 and statistically analyzed in Fig. 6. A 861 significant difference (P<0.05 at 95% CI) of regional burning rates among the 862 versions was observed, and the rates from NDRC report were generally higher. For 863 convenience, six zones were classified by geographic divisions and economic areas 864 865 in China, including the North Plain of China (NPC: Anhui, Shandong, Hebei, Shanxi, Tianjin, Beijing), the Central of China (CC: Hunan, Henan, Hubei), the Yangtze 866 River Delta (YRD: Zhejiang, Jiangsu, Shanghai), the Northeast of China (NC: 867 Heilongjiang, Liaoning, Jilin), the Pan-Pearl River Delta (PRD: Hainan, Guangdong, 868 Fujian, Guangxi, Guizhou, Sichuan, Yunnan, Jiangxi), the West of China (WC: 869 Shannxi, Chongqing, Xinjiang, Qinghai, Ningxia, Tibet, Inner Mongolia, Gansu). 870 871 And the bulk-weighted burning rates that averaged from BAU, EM, and NDRC 872 versions for the six zones were 22.3 % ± 3.1 %, 21.1 % ± 3.3 %, 28.4 % ± 6.2 %, $23.3 \% \pm 9.2 \% 21.4 \% \pm 6.5 \%$, and $14.2 \% \pm 8.0 \%$, respectively. It was obvious 873 that agricultural field burning was most serious in the Yangtze River Delta, 874 especially in the Zhejiang province. The nationwide filed burning rate was 21.4 %, 875 16.3 %, 26.0 %, 14.9 %, and 26.8 % for BAU-I, BAU-II, EM-I, EM-II, and NDRC, 876

respectively, which were comparable with the document values (Daize, 2000; Wei et al., 2004; Zhang et al., 2008a).

879 **3.3.3 Agricultural open burning emissions**

PM_{2.5} emissions from agricultural field burnings based on BAU, EM, and NDRC 880 versions were calculated and geographically presented in Fig. 7 (emissions of 881 detailed individual species in SI). A similar spatial character of regional emission 882 distribution was observed for BAU, EM, and NDRC versions, most emissions were 883 allocated in the North Plain and the Central of China, where the primary agricultural 884 regions locate, echoing the agricultural fire sites in Fig. S2 (SI). Although filed 885 burning rates were higher in the Yangtze River Delta, the crop residue productions in 886 887 this zone were much less, which only contributed 4.3 % of the national straw productions. Take NDRC as the basis, BAU and EM scenarios all underestimated the 888 emissions in the Northeast of China, especially in Heilongjiang. 889

The temporal distributions of field burning emissions also echoed the crop residue 890 891 productions and the agricultural fire sites in summer and autumn harvest. Apart from Henan and Tibet where the main crop straws were produced in summertime, more 892 pollutants were emitted in autumn harvest period to the rest place, which has been 893 confirmed by many studies (He et al., 2011; Wang and Zhang, 2008). And the large 894 895 scale filed burning emissions in autumn exhibited great influence on the haze formation and visibility degradation in the North and East of China (Leng et al., 896 2014; Shi et al., 2014), Huang et al. (2012a) has identified biomass burning together 897 898 with secondary inorganic aerosol (SIA) and dust pollution as three typical haze types in Shanghai. In summertime, filed burning emissions concentrated in the North Plain, 899 the Central, and the South regions. While in autumn, the emissions became more 900 ubiquitous and serious in the Northeast of China. 901

Nationwide emission inventories and flux concentrations were graphically displayed in Fig. 8 and tabular presented in Table 5. The total $PM_{2.5}$ emission from agricultural field burnings was 738.36-1241.69 Gg in 2012, and rice, corn, and

905 wheat straw burnings made up $93.5\% \sim 95.6\%$ of the total emissions. The largest quantities of PM_{2.5} emissions were emitted from Heilongjiang, Shandong, Henan, 906 Jilin, Jiangsu, Anhui and Hebei, distinct difference in the emissions from various 907 scenarios were observed, especially for Heilongjiang province which contributed 908 5.5 % (55.4 Gg) of PM_{2.5} emissions under BAU-II scenarios, while the figure was 909 22.9 % (231.0 Gg) under EM-I scenarios. Annual emissions of PM_{1.0}, OC, and EC 910 911 was 661.81-1111.90, 318.84-533.19, and 98.06-164.97 Gg, respectively, which were comparable with the precious studies (Cao et al., 2006, 2011; Wang et al., 2012). Qin 912 and Xie (2011, 2012) developed national carbonaceous aerosol emission inventories 913 from biomass open burning for multi-years with dynamic burning activity, they 914 believed BC and OC emissions followed an exponential growth from 14.03 and 915 57.37 Gg in 1990 to 116.58 and 476.77 Gg in 2009. Cao et al. (2006, 2011) 916 917 calculated smoke aerosol emissions from biomass burning in China for 2000 and 2007 using the same activity data from BAU-I scenarios, national OC and EC 918 919 emissions were reported to be 425.9 and 103.0 Gg in 2000, however, no evident changes were found for the emissions in 2007, which were assessed to be 433.0 and 920 104.0 Gg. Huang et al. (2012b) estimated crop burning in the fields with unified EFs 921 and burning rate (~6.6%) for all kinds of crops across China in 2006, the estimated 922 annual agricultural fire emissions were about 270, 100, and 30 Gg for PM_{2.5}, OC, 923 and BC, respectively. In present work, agricultural fire PM_{2.5} emissions in 2012 were 924 allocated into six zones, average contribution in percentage for each zone was 925 compared: NPC $(23.1\%) \ge$ NC (21.6%) > PRD $(18.4\%) \ge$ CC (18.2%) > WC 926 (9.8%) >YRD (8.8%). Furtherly, contribution for summertime emissions was: NPC 927 $(35.5\%) > CC (28.8\%) \ge PRD (21.1\%) > YRD (9.1\%) > WC (5.4\%) > NC (0.1\%),$ 928 and for autumn harvest emissions: NC (27.8%) > NPC (19.6%) > PRD (17.6%) >929 CC (15.1%) > WC (11.1%) > YRD (8.8%). It was obviously that the North Plain 930 931 experienced extensive crop fire emissions during the whole harvest periods, where PM_{2.5}, PM_{1.0}, OC, and BC emissions in 2012 were 233.6, 209.8, 102.3, and 29.4 Gg 932 on average. Liu et al. (2015) developed emission inventories from agricultural fires 933 in the North Plain based on MODIS fire radiative power, emission for PM_{2.5}, OC, 934

935 and BC in 2012 was reported to be 102.3, 37.4, and 13.0 Gg, respectively. However, EFs were also treated as unified values (e.g., Crop burning EFs for PM_{2.5}, OC, and 936 BC was 6.3, 2.3, and 0.8 g Kg⁻¹) in the work of Liu et al. (2015) that was cited 937 directly from Akagi et al. (2011) without considering fuel type dependence of EFs. 938 Zhao et al. (2012) established comprehensive anthropogenic emission inventories for 939 Huabei Region including the North Plain, Inner Mongolia, and Liaoning province, 940 941 all crop straws were assumed to be burnt in the field, resulting in much more emissions of 446 Gg OC and 160 Gg BC in 2003. A specific temporal pattern for 942 agricultural fire emissions was observed in the Northeast of China (Heilongjiang, 943 Liaoning, and Jilin), where the open burning were mainly occurred in autumn 944 harvest to produce great amount of pollutants (217.5 Gg PM_{2.5}, 89.4 Gg OC, and 945 29.7 Gg EC), while emissions in the summertime can be neglected. 946

947 In 2012, 20-25 % of national emissions were released from summertime field burnings, that was 226.0 Gg PM_{2.5}, 205.2 Gg PM_{1.0}, 105.9 Gg OC, 28.4 Gg EC, 6.8 948 949 Gg WSOA, 1.0 Gg WSA, 0.1 Gg PAHs, 0.9 Gg phenols, and 2.1 Gg THM on average. The corresponded values for autumn harvest were 781.6, 697.9, 327.3, 950 106.0, 18.4, 4.8, 0.4, 1.9, and 6.6 Gg, respectively. Integrated smoke OC/EC was 3.7 951 from national summertime emission and 3.1 from autumn harvest emission, 952 regarding to different locations, integrated OC/EC in the North Plain was 4.1 in 953 summertime emission and 3.2 in autumn harvest, while OC/EC in the Central of 954 China was 3.1 for both summer and autumn harvest emissions, implying 955 temporal-spatial characters of agricultural field fires exhibit potential influence on 956 composition of smoke emissions and its related physiochemical properties. Zhang et 957 al. (2011) estimated particulate PAHs emissions form three types of crop residues to 958 be 0.46 Gg in 2003. Xu et al. (2006) counted PAHs from all straws with the 959 assumption that burning rates to be unit, and they calculated 5-10 Gg PAHs 960 961 emissions in 2003, which was ten times of our result.

The nationwide flux concentration of smoke $PM_{2.5}$ was 0.7-1.0 µg m⁻³ d⁻¹ in summer harvest and 1.4-3.5 µg m⁻³ d⁻¹ in autumn harvest, while average annual flux concentrations for OC and EC were 0.80 and 0.25 µg m⁻³ d⁻¹. Saikawa et al. (2009) assessed the annual concentrations of OC and BC from biomass burning primary
emission in China using global models of chemical transport (MOZART-2) to be 1.8
and 0.35µg m⁻³. The most polluted areas were Anhui, Henan, Shandong, Jiangsu,
Liaoning, and Hunan.

969

3.3.4 Uncertainties of the emissions

The fuzziness and uncertainties of major pollutants emissions from fuel combustion 970 in China came from the uncertainties in specific-source emission factors and 971 effective consumption of bio- or fossil fuel. Frey et al. analyzed uncertainties in 972 973 emission factors and emissions of air toxic pollutants and technology dependent coal-fire power plants via bootstrap simulation method (Frey and Zhao, 2004; Frey 974 975 and Zheng, 2002). Zhao et al. estimated uncertainties in national anthropogenic pollutants emissions based on Monte Carlo simulation, and they believed activity 976 rates (e.g. fuel consumption) are not the main source of emissions uncertainties at the 977 national level (Zhao et al., 2011; Zhao et al., 2012). The uncertainties in emission 978 979 inventory can also be estimated by comparing different emission inventories for the same region and period (Ma and Van Aardenne, 2004). 980

In this study, we investigated the uncertainties of multi-pollutants emissions for 981 agricultural residue open burning using Monte Carlo Simulation. Detailed 982 983 methodology was referred to Qin and Xie (2011). We followed the assumption: a normal distribution with coefficient of variation (CV) of 30% for the official 984 statistics (e.g., crop production and GDP economic data obtained from Statistic 985 986 Yearbooks, field burning rates for agricultural straw derived from NDRC report, etc.), a normal distribution with 50% CV for open burning rates from literature 987 (BAU-I and BAU-II), and a uniform distribution with \pm 30% deviation for the rest 988 activity data (crop-to-residue ratio, dry matter fraction, and burning efficiency). 989 Regarding the emission factors, Bond et al. (2004) assumed that most particulate 990 EFs followed lognormal distributions with CV of $\pm 50\%$ for domestic EFs, and of \pm 991 150% for EFs obtained from foreign studies. Here, we applied the CV of smoke 992

993 EFs as we measured ones, which were chemical species and fuel type dependent. With randomly selected values within the respective probability density functions 994 (PDFs) of EFs and activity data for each biomass type, Monte Carlo simulation was 995 implemented for 10,000 times, and the uncertainties in national yearly 996 multi-pollutants emissions at 95% CI were obtained for all the 5 versions. 997 Afterwards, uncertainties for the average emission inventories were assessed using 998 999 the propagation of uncertainty calculation that suggested by IPCC (1997) (method in SI), and all the emission uncertainties were presented in percentage in Table 6. 1000 Emissions for water soluble aminiums and organic acids had the vast uncertainties, 1001 due to their large deviation in EFs compared with other smoke species. Besides, 1002 emissions of BAU versions were more accurate than EM versions, because of more 1003 uncertainty addition in the burning rates conversion using economic data for EM 1004 1005 versions. Otherwise, burning rates derived from NDRC report were assumed to have less uncertainty, resulting in the least uncertainties in smoke emission 1006 1007 assessments. On average of all the 5 versions, mean, 2.5th percentile, and 97.5th percentile values for smoke PM2.5 emissions in 2012 were 1005.7, 758.3, and 1008 1344.6 Gg, respectively. As to OC emissions, mean, 2.5th percentile, and 97.5th 1009 percentile values were 432.4, 327.8, and 576.4 Gg, the figure for EC was 134.2, 1010 100.9, and 187.9 Gg. Therefore, the overall propagation of uncertainties for smoke 1011 PM_{2.5}, OC, and EC at 95% CI was (-24.6%, 33.7%), (-24.4%, 33.5%), and (-24.2%, 1012 33.3%), respectively. The uncertainties for OC and EC emissions were much less 1013 than the work of Qin and Xie (2011), in which emission and uncertainties were 1014 266.7 Gg (-55.9%, 96.1%) for OC and 66. 9 Gg (-53.9%, 92.6%) for EC in 2005. 1015

- 1016 **3.4 Health and health-related economic impacts**
- 1017 3.4.1 Carcinogenic risk

1018 Calculated CR_{SPM} for smoke $PM_{2.5}$ from wheat, corn, rice, cotton, and soybean 1019 straw burning were 5.3×10^{-6} , 3.8×10^{-6} , 2.6×10^{-6} , 0.7×10^{-6} , and 1.3×10^{-6} per µg m⁻³, 1020 respectively. And the corresponded one in million PEL was 0.2, 0.3, 0.4, 1.4, and 0.8 1021 μ g m⁻³. Wu et al. (2009) ever assessed unit risk of wood and fuel burning particles 1022 using metals merely, the results were 3.2×10^{-6} and 1.5×10^{-6} per μ g m⁻³, which were 1023 close to that in our study. In actual application, PEL of smoke particles should be 1024 bulk mass concentration of mixed aerosols.

It was noticeable that apart from Tibet and Qinghai, the flux concentration of smoke $PM_{2.5}$ among all the five emission versions in other regions far surpassed the PEL, especially the North Plain and the Central of China, exhibiting great potential inhalable cancer risk. For the health care, emission flux concentration should be constrained within the PEL of crop straw burning aerosol. Thus the critical filed burning rates can be derived to ensure risk aversion following Eq. (11):

1 $R_k \leq \frac{10^{-6} \times S_k \times h \times T_k}{\sum_j \sum_i P_{t,k,i} \times r_i \times H_{t,k,i} \times D_i \times f_i \times EF_{i,j} \times CRF_i}$ (11) 2 The conservative values of regional field burning rates from Eq. (11) were named

The conservative values of regional field burning rates from Eq. (11) were named as Carcinogenic Risk Control scenarios (CRC) and listed in Table S11 (SI), which would be instructive in emission control. Under CRC, national crop straw field burning rate was less than 3%, emissions of $PM_{2.5}$ were geographically presented in Fig. S4 (SI), and 146.3 Gg yr⁻¹ smoke $PM_{2.5}$ should be released at largest in China, the corresponded annual flux concentration of $PM_{2.5}$ was within 0.3 µg m⁻³ d⁻¹ (detailed emission inventories under CRC version see in SI).

1039 3.4.2 Health impacts

Regional health impacts from acute exposure of agricultural residue burning aerosol 1040 were assessed using average daily flux concentrations of smoke PM_{2.5}, the result was 1041 tabulated in Table S12 (SI). The impacts from smoke PM_{2.5} exposure were severest 1042 1043 in Jiangsu, Shandong, and Henan province, where annual premature mortality was over one thousand. Overall, China suffered from 7836 (95% CI: 3232, 12362) 1044 premature death, 31181 (95% CI: 21145, 40881) respiratory hospital admissions, 1045 29520 (95% CI: 12873, 45602) cardiovascular hospital admissions, and 7267237 (95% 1046 CI: 2961487, 1130784) chronic bronchitis related to agricultural fire smoke in 2012 1047 from Table 7. According to national health statistical reports (NHFPC, 2013), the 1048

1049 hospital admission due to respiratory and cardiovascular disease was 5071523 in China in 2012, and smoke $PM_{2.5}$ exposure might contribute ~1.2% of the hospital 1050 1051 admissions from this study. Saikawa et al. (2009) ever reported 70000 premature deaths in China and an additional 30000 deaths globally due to OC, EC, and sulfate 1052 exposure that were primarily emitted from biofuel combustion in China in 2000, 1053 1054 however, the results should be overestimated not only in the exaggerated pollutant 1055 emissions but also in the iterative operations of respective species induced mortality, besides, the exposure-response coefficient β and incidence rate he applied from Pope 1056 et al. (2002) and WHO (2000) were higher than the practical values from local 1057 research (Cao et al., 2012; Chen et al., 2011; Hou et al., 2012). From Table 7, under 1058 1059 CRC version, over 92 % mortality and morbidity can be avoided.

1060 **3.4.3 Health-related economic losses**

Health-related total economic losses from straw open burning smoke PM2.5 exposure 1061 were assessed to be 8822.4 (95% CI: 3574.4, 13034.2) million US\$ on average from 1062 1063 Table 8, accounting for 0.1% of the total GDP in 2012, and detailed regional economic losses were listed in Table S13. Economic losses from premature death 1064 contributed about 17% of total losses, and loss from chronic bronchitis dominated. 1065 Hou et al. (2012) ever estimated 106.5 billion US\$ lost due to ambient PM_{10} 1066 exposure in China in 2009; even a severe haze episode (PM_{2.5} be focused on) in 1067 1068 January 2013 may cause 690 premature death and 253.8 million US\$ loss in Beijing, and source-specification analysis stressed the emission from biomass burning (Yang 1069 et al., 2015; Gao et al., 2015). It was obvious that smoke $PM_{2.5}$ contributed a 1070 noticeable damage to public health and social welfare. According to CRC version 1071 estimation, the carcinogenic risk control policy can save over 92 % of the economic 1072 1073 loss.

1074 **4 Conclusion**

1075 Detailed chemical compositions of smoke aerosol from five major agricultural

straws burning were characterized using an aerosol chamber system. And
corresponded emission factors for particulate OC-EC, WSI, WSOA, WSA, PAHs,
Phenols, and THM in smoke PM_{2.5} and PM_{1.0} were established.

1079 Permissible exposure limits (PEL) of the smoke particles were assessed for 1080 carcinogenic risk concern based on selected hazard pollutants including PAHs and 1081 THM in smoke $PM_{2.5}$. Daily exposure concentration should be constrained within 1082 0.2, 0.3, 0.4, 1.4, and 0.8 μ g m⁻³ for wheat, corn, rice, cotton, and soybean straw, 1083 respectively.

Emission inventories of primary particulate pollutants from agricultural field 1084 burning in 2012 were estimated based on BAU-I, BAU-II, EM-I, EM-II, and NDRC 1085 scenarios, which were further allocated into different regions at summer and autumn 1086 open burning periods. The estimated total emissions were 1005.7 Gg PM_{2.5} (95%CI: 1087 1088 -24.6% , 33.7%), 901.4 Gg PM_{1.0} (95%CI: -24.4%, 33.5%), 432.4 Gg OC (95%CI: -24.2%, 33.5%), 134.2 Gg EC (95%CI: -24.8%, 34.0%), 249.8 Gg WSI (95%CI: 1089 -25.4%, 34.9%), 25.1 Gg WSOA (95%CI: -33.3%, 41.4%), 5.8 Gg WSA (95%CI: 1090 -30.1%, 38.5%), 8.7 Gg THM (95%CI: -26.6%, 35.6%), 0.5 Gg PAHs (95%CI: 1091 -26.0%, 34.9%), and 2.7 Gg Phenols (95%CI: -26.1%, 35.1%), respectively. The 1092 spatial and temporal distributions of the five versions have similar characters that 1093 echo to the agricultural fires sites from satellite remote sensing. Less than 25 % of 1094 the emissions were released from summer field burnings that were mainly 1095 contributed by the North Plain and the Central of China. Flux concentrations of 1096 annual smoke PM2.5 that were calculated using box-model method based on five 1097 versions all exceed the PEL. From assessment of health impacts and health-related 1098 economic losses due to smoke PM2.5 short-term exposure, China suffered from 7836 1099 (95%CI: 3232, 12362) premature mortality and 7267237 (95% CI: 2961487, 1100 1130784) chronic bronchitis in 2012, which led to 8822.4 (95%CI: 3574.4, 13034.2) 1101 1102 million US\$, or 0.1 % of the total GDP losses.

Percentage of open burned crop straws at post-harvest period should cut down to less than 3% to ensure risk aversion from carcinogenicity, especially the North Plain and the Northeast, where the emissions should decease at least by 94% to meet the

1106 PEL. And by applying such emission control policy, over 92% of the mortality and

1107 morbidity attributed to agricultural fire smoke $PM_{2.5}$ can be avoided in China.

1108 Supplementary material related to this article is available online at:

Acknowledgment. This work is supported by National Natural Science Foundation of
China (No. 21190053, 21177025), Cyrus Tang Foundation (No. CTF-FD2014001),
Shanghai Science and Technology Commission of Shanghai Municipality (No.
13XD1400700, 12DJ1400100), Priority fields for Ph.D. Programs Foundation of
Ministry of Education of China (No. 20110071130003) and Strategic Priority
Research Program of the Chinese Academy of Sciences (Grant No. XDB05010200).

1115

1116 Ackerman, A. S.: Reduction of Tropical Cloudiness by Soot, Science, 5468, 1042-1047, 2000.

1117 Adams, P. J., Seinfeld, J. H. and Koch, D. M.: Global concentrations of tropospheric sulfate, nitrate,

References:

and ammonium aerosol simulated in a general circulation model, J. Geophys Res, D11:13791-13823,

1119 1999.

1120 Akagi, S. K., Yokelson, R. J., Wiedin myer, C., Alvarado, M. J., Reid, J. S., Karl, T., Crounse, J. D. and

Wennberg, P. O.: Emission factors for open and domestic biomass burning for use in atmosphericmodels, Atmos Chem. Phys., 9, 4039-4072, 2011.

Amdur, M. O. and Chen, L. C.: Furnace-Generated Acid Aerosols: Speciation and Pulmonary Effects,
Environ. Health Persp., 79, 147-150, 1989.

Andreae, M. O. and Gelencs ér, A.: Black carbon or brown carbon? The nature of light-absorbing
carbonaceous aerosols, Atmos Chem. Phys., 10, 3131-3148, 2006.

Andreae, M. O. and Merlet, P.: Emission of trace gases and aerosols from biomass burning, GlobalBiogeochem Cy., 4, 955-966, 2001.

1129 Andreae, M. O., Andreae, T. W., Annegarn, H., Beer, J., Cachier, H., le Canut, P., Elbert, W.,

Maenhaut, W., Salma, I., Wienhold, F. G., and Zenke, T. :, Airborne studies of aerosol emissions from
savanna fires in southern Africa: 2. Aerosol chemical composition, J. Geophys Res., D24, 32119-32128,

1132 1998.

1133 Araujo, J. A., Barajas, B., Kleinman, M., Wang, X., Bennett, B. J., Gong, K. W. Navab, M., Harkema,

J., Sioutas, C., Lusis, A. J., and Nel, A. E.: A mbient particulate pollutants in the ultrafine range promote
early atherosclerosis and systemic oxidative stress, Circ Res., 5, 589-596, 2008.

Arey, J. and Atkinson, R.: Photochemical reactions of PAHs in the atmosphere, PAHs: An
Ecotoxicological Persp., 47- 63, doi: 10.1002/0470867132.ch4, 2003.

1138 Arora, P. and Jain, S.: Estimation of Organic and Elemental Carbon Emitted from Wood Burning in

1139 Traditional and Improved Cookstoves Using Controlled Cooking Test, Environ. Sci. Technol., 6,1140 3958-3965, 2015.

- Aunan, K. and Pan, X.: Exposure-response functions for health effects of ambient air pollution
 applicable for China-a meta-analysis, Sci. Total Environ., 329, 3-16, 2004.
- 1143 Aurell, J., Gullett, B. K. and Tabor, D.: Emissions from southeastern U.S. Grasslands and pine

1144 savannas: Comparison of aerial and ground field measurements with laboratory burns, Atmos. Environ.,

1145 111, 170-178, 2015.

- Bell, R. W. and Hipfner, J. C.: Airborne Hexavalent Chromium in Southwestern Ontario, J. Air WasteManage, 8, 905-910, 1997.
- Berndt, T. and Böge, O.: Formation of phenol and carbonyls from the atmospheric reaction of OH
 radicals with benzene, Phys. Chem. Chem. Phys., 10, 1205-1214, doi:10.1039/B514148F, 2006.
- 1150 Bølling, A. K., Pagels, J., Yttri, K. E., Barregard, L., Sallsten, G., Schwarze, P. E. and Boman, C.:
- 1151 Health effects of residential wood smoke particles: the importance of combustion conditions and
- 1152 physicochemical particle properties, Part. Fibre Toxicol., 29, doi:10.1186/1743-8977-6-29, 2009.
- Bond, T. C.: A technology-based global inventory of black and organic carbon emissions from
 combustion, J. Geophys Res., 109, D14203, doi:10.1029/2003JD003697, 2004.
- 1155 Bond, T. C., Doherty, S. J., Fahey, D. W., Forster, P. M., Berntsen, T., DeAngelo, B. J. Flanner, M. G.,
- 1156 Ghan, S., Kärcher, B., Koch, D., Kinne, S., Kondo, Y., Quinn, P. K., Sarofim, M. C., Schultz, M. G.,
- 1157 Schulz, M., Venkataraman, C., Zhang, H., Zhang, S., Bellouin, N., Guttikunda, S. K., Hopke, P. K.,
- 1158 Jacobson, M. Z., Kaiser, J. W., Klimont, Z., Lohmann, U., Schwarz, J., PShindell, D., Storelvmo, T.,
- Warren, S. G., and Zender, C. S.: Bounding the role of black carbon in the climate system: Ascientific assessment, J. Geophys Res: Atmos, 11, 5380-5552, 2013.
- Bruce, R. M., Santodonato, J. and Neal, M. W.: Summary Review of the Health Effects AssociatedWith Phenol, Toxicol Ind. Health, 4, 535-568, 1987.
- Buha, J., Mueller, N., Nowack, B., Ulrich, A., Losert, S. and Wang, J.: Physical and Chemical
 Characterization of Fly Ashes from Swiss Waste Incineration Plants and Determination of the Ash
 Fraction in the Nanometer Range, Environ. Sci. Technol., 9, 4765-4773, 2014.
- Burkart, K., Nehls, I., Win, T. and Endlicher, W.: The carcinogenic risk and variability of
 particulate-bound polycyclic aromatic hydrocarbons with consideration of meteorological conditions,
 Air Quality, Atmos. Health, 1, 27-38, 2013.
- 1169 Bzdek, B. R., Ridge, D. P. and Johnston, M. V.: A mine reactivity with charged sulfuric acid clusters,
- 1170 Atmos. Chem. Phys., 16, 8735-8743, 2011.
- Bzdek, B. R., Ridge, D. P. and Johnston, M. V., Amine exchange into ammonium bisulfate andammonium nitrate nuclei, Atmos Chem Phys, 8:3495-3503, 2010.
- 1173 Cao, G., Zhang, X. and Zheng, F.: Inventory of black carbon and organic carbon emissions from China,
 1174 Atmos. Environ., 34, 6516-6527, 2006.
- 1175 Cao, G., Zhang, X., Gong, S., An, X. and Wang, Y.: Emission inventories of primary particles and
 1176 pollutant gases for China, Chinese Sci. Bull, 8, 781-788, 2011.
- 1177 Cao, G., Zhang, X., Wang, D. and Zheng, F.: Inventory of Emissions of Pollutants from Open Burning
 1178 Crop Residues, J. Agro-Environ. Sci., 4, 800-804, 2005.
- 1179 Cao, J., Xu, H., Xu, Q., Chen, B. and Kan, H.: Fine particulate matter constituents and 1180 cardiopulmonary mortality in a heavily polluted Chinese city, Environ. Health Persp., 3, 373, 2012.
- 1181 Carlton, A. G., Turpin, B. J., Lim, H., Altieri, K. E. and Seitzinger, S.: Link between isoprene and
- secondary organic aerosol (SOA): Pyruvic acid oxidation yields low volatility organic acids in clouds,
 Geophys. Res. Lett., Lo6822, doi:10.1029/2005GL025374, 2006.
- 1184 Cermak, J. and Knutti, R.: Beijing Olympics as an aerosol field experiment, Geophys. Res. Lett., 36,
 1185 L10806, doi:10.1029/2009GL038572, 2009.
- 1186 Chakrabarty, R. K., Moosmüller, H., Chen, L. W. A., Lewis, K., Arnott, W. P., Mazzoleni, C., Dubey,
- M. K., Wold, C. E., Hao, W. M., and Kreidenweis, S. M.: Brown carbon in tar balls from smoldering
 biomass combustion, Atmos. Chem. Phys., 13, 6363-6370, 2010.
- 1189 Chan, M. N., Choi, M. Y., Ng, N. L. and Chan, C. K.: Hygroscopicity of Water-Soluble Organic

- 1190 Compounds in Atmospheric Aerosols: Amino Acids and Biomass Burning Derived Organic Species,
- 1191 Environ. Sci. Technol., 6, 1555-1562, 2005.
- Chen, H., Hu, D., Wang, L., Mellouki, A. and Chen, J.: Modification in light absorption cross section
 of laboratory-generated black carbon-brown carbon particles upon surface reaction and hydration,
- 1194 Atmos. Environ., 116, 253-261, 2015.
- 1195 Chen, R., Li, Y., Ma, Y., Pan, G., Zeng, G., Xu, X., Chen, B. and Kan, H.: Coarse particles and
- mortality in three Chinese cities: The China Air Pollution and Health Effects Study (CAPES), Sci.
 Total Environ., 23, 4934-4938, 2011.
- 1198 Cheng, Y., Ho, K. F., Lee, S. C. and Law, S. W.: Seasonal and diurnal variations of $PM_{1.0}$, $PM_{2.5}$ and 1199 PM_{10} in the roadside environment of Hong Kong, China Particuology, 06:312-315, 2006.
- 1200 China Ministry of Health (CMH): China statistical yearbook of public health, Peking Union Medical
 1201 College Press, 172–189, 2009. (*In Chinese*)
- 1202 Christopher, S. A., Chou, J., Zhang, J., Li, X., Berendes, T. and Welch, R. M.: Shortwave direct1203 radiative forcing of biomass burning aerosols estimated using VIRS and CERES data, Geophys. Res.
- 1204 Lett., 15, 2197-2200, 2000.
- 1205 Clarke, A., McNaughton, C., Kapustin, V., Shinozuka, Y., Howell, S., Dibb, J., Zhou, J., Anderson, B.,
- Brekhovskikh, V., Turner, H. and Pinkerton, M.: Biomass burning and pollution aerosol over North
 America: Organic components and their influence on spectral optical properties and humidification
 response, J. Geophys. Res., D12, doi:10.1029/2006JD007777, 2007.
- 1209 Cwiertny, D. M., Baltrusaitis, J., Hunter, G. J., Laskin, A., Scherer, M. M. and Grassian, V. H.:
 1210 Characterization and acid-mobilization study of iron-containing mineral dust source materials, J.
 1211 Geophys. Res.: Atmos., D5, doi:10.1029/2007JD009332, 2008.
- 1212 Daize, H.: The Utilizing Status and Prospects of the Crop Straw Resources in China, Resource1213 Development & Market:12, 2000.
- 1214 Davidson, C. I., Phalen, R. F. and Solomon, P. A.: Airborne Particulate Matter and Human Health: A
 1215 Review, Aerosol Sci. Tech., 8, 737-749, 2005.
- 1216 Delfino, R. J., Sioutas, C. and Malik, S.: Potential role of ultrafine particles in associations between
 1217 airborne particle mass and cardiovascular health, Environ. Health Perspect., 8, 934-946, 2005.
- 1218 Dentener, F. J., Carmichael, G. R., Zhang, Y., Lelieveld, J. and Crutzen, P. J.: Role of mineral aerosol
 1219 as a reactive surface in the global troposphere, J. Geophys. Res., D17, 22869-22889, doi:
 10.1029/96JD01818, 1996.
- Dhammapala, R., Claiborn, C., Corkill, J. and Gullett, B.: Particulate emissions from wheat and
 Kentucky bluegrass stubble burning in eastern Washington and northern Idaho, Atmos. Environ., 6,
 1007-1015, 2006.
- 1224 Dhammapala, R., Claiborn, C., Jimenez, J., Corkill, J., Gullett, B., Simpson, C. and Paulsen, M.:
 1225 Emission factors of PAHs, methoxyphenols, levoglucosan, elemental carbon and organic carbon from
- simulated wheat and Kentucky bluegrass stubble burns, Atmos. Environ., 12, 2660-2669, 2007a.
- 1227 Dhammapala, R., Claiborn, C., Simpson, C. and Jimenez, J.: Emission factors from wheat and
 1228 Kentucky bluegrass stubble burning: Comparison of field and simulated burn experiments, Atmos.
 1229 Environ., 7, 1512-1520, 2007b.
- 1230 Dusek, U., Frank, G. P., Hildebrandt, L., Curtius, J., Schneider, J., Walter, S., Chand, D., Drewnick, F.,
- Hings, S., Jung, D.: Size matters more than chemistry for cloud-nucleating ability of aerosol particles,
 Science, 5778, 1375-1378, 2006.
- 1233 Echalar, F., Gaudichet, A., Cachier, H. and Artaxo, P.: Aerosol emissions by tropical forest and

- savanna biomass burning: Characteristic trace elements and fluxes, Geophys. Res. Lett., 22, 3039-3042,
 doi:10.1029/95GL03170, 1995.
- 1236 Engelhart, G. J., Hennigan, C. J., Miracolo, M. A., Robinson, A. L. and Pandis, S. N.: Cloud
 1237 condensation nuclei activity of fresh primary and aged biomass burning aerosol, Atmos. Chem. Phys.,
 1238 15, 7285-7293, doi:10.5194/acp- 12- 7285- 2012, 2012.

Falkovich, A. H., E., R. G., G., S., Y., R., Maenhaut, W. and Artaxo, P.: Low molecular weight organic
acids in aerosol particles from Rondônia, Brazil, during the biomass-burning, transition and wet
periods, Atmos. Chem. Phys., 5, 781-797, doi:10.5194/acp-5-781-2005, 2005.

- Frey, H. C. and Zhao, Y.: Quantification of Variability and Uncertainty for Air Toxic Emission
 Inventories with Censored Emission Factor Data, Environ. Sci. Technol., 22, 6094-6100, 2004.
- Frey, H. and Zheng, J.: Quantification of variability and uncertainty in air pollutant emission
 inventories: method and case study for utility NOx emissions, J. Air Waste Manag. Assoc., 9,
 1083-1095, 2002.
- Fu, H., Zhang, M., Li, W., Chen, J., Wang, L., Quan, X. and Wang, W.: Morphology, composition and
 mixing state of individual carbonaceous aerosol in urban Shanghai, Atmos. Chem. Phys., 2, 693-707,
- 1249 2012.
- Galarneau, E.: Source specificity and atmospheric processing of airborne PAHs: Implications forsource apportionment, Atmos. Environ., 35, 8139-8149, 2008.
- 1252 Gao, M., Guttikunda, S. K., Carmichael, G. R., Wang, Y., Liu, Z., Stanier, C. O., Saide, P. E. and Yu,
- M.: Health impacts and economic losses assessment of the 2013 severe haze event in Beijing area, Sci.
 Total Environ., 511, 553-561, 2015.
- Gao, S., Hegg, D. A., Hobbs, P. V., Kirchstetter, T. W., Magi, B. I. and Sadilek, M.: Water-soluble
 organic components in aerosols associated with savanna fires in southern Africa: Identification,
 evolution, and distribution, J. Geophys.l Res.: Atmos., D13, doi:10.1029/2002JD002324, 2003.
- 1258 Ge, X., Wexler, A. S. and Clegg, S. L.: Atmospheric amines-Part I. A review, Atmos. Environ., 3,
 1259 524-546, 2011.
- 1260 Ghorai, S., Wang, B., Tivanski, A. and Laskin, A.: Hygroscopic Properties of Internally Mixed
 1261 Particles Composed of NaCl and Water-Soluble Organic Acids, Environ. Sci. Technol., doi:
 1262 10.1021/es404727u, 2014.
- Giordano, M., Espinoza, C. and Asa-Awuku, A.: Experimentally measured morphology of biomass
 burning aerosol and its impacts on CCN ability, Atmos. Chem. Phys., 4, 1807-1821, 2015.
- 1265 Grieshop, A. P., Logue, J. M., Donahue, N. M. and Robinson, A. L.: Laboratory investigation of 1266 photochemical oxidation of organic aerosol from wood fires 1: measurement and simulation of organic
- 1267 aerosol evolution, Atmos. Chem. Phys., 4, 1263-1277, 2009.
- 1268 Guttikunda, S. K. and Kopakka, R. V.: Source emissions and health impacts of urban air pollution in
 1269 Hyderabad, India, Air Quality, Atmos. & Health, 2, 195-207, 2014.
- 1270 Han, Y. M., Chen, L., Huang, R., Chow, J. C., Watson, J. G., Ni, H. Y., Liu, S. X., Fung, K. K., Shen,
- 1271 Z. X. and Wei, C.: Carbonaceous aerosols in megacity Xi'an, China: Implications of thermal/optical
 1272 protocols comparison, Atmos. Environ., 132, 58-68, 2016.
- 1273 Han, Y. M., Lee, S. C., Cao, J. J., Ho, K. F. and An, Z. S.: Spatial distribution and seasonal variation of
- 1274 char-EC and soot-EC in the atmosphere over China, Atmos. Environ., 38, 6066-6073, 2009.
- 1275 Han, Y., Cao, J., Chow, J. C., Watson, J. G., An, Z., Jin, Z., Fung, K. and Liu, S.: Evaluation of the
- 1276 thermal/optical reflectance method for discrimination between char- and soot-EC, Chemosphere, 4,
- **1277** 569-574, 2007.

- 1278 Hayashi, K., Ono, K., Kajiura, M., Sudo, S., Yonemura, S., Fushimi, A., Saitoh, K., Fujitani, Y. and
- 1279 Tanabe, K.: Trace gas and particle emissions from open burning of three cereal crop residues: Increase1280 in residue moistness enhances emissions of carbon monoxide, methane, and particulate organic carbon,
- 1281 Atmos. Environ., 95, 36-44, 2014.

Hays, M. D., Fine, P. M., Geron, C. D., Kleeman, M. J. and Gullett, B. K.: Open burning of agricultural
biomass: physical and chemical properties of particle-phase emissions, Atmos. Environ., 36,

- 1284 6747-6764, 2005.
- He, K., Zhao, Q., Ma, Y., Duan, F. and Yang, F.: Spatial and seasonal variability of PM_{2.5} acidity at two Chinese megacities: insights into the formation of secondary inorganic aerosols, Atmos. Chem.
 Phys. Dis., 25557-25603, doi:10.5194/acpd- 11- 25557- 2011, 2011a.
- He, M., Zheng, J., Yin, S. and Zhang, Y.: Trends, temporal and spatial characteristics, and uncertainties
 in biomass burning emissions in the Pearl River Delta, China, Atmos. Environ., 24, 4051-4059, 2011b.
- 1290 Ho, K. F., Ho, S. S. H., Huang, R., Liu, S. X., Cao, J., Zhang, T., Chuang, H., Chan, C. S., Hu, D. and
- 1291 Tian, L.: Characteristics of water-soluble organic nitrogen in fine particulate matter in the continental

1292 area of China, Atmos. Environ., 106, 252-261, 2015.

- Hou, Q., An, X., Wang, Y., Tao, Y. and Sun, Z.: An assessment of China's PM₁₀-related health
 economic losses in 2009, Sci. Total. Environ., 61-65, 2012.
- Hu, Y., Lin, J., Zhang, S., Kong, L., Fu, H. and Chen, J.: Identification of the typical metal particles
 among haze, fog, and clear episodes in the Beijing atmosphere, Sci. Total Environ., 369-380, 2015.
- 1297 Huang, K., Zhuang, G., Lin, Y., Fu, J. S., Wang, Q., Liu, T., Zhang, R., Jiang, Y., Deng, C. and Fu, Q.:
- 1298 Typical types and formation mechanisms of haze in an Eastern Asia megacity, Shanghai, Atmos. Chem.1299 Phys., 2012a.
- Huang, K., Zhuang, G., Lin, Y., Wang, Q., Fu, J. S., Fu, Q., Liu, T. and Deng, C.: How to improve the
 air quality over megacities in China: pollution characterization and source analysis in Shanghai before,
 during, and after the 2010 World Expo, Atmos. Chem. Phys., 12, 5927-5942, 2013.
- 1303 Huang, R., Zhang, Y., Bozzetti, C., Ho, K., Cao, J., Han, Y., Daellenbach, K. R., Slowik, J. G., Platt, S.
- 1304 M., Canonaco, F., Zotter, P., Wolf, R., Pieber, S., Bruns, E., Crippa, M., Ciarelli, G., Piazzalunga, A.,
- 1305 Schnelle-Kreis, J., Zimmermann, R., An, Z., Szidat, S., Baltensperger, U., Haddad, I. and Prevot, A.:
- 1306 High secondary aerosol contribution to particulate pollution during haze events in China, Nature, 2014.
- Huang, S., Hsu, M. and Chan, C.: Effects of submicrometer particle compositions on cytokine
 production and lipid peroxidation of human bronchial epithelial cells., Environ. Health Persp., 4, 478,
- 1309 2003.
- Huang, X., Li, M., Li, J. and Song, Y.: A high-resolution emission inventory of crop burning in fields
 in China based on MODIS Thermal Anomalies/Fire products, Atmos. Environ., 9-15, 2012b.
- Huo, J., Lu, X., Wang, X., Chen, H., Ye, X., Gao, S., Gross, D. S., Chen, J. and Yang, X.: Online
 single particle analysis of chemical composition and mixing state of crop straw burning particles: from
 laboratory study to field measurement, Front Env. Sci. Eng., 2, 244-252, 2016.
- 1315 IBRD and SEPA, Cost of pollution in China: economic estimates of physical damages, 2007, pp.1316 1-128.
- 1317 IPCC, Greenhouse Gas Inventory Reference Manual: Revised 2006 IPCC Guidelines for National
 1318 Greenhouse Gas Inventories. IPCC/OECD/IES, UK. Meteorological Office, Bracknell, UK., 2007.
- 1319 IPCC, Quantifying Uncertainties in Practice, Chapter 6: Good Practice Guidance and Uncertainty
- 1919 In Ce, Quantifying Oncertaintees in Flattice, Chapter 6. Good Flattice Guidance and Oncertainty
- 1320 Management in National Greenhouse Gas Inventories: In: IES, IPCC, OECD, et al. Bracknell, UK,
- **1321** 1997.

- 1322 Janssen, N. A. H., Hoek, G., Simic-Lawson, M., Fischer, P., van Bree, L., Ten Brink, H., Keuken, M.,
- 1323 Atkinson, R., Anderson, H., Brunekreef, B. and Cassee, F.: Black Carbon as an Additional Indicator of
- 1324 the Adverse Health Effects of Airborne Particles Compared with PM_{10} and $PM_{2.5}$, Environ. Health
- 1325 Persp., 12, 1691-1699, 2011.
- Jayarathne, T., Stockwell, C. E., Yokelson, R. J., Nakao, S. and Stone, E. A.: Emissions of fine particle
 fluoride from biomass burning, Environ. Sci. Technol., 21, 12636-12644, 2014.
- 1328 Jenkins, B. M., Jones, A. D., Turn, S. Q. and Williams, R. B.: Emission factors for polycyclic aromatic
- 1329 hydrocarbons from biomass burning, Environ. Sci. Technol., 8, 2462-2469, 1996.
- 1330 Jickells, T. D., An, Z. S., Andersen, K. K., Baker, A. R., Bergametti, G., J, N. B. J., Duce, R. A.,
- Hunter, H., Mahowald, N. and Prospero, A.: Global Iron Connections Between Desert Dust, OceanBiogeochemistry, and Climate, Science, 308, 67-71, 2005.
- Kennedy, I. M.: The health effects of combustion-generated aerosols, P. Combust Inst, 2, 2757-2770,2007.
- 1335 Kim, K., Jahan, S. A., Kabir, E. and Brown, R. J.: A review of airborne polycyclic aromatic
 1336 hydrocarbons (PAHs) and their human health effects, Environ Int., 71-80, 2013.
- 1337 Kong, L., Yang, Y., Zhang, S., Zhao, X., Du, H., Fu, H., Zhang, S., Cheng, T., Yang, X. and Chen, J.:
- Observations of linear dependence between sulfate and nitrate in atmospheric particles, J. Geophys.
 Res.: Atmos., 1, 341-361, doi:10.1002/2013JD020222, 2014.
- Koopmans, A. and Koppejan, J.: Agricultural and forest residue-generation, utilization and availability,
 Modern Applications of Biomass Energy, 1997.
- 1342 Korenaga, T., Liu, X. and Huang, Z.: The influence of moisture content on polycyclic aromatic
 1343 hydrocarbons emission during rice straw burning, Chemosphere-Global Change Science, 1, 117-122,
 1344 2001
- Kundu, S., Kawamura, K., Andreae, T. W., Hoffer, A. and Andreae, M. O.: Molecular distributions of
 dicarboxylic acids, ketocarboxylic acids and α-dicarbonyls in biomass burning aerosols: implications
- for photochemical production and degradation in smoke layers, Atmos. Chem. Phys., 5, 2209-2225,2010.
- Laskin, A., Laskin, J. and Nizkorodov, S. A.: Chemistry of atmospheric brown carbon, Chem. Rev., 10,
 4335-4382, 2015.
- Lavanchy, V. M. H., G Ggeler, H. W., Nyeki, S. and Baltensperger, U.: Elemental carbon (EC) and
 black carbon (BC) measurements with a thermal method and an aethalometer at the high-alpine
 research station Jungfraujoch, Atmos. Environ., 17, 2759-2769, 1999.
- Lee, A. K. Y., Willis, M. D., Healy, R. M., Wang, J. M., Jeong, C. H., Wenger, J. C., Evans, G. J. and
 Abbatt, J.: Single particle characterization of biomass burning organic aerosol (BBOA): evidence for
 non-uniform mixing of high molecular weight organics and potassium, Atmos. Chem. Phys. Dis., 22,
- 1357 32157-32183, 2015.
 1358 Lee, D. and Wexler, A. S.: Atmospheric amines-Part III: Photochemistry and toxicity, Atmos. Environ.,
- **1359** 95-103, 2013.
- 1360 Lee, R. G. M., Coleman, P., Jones, J. L., Jones, K. C. and Lohmann, R.: Emission Factors and
- 1361 Importance of PCDD/Fs, PCBs, PCNs, PAHs and PM₁₀ from the Domestic Burning of Coal and Wood
- 1362 in the U.K., Environ. Sci. Technol., 6, 1436-1447, 2005.
- 1363 Leng, C., Zhang, Q., Zhang, D., Xu, C., Cheng, T., Zhang, R., Tao, J., Chen, J., Zha, S. and Zhang, Y.:
- 1364 Variations of cloud condensation nuclei (CCN) and aerosol activity during fog-haze episode: a case
- 1365 study from Shanghai, Atmos. Chem. Phys., 22, 12499-12512, doi:10.5194/acp-14-12499-2014, 2014.

- 1366 Lewis, K. A., Arnott, W. P., Moosmuller, H., Chakrabarty, R. K., Carrico, C. M., Kreidenweis, S. M.,
- 1367 Day, D. E., Malm, W., Laskin, A., Jimenez, J., Ulbrich, I., Huffman, J., Onasch, T., Trimborn, A., Liu,
- L. and Mishchenko, M.: Reduction in biomass burning aerosol light absorption upon humidification:roles of inorganically-induced hygroscopicity, particle collapse, and photoacoustic heat and mass
- 1370 transfer, Atmos. Chem. Phys., 9, 8949-8966, 2009.
- Levin, E. J. T., McMeeking, G. R., Carrico, C. M., Mack, L. E., Kreidenweis, S. M., Wold, C. E.,
 Moosmüller, H., Arnott, W., Hao, W., Collett, J. and Malm, W.: Biomass burning smoke aerosol
 properties measured during Fire Laboratory at Missoula Experiments (FLAME), J. Geophys Res,
 D18210, doi:10.1029/2009JD013601, 2010.
- Li, C., Ma, Z., Chen, J., Wang, X., Ye, X., Wang, L., Yang, X., Kan, H., Donaldson, D. and Mellouki,
 A.: Evolution of biomass burning smoke particles in the dark, Atmos. Environ., 120, 244-252, 2015.
- Li, C., Hu, Y., Chen, J., Ma, Z., Ye, X., Yang, X., Wang, L., Wang, X. and Mellouki, A.:
 Physiochemical properties of carbonaceous aerosol from agricultural residue burning: Density,
 volatility, and hygroscopicity, Atmos. Environ., 2016.
- Li, J., Pósfai, M., Hobbs, P. V. and Buseck, P. R.: Individual aerosol particles from biomass burning in
 southern Africa: 2, Compositions and aging of inorganic particles, J. Geophys. Res.: Atmos.,
 (1984-2012), D13, doi:10.1029/2002JD002310, 2003.
- Li, X., Wang, S., Duan, L., Hao, J., Li, C., Chen, Y. and Yang, L.: Particulate and trace gas emissions
 from open burning of wheat straw and corn stover in China, Environ. Sci. Techno., 17, 6052-6058,
 2007.
- Lima, A. L. C., Farrington, J. W. and Reddy, C. M.: Combustion-Derived Polycyclic Aromatic
 Hydrocarbons in the Environment-A Review, Environ. Forensics., 2, 109-131, 2005.
- Lin, J., Nielsen, C. P., Zhao, Y., Lei, Y., Liu, Y. and McElroy, M. B.: Recent changes in particulate air
 pollution over China observed from space and the ground: effectiveness of emission control, Environ.
 Sci. Technol., 20, 7771-7776, 2010.
- Lin, P., Aiona, P. K., Li, Y., Shiraiwa, M., Laskin, J., Nizkorodov, S. A. and Laskin, A.: Molecular
 Characterization of Brown Carbon in Biomass Burning Aerosol Particles, Environ. Sci. Technol., 21,
 11815-11824, 2016.
- Liu, M., Song, Y., Yao, H., Kang, Y., Li, M., Huang, X. and Hu, M.: Estimating emissions from
 agricultural fires in the North China Plain based on MODIS fire radiative power, Atmos. Environ.,
 326-334, 2015.
- Liu, Q. and Bei, Y.: Impacts of crystal metal on secondary aliphatic amine aerosol formation during
 dust storm episodes in Beijing, Atmos. Environ., 227-234, 2016.
- Lobert, J. M., Scharffe, D. H., Hao, W. M. and Crutzen, P. J.: Importance of biomass burning in the
 atmospheric budgets of nitrogen-containing gases, 6284, 552-554, 1990.
- 1401 Lu, Z., Zhang, Q. and Streets, D. G.: Sulfur dioxide and primary carbonaceous aerosol emissions in
- 1402 China and India, 1996-2010, Atmos. Chem. Phys., 11, 9839-9864, doi:10.5194/acp-11-9839-2011,
 1403 2011.
- Ma, J. and Van Aardenne, J. A.: Impact of different emission inventories on simulated tropospheric
 ozone over China: a regional chemical transport model evaluation, Atmos. Chem. Phys., 4, 877-887,
 2004.
- 1407 May, A. A., McMeeking, G. R., Lee, T., Taylor, J. W., Craven, J. S., Burling, I., Sullivan, A. P., Akagi,
- 1408 S., Collett, J., Flynn, M., Coe, H., Urbanski, S., Seinfeld, J., Yokelson, R. and Kreidenweis, S.: Aerosol
- 1409 emissions from prescribed fires in the United States: A synthesis of laboratory and aircraft

- 1410 measurements, J. Geophys. Res.: Atmos., 20, 11826-11849, doi:10.1002/2014JD021848, 2014.
- Meskhidze, N.: Dust and pollution: A recipe for enhanced ocean fertilization? J. Geophys. Res, D3,
 doi:10.1029/2004JD005082, 2005.
- 1112 doi.10.102)/20013D003002, 2003.
- 1413 Mikhailov, E. F., Vlasenko, S. S., Podgorny, I. A., Ramanathan, V. and Corrigan, C. E.: Optical
- properties of soot-water drop agglomerates: An experimental study, J. Geophys. Re., 111, D07209,
 doi:10.1029/2005JD006389, 2006.
- 1416 Moreno, T., Karanasiou, A., Amato, F., Lucarelli, F., Nava, S., Calzolai, G., Chiari, M., Coz, E.,
- 1417 Artinano, B., Lumbreras, J., Borge, R., Boldo, R., Linares, C., Alastursy, A., Querol, X. and Gibbons,
- 1418 X.: Daily and hourly sourcing of metallic and mineral dust in urban air contaminated by traffic and
- 1419 coal-burning emissions, Atmos. Environ., 68, 33-44, 2013.
- 1420 NBSC, China Statistical Yearbook 2013: China Statistics Press Beijing, China, 2013.
- 1421 NHFPC, National Health and Family Planning Yearbook: Peking Union Medical College Press, 2013,1422 p. 415.
- 1423 Oanh, N. T. K., Ly, B. T., Tipayarom, D., Manandhar, B. R., Prapat, P., Simpson, C. D. and Liu, L. S.:
- 1424 Characterization of particulate matter emission from open burning of rice straw, Atmos. Environ., 2,
 1425 493-502, 2011.
- Ostro, B. and Chestnut, L: Assessing the health benefits of reducing particulate matter air pollution in
 the United States, Environ. Res., 2, 94-106, 1998.
- Pope III, C. A., Burnett, R. T., Thun, M. J., Calle, E. E., Krewski, D., Ito, K. and Thurston, G. D.: Lung
 cancer, cardiopulmonary mortality, and long-term exposure to fine particulate air pollution, Jama, 9,
 1132-1141, 2002.
- 1431 Pope, C. A., Burnett, R. T., Thurston, G. D., Thun, M. J., Calle, E. E., Krewski, D. and Godleski, J. J.:
- 1432 Cardiovascular mortality and long-term exposure to particulate air pollution epidemiological evidence
 1433 of general pathophysiological pathways of disease, Circulation, 1, 71-77, 2004.
- Pósfai, M.: Atmospheric tar balls: Particles from biomass and biofuel burning, J. Geophys. Res., 109,
 D06213, doi:10.1029/2003JD004169, 2004.
- 1436 Qin, Y. and Xie, S. D.: Spatial and temporal variation of anthropogenic black carbon emissions in
 1437 China for the period 1980-2009, Atmos. Chem. Phys., 11, 4825-4841, 2012.
- Qin, Y. and Xie, S. D.: Historical estimation of carbonaceous aerosol emissions from biomass open
 burning in China for the period 1990-2005, Environ Pollut., 12, 3316-3323, 2011.
- Qiu, C. and Zhang, R.: Physiochemical Properties of Alkylaminium Sulfates: Hygroscopicity,
 Thermostability, and Density, Environ. Sci. Technol., 8, 4474-4480, 2012.
- Qiu, C., Wang, L., Lal, V., Khalizov, A. F. and Zhang, R.: Heterogeneous Reactions of Alkylamines
 with Ammonium Sulfate and Ammonium Bisulfate, Environ. Sci. Technol., 11, 4748-4755, 2011.
- 1444 Ram, K., Sarin, M. M. and Tripathi, S. N.: Temporal Trends in Atmospheric PM_{2.5}, PM₁₀, Elemental
- 1445 Carbon, Organic Carbon, Water-Soluble Organic Carbon, and Optical Properties: Impact of Biomass
 1446 Burning Emissions in The Indo-Gangetic Plain, Environ. Sci. Technol., 2, 686-695, 2011.
- 1447 Ram, K. and Sarin, M. M.: Day-night variability of EC, OC, WSOC and inorganic ions in urban
- 1448 environment of Indo-Gangetic Plain: implications to secondary aerosol formation, Atmos. Environ., 2,
 1449 460-468, 2011.
- 1450 Reddy, M. S. and Venkataraman, C.: Atmospheric optical and radiative effects of anthropogenic
 1451 aerosol constituents from India, Atmos. Environ., 34, 4511-4523, 2000.
- 1452 Reid, J. S., Eck, T. F., Christopher, S. A., Koppmann, R., Dubovik, O., Eleuterio, D. P., Holben, B. N.,
- 1453 Reid, E. A. and Zhang, J.: A review of biomass burning emissions part III: intensive optical properties

- 1454 of biomass burning particles, Atmos. Chem. Phys., 5, 827-849, 2005a.
- 1455 Reid, J. S., Koppmann, R., Eck, T. F. and Eleuterio, D. P.: A review of biomass burning emissions part
- 1456 II: intensive physical properties of biomass burning particles, Atmos. Chem. Phys., 3, 799-825, 2005b.
- 1457 Richter, H. and J, H.: Formation of polycyclic aromatic hydrocarbons and their growth to soot-a review1458 of chemical reaction pathways, Prog Energ Combust, 4, 565-608, 2000.
- Ripoll, A., Minguillón, M. C., Pey, J., Pérez, N., Querol, X. and Alastuey, A.: Joint analysis of continental and regional background environments in the western Mediterranean: PM₁ and PM₁₀
 concentrations and composition, Atmos. Chem. Phys., 2, 1129-1145, doi:10.5194/acp-15-1129-2015, 2015.
- Roemer, W. H. and van Wijnen, J. H.: Differences among Black Smoke, PM₁₀, and PM_{1.0} Levels at
 Urban Measurement Sites, Environ. Health Persp., 2, 151-153, 2001.
- 1465 Rose, D., Gunthe, S. S., Su, H., Garland, R. M., Yang, H., Berghof, M., Cheng, Y. F., Wehner, B.,
 1466 Achtert, P., Nowak, A., Wiedensohler, A., Takegawa, N., Kondo, Y., Hu, M., Zhang, Y., Andreae, M.
- and Poschl, U.: Cloud condensation nuclei in polluted air and biomass burning smoke near the
 mega-city Guangzhou, China-Part 2: Size-resolved aerosol chemical composition, diurnal cycles, and
 externally mixed weakly CCN-active soot particles, Atmos. Chem. Phys., 6, 2817-2836,
 doi:10.5194/acp-11-2817-2011, 2011.
- 1471 Rosenfeld, D.: Atmosphere: Aerosols, Clouds, and Climate, Science, 5778, 1323-1324, 2006.
- Safai, P. D., Raju, M. P., Budhavant, K. B., Rao, P. and Devara, P.: Long term studies on
 characteristics of black carbon aerosols over a tropical urban station Pune, India, Atmos. Res., 173-184,
 2013.
- Saffari, A., Daher, N., Samara, C., Voutsa, D., Kouras, A., Manoli, E., Karagkiozidou, O.,
 Vlachokostas, C., Moussiopoulos, N., Shafer, M., Schauer, J. and Sioutas, C.: Increased Biomass
 Burning Due to the Economic Crisis in Greece and Its Adverse Impact on Wintertime Air Quality in
 Thessaloniki, Environ. Sci. Technol., 23, 13313-13320, 2013.
- Saikawa, E., Naik, V., Horowitz, L. W., Liu, J. and Mauzerall, D. L.: Present and potential future
 contributions of sulfate, black and organic carbon aerosols from China to global air quality, premature
 mortality and radiative forcing, Atmos. Environ., 17, 2814-2822, 2009.
- Samy, S. and Hays, M. D.: Quantitative LC-MS for water-soluble heterocyclic amines in fine aerosols
 (PM_{2.5}) at Duke Forest, USA, Atmos. Environ., 77-80, 2013.
- Santodonato, J.: Review of the estrogenic and antiestrogenic activity of polycyclic aromatic
 hydrocarbons: relationship to carcinogenicity, Chemosphere, 4, 835-848, 1997.
- Schade, G. W. and Crutzen, P. J.: Emission of aliphatic amines from animal husbandry and their
 reactions: Potential source of N₂O and HCN, J. Atmos. Chem., 3, 319-346, 1995.
- 1488 Schauer, J. J., Kleeman, M. J., Cass, G. R. and Simoneit, B. R. T.: Measurement of Emissions from Air
- Pollution Sources. 3. C1–C29 Organic Compounds from Fireplace Combustion of Wood, Environ. Sci.
 Technol., 9, 1716-1728, 2001.
- 1491 Schlesinger, R. B.: Comparative deposition of inhaled aerosols in experimental animals and humans: a
- review, Journal of Toxicology and Environmental Health, Part A Current Issues, 2, 197-214, 1985.
- Seinfeld, J. H. and Pandis, S. N.: Atmospheric chemistry and physics: from air pollution to climatechange: John Wiley & Sons, 2012.
- 1495 Sen, A., Mandal, T. K., Sharma, S. K., Saxena, M., Gupta, N. C., Gautam, R., Gupta, A., Gill, T., Rani,
- 1496 S., Saud, T., Singh, D. and Gadi, R.: Chemical properties of emission from biomass fuels used in the
- rural sector of the western region of India, Atmos. Environ., 411-424, 2014.

- Shi, Y., Chen, J., Hu, D., Wang, L., Yang, X. and Wang, X.: Airborne submicron particulate (PM₁)
 pollution in Shanghai, China: Chemical variability, formation/dissociation of associated semi-volatile
 components and the impacts on visibility, Sci. Total. Environ., 199-206, 2014.
- 1501 Shindell, D., Kuylenstierna, J. C. I., Vignati, E., van Dingenen, R., Amann, M., Klimont, Z., Anenberg,
- 1502 S. C., Muller, N., Janssens-Maenhout, G., Raes, F., Schwartz, J., Faluvegi, G., Pozzoli, L., Kupiainen,
- 1503 K., Hoglund-Isaksson, L., Emberson, L., Streets, D., Ramanathan, V., Hicks, K., Oanh, N., Milly, G.,
- 1504 Williams, M., Demkin, V. and Fowler, D.: Simultaneously Mitigating Near-Term Climate Change and
- 1505 Improving Human Health and Food Security, Science, 6065, 183-189, 2012.
- 1506 Simcik, M. F., Eisenreich, S. J. and Lioy, P. J.: Source apportionment and source/sink relationships of
- PAHs in the coastal atmosphere of Chicago and Lake Michigan, Atmos. Environ., 30, 5071-5079,1999.
- 1509 Simoneit, B. R. T., Rushdi, A. I., Bin Abas, M. R. and Didyk, B. M.: Alkyl Amides and Nitriles as1510 Novel Tracers for Biomass Burning, Environ. Sci. Technol, 1, 16-21, 2003.
- Streets, D. G.: Dissecting future aerosol emissions: Warming tendencies and mitigation opportunities,
 Climatic Change, 3-4, 313-330, 2007.
- 1513 Sun, J., Peng, H., Chen, J., Wang, X., Wei, M., Li, W., Yang, L., Zhang, L., Wang, W. and Mellouki,
- 1514 A.: An estimation of CO_2 emission via agricultural crop residue open field burning in China from 1996
- to 2013, J. Clean Prod, 2625-2631, 2016.
- Takegawa, N., Miyakawa, T., Kawamura, K. and Kondo, Y.: Contribution of Selected Dicarboxylic
 and ω-Oxocarboxylic Acids in Ambient Aerosol to the m/z 44 Signal of an Aerodyne Aerosol Mass
 Spectrometer, AerosolSci. Tech., 41, 418-437, 2007.
- Tao, Y., Ye, X., Jiang, S., Yang, X., Chen, J., Xie, Y. and Wang, R.: Effects of amines on particle
 growth observed in new particle formation events, J. Geophys. Res.: Atmos., 121, 324-335,
 doi:10.1002/2015JD024245, 2016.
- 1522 Tian, D., Hu, Y., Wang, Y., Boylan, J. W., Zheng, M. and Russell, A. G.: Assessment of Biomass
- Burning Emissions and Their Impacts on Urban and Regional PM_{2.5}: A Georgia Case Study, Environ
 Sci. Technol., 2, 299-305, 2008.
- Tóth, A., Hoffer, A., Nyirő-Kósa, I., Pósfai, M. and Gelencs ér, A.: Atmospheric tar balls: aged primary
 droplets from biomass burning? Atmos. Chem. Phys. Dis., 12, 33089-33104, doi:10.5194/acp-146669-2014, 2013.
- Tsai, P. J., Shieh, H. Y., Lee, W. J. and Lai, S. O.: Health-risk assessment for workers exposed to
 polycyclic aromatic hydrocarbons (PAHs) in a carbon black manufacturing industry, Sci. Total.
 Environ., 1-3, 137-150, 2001.
- Urban, R. C., Alves, C. A., Allen, A. G., Cardoso, A. A. and Campos, M.: Organic aerosols in a
 Brazilian agro-industrial area: Speciation and impact of biomass burning, Atmos. Res., 271-279, 2016.
- 1533 Veres, P., Roberts, J. M., Burling, I. R., Warneke, C., de Gouw, J. and Yokelson, R. J.: Measurements
- of gas-phase inorganic and organic acids from biomass fires by negative-ion proton-transfer
 chemical-ionization mass spectrometry, J. Geophys Res, 115, D23302, doi:10.1029/2010JD014033,
 2010.
- 1537 Wang, L., Li, X. and Xu, Y.: The Economic Losses Caused By Crop Residues Burnt in Open Field in
- 1538 China, J. Arid Land Resources and Environment, 2, 170-175, 2008a.
- 1539 Wang, R., Tao, S., Wang, W., Liu, J., Shen, H., Shen, G., Wang, B., Liu, X., Li, W., Huang, Y., Zhang,
- 1540 Y., Lu, Y., Chen, H., Chen, Y., Wang, C., Zhu, D., Wang, X., Li, B., Liu, W., Ma, J. and Prospero, A.:
- 1541 Black carbon emissions in China from 1949 to 2050, Environ. Sci. Technol., 14, 7595-7603, 2012.

- 1542 Wang, S. and Zhang, C.: Spatial and temporal distribution of air pollutant emissions from open burning
- 1543 of crop residues in China, Sciencepaper Online, 5:329-333, 2008b.(*in chinese*)
- Wang, S., Zhao, M., Xing, J., Wu, Y., Zhou, Y., Lei, Y., He, K., Fu, L. and Hao, J.: Quantifying the air
 pollutants emission reduction during the 2008 Olympic Games in Beijing, Environ. Sci. Technol., 7,
 2490-2496, 2010.
- 1547 Wang, W., Jariyasopit, N., Schrlau, J., Jia, Y., Tao, S., Yu, T., Dashwood, R. H., Zhang, W., Wang, X.
- 1548 and Simonich, S.: Concentration and photochemistry of PAHs, NPAHs, and OPAHs and toxicity of
- 1549 PM_{2.5} during the Beijing Olympic Games, Environ. Sci. Technol., 16, 6887-6895, 2011.
- Wei, B. and Yang, L.: A review of heavy metal contaminations in urban soils, urban road dusts andagricultural soils from China, Microchem J., 2, 99-107, 2010.
- 1552 Wei, W., Jitao, Y., Qingling, Z. and Bailiang, Z.: Current Situation and Developing Direction of Straw
- 1553 Utilization Technology in China, China Resources Comprehensive Utilization, 11, 2004.
- 1554 WHO, Life Database in 2000, World Health Organization, 2000.
- 1555 Wilson, J. M., Baeza-Romero, M. T., Jones, J. M., Pourkashanian, M., Williams, A., Lea-Langton, A.
- 1556 R., Ross, A. B. and Bartle, K.: Soot Formation from the Combustion of Biomass Pyrolysis Products
- and a Hydrocarbon Fuel, n-Decane: An Aerosol Time Of Flight Mass Spectrometer (ATOFMS) Study,
 Energ Fuel, 3, 1668-1678, 2013.
- 1559 Wong, C., Vichit-Vadakan, N., Kan, H., Qian, Z. and Teams, T. P. P.: Public Health and Air Pollution
- in Asia (PAPA): A Multicity Study of Short-Term Effects of Air Pollution on Mortality, Environ.
 Health Persp., 9, 1195-1202, 2008.
- Wu, C., Liu, L. J. S., Cullen, A., Westberg, H. and Williamson, J., Spatial-temporal and cancer risk
 assessment of selected hazardous air pollutants in Seattle, Environ Int., 1, 11-17, 2011.
- Wu, C., Wu, S., Wu, Y., Cullen, A. C., Larson, T. V., Williamson, J. and Liu, L. J. S.: Cancer risk
 assessment of selected hazardous air pollutants in Seattle, Environ Int., 3, 516-522, 2009.
- 1566 Xu, S., Liu, W. and Tao, S.: Emission of Polycyclic Aromatic Hydrocarbons in China, Environ Sci.
 1567 Technol., 3, 702-708, 2006.
- Yang, C., Peng, X., Huang, W., Chen, R., Xu, Z., Chen, B. and Kan, H. A time-stratified
 case-crossover study of fine particulate matter air pollution and mortality in Guangzhou, China, Int
 Arch. Occ. Env. Hea., 5, 579-585, 2012.
- Yang, M., Howell, S. G., Zhuang, J. and Huebert, B. J.: Attribution of aerosol light absorption to black
 carbon, brown carbon, and dust in China-interpretations of atmospheric measurements during
 EAST-AIRE, Atmos. Chem. Phys., 6, 2035-2050, doi:10.5194/acp-9-2035-2009, 2009.
- Yang, Y., Liu, X., Qu, Y., Wang, J., An, J., Zhang, Y. and Zhang, F.: Formation mechanism of
 continuous extreme haze episodes in the megacity Beijing, China, in January 2013, Atmos. Res.,
 192-203, 2015.
- 1577 Yokelson, R. J., Karl, T., Artaxo, P. and Blake, D. R.: The Tropical Forest and Fire Emissions
- 1578 Experiment: overview and airborne fire emission factor measurements, Atmos. Chem. Phys., 7,
 1579 5175-5196, doi:10.5194/acp-7-5175-2007, 2007.
- Yunker, M. B., Macdonald, R. W., Vingarzan, R., Mitchell, R. H., Goyette, D. and Sylves tre, S.: PAHs
 in the Fraser River basin: a critical appraisal of PAH ratios as indicators of PAH source and
 composition, Org Geochem, 4, 489-515, 2002.
- 1583 Zhang, H., Hu, D., Chen, J., Ye, X., Wang, S. X., Hao, J. M., Wang, L., Zhang, R. and An, Z.: Particle
- Size Distribution and Polycyclic Aromatic Hydrocarbons Emissions from Agricultural Crop Residue
 Burning, Environ. Sci. Technol., 13, 5477-5482, 2011.

- 1586 Zhang, H., Wang, S., Hao, J., Wan, L., Jiang, J., Zhang, M., Mestl, H. E., Mestl, H., Alnes, L., Aunan,
- 1587 K. and Mellouki, A.: Chemical and size characterization of particles emitted from the burning of coal1588 and wood in rural households in Guizhou, China, Atmos. Environ., 94-99, 2012.
- 1589 Zhang, H., Ye, X., Cheng, T., Chen, J., Yang, X., Wang, L. and Zhang, R.: A laboratory study of
 agricultural crop residue combustion in China: Emission factors and emission inventory, Atmos.
 1591 Environ., 36, 8432-8441, 2008a.
- Zhang, R., Khalizov, A. F., Pagels, J., Zhang, D., Xue, H. and McMurry, P. H.: Variability in
 morphology, hygroscopicity, and optical properties of soot aerosols during atmospheric processing,
 Proc Natl Acad Sci USA, 30, 10291-10296, 2008b.
- Zhang, R., Suh, I., Zhao, J., Zhang, D., Fortner, E. C., Tie, X., Molina, L. T. and Molina, M. J.:
 Atmospheric new particle formation enhanced by organic acids, Science, 5676, 1487-1490, 2004.
- Zhao, B., Wang, P., Ma, J. Z., Zhu, S., Pozzer, A. and Li, W.: A high-resolution emission inventory of
 primary pollutants for the Huabei region, China, Atmos. Chem. Phys., 1, 481-501, 2012.
- 1599 Zhao, Y., Nielsen, C. P., Lei, Y., McElroy, M. B. and Hao, J.: Quantifying the Uncertainties of a
- 1600 Bottom-Up Emission Inventory of Anthropogenic Atmospheric Pollutants in China, Atmos. Chem.
- 1601 Phys., 11, 2295-2308, 2011.
- 1602 Zheng, J., Ma, Y., Chen, M., Zhang, Q., Wang, L., Khalizov, A. F., Yao, L., Wang, Z., Wang, X. and
- 1603 Chen, L.: Measurement of atmospheric amines and ammonia using the high resolution time -of-flight1604 chemical ionization mass spectrometry, 249-259, 2015.
- 1605

1607 **Tables and figure captions**

- **Table 1.** Emission factors of particulate chemical species in smoke $PM_{2.5}$ from agricultural residue burning (mean value \pm standard deviation).
- 1610 **Table 2.** Emission factors of particulate chemical species in smoke $PM_{1.0}$ from 1611 agricultural residue burning (mean value \pm standard deviation).
- 1612 **Table 3.** Comparison of emission factors with literature (specific chemical materials 1613 in form of $PM_{2,5}$)
- 1614 Table 4. Summary of field burning rates and economic data in China
- 1615 Table 5. National agricultural field burning emissions of BAU, EM, and NDRC
- 1616 scenarios in China, 2012.
- 1617 **Table 6.** Uncertainties for national smoke aerosol emissions in 2012.
- 1618 **Table 7.** Estimated number of cases (95% CI) attributable to agricultural fire smoke
- 1619 $PM_{2.5}$ exposure in China, 2012
- 1620 Table 8. Health-related economic loss (95% CI) from agricultural fire smoke PM_{2.5}
- 1621 exposure in China, 2012
- 1622 **Figure 1.** Schematic methodology for developing emission estimations
- **Figure 2.** Chemical profiles of smoke $PM_{2.5}$ and $PM_{1.0}$ from 5 types agricultural residue burnings. OM (organic matter = $1.3 \times OC$). OWSI, other water soluble ions including F⁻, NO₂⁻, Na⁺, Ca²⁺, Mg²⁺.
- **Figure 3.** a) Emission factors of 16 USEPA priority PAHs in smoke PM_{2.5} and PM_{1.0};
- 1627 b) expulsion-accumulation of PAHs in OC-EC of smoke $PM_{2.5}$ and $PM_{1.0}$
- **Figure 4.** Transmission electron microscope (TEM) images and EDX analysis of fresh agricultural residue burning particles. (a)-(c) Crystal and amorphous KCl particles internally mixed with sulfate, nitrate, and carbonaceous materials. (d)-(f) Heavy metal-bearing fractal-like fly ash particles. (e)-(g) Chain-like soot particles and tar ball.
- Figure 5. Annual agricultural residue production of five major crops and allocated
 into two harvest (summer and autumn harvest) based on agricultural yield in China,
 2012.
- 1636 Figure 6. Statistical analysis of field burning rates from BAU, EM, and NDRC1637 versions
- 1638 **Figure 7.** Spatial and temporal distribution of smoke $PM_{2.5}$ emissions and flux 1639 concentrations from agricultural field burning over China, 2012

- 1640 Figure 8. Nationwide PM_{2.5} emissions and flux concentrations based on different
- 1641 burning versions. The inset pie-graphs are chemical compositions of integrated $PM_{2.5}$
- 1642 from five major agricultural residue burning.

Chemical Species (g kg ⁻¹)	wheat straw	corn straw	rice straw	cotton residue	soybean residue
PM _{2.5}	5.803 ±0.363	5.988 ±0.723	14.732 ±2.417	15.162 ±2.053	3.249 ± 0.350
OC	2.813 ± 0.147	2.393 ± 0.351	6.882 ± 0.689	7.415 ± 0.547	1.539 ± 0.253
EC	0.676 ± 0.027	0.778 ± 0.152	2.182 ± 0.278	1.192 ± 0.171	0.614 ± 0.190
Inorganic ions (g kg ⁻¹)	1.273 ±0.072	1.810 ± 0.030	3.086 ±0.266	3.810 ±0.246	0.523 ±0.149
SO ₄ ²⁻	0.084 ± 0.028	0.217 ± 0.041	$0.409\ \pm 0.127$	$0.701\ \pm 0.081$	0.073 ± 0.014
Cl	0.576 ± 0.038	$0.709\ \pm 0.034$	1.158 ± 0.232	1.351 ± 0.114	0.178 ± 0.030
F	0.023 ± 0.061	0.061 ± 0.005	0.073 ± 0.024	0.265 ± 0.012	0.009 ± 0.004
NO ₃ ⁻	0.023 ± 0.000	0.032 ± 0.002	0.051 ± 0.025	0.072 ± 0.004	0.009 ± 0.004
NO_2^-	0.006 ± 0.001	0.016 ± 0.002	0.018 ± 0.002	0.036 ± 0.001	0.004 ± 0.003
Ca ²⁺	0.030 ± 0.011	0.036 ± 0.003	0.046 ± 0.007	0.060 ± 0.003	$0.010\ \pm 0.002$
Na^+	0.005 ± 0.001	0.012 ± 0.001	0.028 ± 0.004	0.050 ± 0.004	$0.005 \ \pm 0.001$
$\mathbf{NH_4}^+$	0.152 ± 0.005	0.197 ± 0.010	0.542 ± 0.107	0.347 ± 0.008	0.029 ± 0.004
Mg^{2+}	0.005 ± 0.000	0.017 ± 0.002	0.023 ± 0.004	$0.032 \ \pm 0.002$	$0.005 \ \pm 0.001$
\mathbf{K}^{+}	0.368 ± 0.041	0.514 ± 0.009	0.739 ± 0.049	0.947 ± 0.070	0.200 ± 0.023
Organic Acids (mg kg ⁻¹)	156.680 ±81.830	46.670 ±9.000	557.130 ±269.380	769.990 ±317.550	143.310 ±39.770
CH ₃ COOH	148.900 ±79.290) 36.640 ±8.210	417.930 ±186.140	743.320 ±159.600	135.500 ±62.320
MSA	7.170 ± 2.110	10.030 ± 30.000	136.990 ± 81.700	12.980 ± 1.530	3.200 ± 1.530
$H_2C_2O_4$	2.610 ± 0.430	ND	2.210 ± 1.560	4.760 ± 2.640	2.170 ± 2.380
НСООН	ND	ND	ND	8.930 ± 2.630	$2.440\ \pm 1.450$
Amine salts (mg kg ⁻¹)	19.246 ±9.368	32.877 ±19.141	104.787 ±15.635	102.409 ±13.379	4.514 ±1.776
$MeOH^+ + MMAH^+$	1.322 ± 0.086	$5.735\ {\pm}0.102$	$17.226\ \pm 1.454$	$19.888\ \pm 0.351$	$0.456\ {\pm}0.196$
$\mathbf{MEAH}^{\scriptscriptstyle +}$	0.201 ± 0.055	0.675 ± 0.135	$4.175\ \pm 0.920$	3.690 ± 1.959	ND
$\mathrm{TEOH}^{\scriptscriptstyle +}$	2.562 ± 0.962	4.118 ± 0.741	25.129 ± 0.343	14.376 ± 8.688	0.672 ± 0.558
$\mathbf{DEAH}^{+} + \mathbf{TMAH}^{+}$	13.728 ± 7.512	18.973 ± 0.466	46.148 ±12.185	28.568 ± 5.321	2.012 ± 0.878
\mathbf{DMAH}^{+}	1.434 ± 0.925	3.376 ± 0.674	12.110 ± 6.166	35.887 ± 2.940	1.374 ± 0.144
Elemental Species (mg kg ⁻¹)	53.813 ±18.860	53.546 ±9.070	131.612 ±5.920	27.577 ±3.700	14.003 ±8.710
Phenols (mg kg ⁻¹)	26.785 ±8.582	16.390 ±2.652	27.238 ±4.861	41.481 ±5.517	9.673 ±2.272
PAHs (mg kg ⁻¹)	1.814 ± 0.348	2.706 ± 0.798	7.267 ±1.722	8.302 ±2.856	1.832 ± 0.353

Table 1. Emission factors of particulate chemical species in smoke $PM_{2.5}$ from agricultural residue burning (mean value \pm standard deviation).

1646 ND means not detected

Chemical Species (g kg ⁻¹)	wheat straw	corn straw	rice straw	cotton residue	soybean residue
$PM_{1.0}$	5.298 ± 0.295	5.360 ± 0.551	13.200 ± 1.440	12.635 ± 1.243	3.036 ± 0.257
OC	2.419 ± 0.126	2.063 ± 0.340	6.024 ± 0.602	6.036 ± 0.360	1.338 ± 0.128
EC	0.650 ± 0.037	0.728 ± 0.122	2.083 ± 0.413	1.023 ± 0.205	0.575 ± 0.260
Inorganic ions (g kg ⁻¹)	1.215 ±0.040	1.768 ± 0.010	2.940 ±0.249	3.516 ±0.145	0.510 ± 0.156
SO4 ²⁻	0.078 ± 0.011	0.199 ± 0.032	0.333 ± 0.107	0.581 ± 0.054	0.073 ± 0.056
Cl	0.544 ± 0.033	0.712 ± 0.027	1.145 ± 0.118	1.243 ± 0.067	0.175 ± 0.031
F	0.022 ± 0.007	0.041 ± 0.004	0.078 ± 0.030	0.151 ± 0.011	$0.001 \ \pm 0.001$
NO ₃ ⁻	$0.021 \ \pm 0.005$	$0.027 \ \pm 0.002$	0.043 ± 0.016	0.061 ± 0.003	$0.009\ \pm 0.002$
NO_2^-	0.006 ± 0.001	0.010 ± 0.003	0.013 ± 0.004	$0.019\ \pm 0.002$	0.004 ± 0.003
Ca ²⁺	0.027 ± 0.013	0.028 ± 0.002	0.045 ± 0.008	0.067 ± 0.005	$0.010\ \pm 0.002$
Na ⁺	0.004 ± 0.000	0.012 ± 0.000	0.027 ± 0.003	0.056 ± 0.006	0.005 ± 0.002
$\mathrm{NH_4}^+$	0.147 ± 0.005	0.191 ± 0.009	0.511 ± 0.067	0.401 ± 0.004	$0.031\ {\pm}0.005$
Mg^{2+}	0.005 ± 0.001	0.035 ± 0.001	0.024 ± 0.006	0.033 ± 0.002	$0.005 \ \pm 0.001$
\mathbf{K}^{+}	0.359 ± 0.040	0.513 ± 0.015	0.721 ± 0.073	0.994 ± 0.067	$0.197\ {\pm}0.035$
Organic Acids (mg kg ⁻¹)	124.310 ±25.170	47.830 ±10.610	427.400 ±221.270	639.820 ±244.960	130.760 ±59.310
CH ₃ COOH	115.790 ±21.940	38.960 ± 9.610	383.360 ±179.050	615.790 ±232.860	124.310 ±69.000
MSA	6.830 ± 2.030	8.870 ± 2.730	$41.380\ \pm 38.480$	$11.380\ {\pm}2.360$	3.200 ± 1.730
$H_2C_2O_4$	1.690 ± 1.200	ND	$2.660\ \pm 1.760$	3.620 ± 1.250	$1.560\ \pm 1.670$
НСООН	ND	ND	ND	9.030 ± 7.710	$1.690\ \pm 1.390$
Amine salts (mg kg ⁻¹)	18.191 ±5.351	29.891 ±13.480	81.726 ± 11.455	85.720 ±21.337	4.385 ±1.445
$MeOH^+ + MMAH^+$	1.300 ± 0.282	5.647 ± 0.342	16.627 ± 0.104	18.834 ± 1.991	0.464 ± 0.265
\mathbf{MEAH}^{+}	0.157 ± 0.037	0.787 ± 0.211	3.581 ± 0.602	2.771 ±1.304	ND
$\mathbf{T}\mathbf{EOH}^{+}$	1.719 ± 0.283	5.115 ± 0.732	17.575 ± 0.844	11.441 ± 3.229	0.529 ± 0.304
$DEAH^+ + TMAH^+$	13.716 ±9.047	15.921 ± 1.620	33.565 ± 6.795	$29.057\ \pm 3.793$	2.278 ± 0.533
\mathbf{DMAH}^{+}	1.300 ± 0.702	2.420 ± 0.575	10.377 ±4.521	23.617 ± 20.086	1.115 ± 0.343
Elemental Species (mg kg ⁻¹)	31.586 ±10.630	29.265 ±4.240	51.062 ± 5.920	16.738 ± 3.480	11.817 ±6.650
Phenols (mg kg ⁻¹)	20.774 ±4.972	13.193 ±2.181	20.480 ± 1.403	23.521 ±8.521	7.689 ±1.356
PAHs (mg kg ⁻¹)	1.257 ± 0.398	1.420 ± 0.232	3.967 ± 0.970	4.359 ±1.373	1.123 ± 0.205

Table 2. Emission factors of particulate chemical species in smoke $PM_{1.0}$ from agricultural residue burning (mean value \pm standard deviation).

1649 ND means not detected

See at a s		Emission factors (g kg ⁻¹)	Deferment		
Species	This work	Reference value	Kelerence		
PM _{2.5}	8.99 ±5.55	7.6~11.7(AR), 6.26~15.3 (TL), ~3.0 (AR), 2.2~15.0 (AR)	Li et al., 2007; Akagi et al., 2011; Dhammapala et al., 2007; Hayashi et al., 2014		
PM _{1.0}	7.91 ± 4.67	4.4.3~12.1 (TL)	May et al., 2014		
OC	4.21 ±2.73	2.7~3.9 (AR), 2.3~9.7(TL), ~1.9(AR) , 1.0~9.3 (AR), 0.8~5.9 (TL)	Li et al., 2007; Akagi et al., 2011; Dhammapala et al., 2007; Hayashi et al., 2014; May et al., 2014		
EC	1.09 ± 0.65	0.35~0.49 (AR), 0.37~0.91(TL), ~0.4(AR), 0.21~0.81(AR), 1.13~1.73 (TL)	Li et al., 2007; Akagi et al., 2011; Dhammapala et al., 2007; Hayashi et al., 2014; May et al., 2014		
WSOA	0.33 ± 0.31	0.039~0.109 (TL)	Akagi et al., 2011		
WSA	0.05 ± 0.05	0.08~0.13 (TL), ~0.55 (TL)	Akagi et al., 2011; Andreae et al., 2001		
WSI	2.10 ±1.34	1.84~4.9 (AR),0.8~1.31(TL), 0.43~1.63 (AR)	Li et al., 2007; Akagi et al., 2011; Hayashi et al., 2014		
THM	0.06 ± 0.05	0.06~0.09 (AR)	Li et al., 2007		
PAHs (×10 ³)	4.38 ± 3.15	~17(AR), 0.72~1.64(AR), ~9.0 (W)	Dhammapala et al., 2007; Zhang et al., 2011; Lee et al.2005		
Phenols $(\times 10^3)$	24.31 ±12.11	~35(AR), ~5 (AR), ~13 (TL)	Dhammapala et al., 2007; Hays et al., 2005; Andreae et al., 2001		

Table 3. Comparison of emission factors with literature (specific chemical materials in form of $PM_{2.5}$).

AR: agricultural residue; TL: total, including forest fires and straw burning; W: wood

Duarinaa	Burning rate from literature		Agricultural income ratio ^c			Estimated	burning rate	NDRC report ^d	Avanaga nata
Province	BAU-I ^a	BAU-II ^b	2000	2006	2012	EM-I	EM-II	NDRC	Average rate
Beijing	0.00	0.17	0.08	0.06	0.06	0.00	0.19	0.13	$0.10\ \pm 0.08$
Tianjin	0.00	0.17	0.10	0.14	0.12	0.00	0.20	0.30	$0.13\ \pm 0.12$
Hebei	0.20	0.17	0.27	0.22	0.24	0.22	0.16	0.19	$0.19\ \pm 0.02$
Shanxi	0.20	0.17	0.20	0.21	0.25	0.16	0.14	0.22	$0.18\ \pm 0.03$
Inner Mongolia	0.00	0.12	0.44	0.49	0.66	0.00	0.09	0.27	$0.10\ \pm 0.10$
Liaoning	0.20	0.12	0.30	0.29	0.39	0.16	0.09	0.34	$0.18\ \pm 0.09$
Jilin	0.30	0.12	0.73	0.73	0.77	0.28	0.11	0.25	$0.21\ \pm 0.08$
Heilongjiang	0.30	0.12	0.99	0.83	0.59	0.50	0.17	0.25	$0.27\ \pm 0.13$
Shanghai	0.00	0.32	0.10	0.08	0.09	0.00	0.29	0.12	$0.15\ \pm 0.14$
Jiangsu	0.30	0.32	0.32	0.22	0.30	0.32	0.23	0.19	$0.27\ \pm 0.05$
Zhejiang	0.30	0.32	0.19	0.08	0.09	0.64	0.28	0.22	$0.35\ \pm 0.15$
Anhui	0.20	0.32	0.44	0.39	0.43	0.21	0.29	0.43	$0.29\ \pm 0.08$
Fujian	0.30	0.32	0.18	0.10	0.14	0.39	0.22	0.17	$0.28\ \pm 0.08$
Jiangxi	0.20	0.11	0.45	0.31	0.44	0.20	0.08	0.25	$0.17\ \pm 0.06$
Shandong	0.30	0.17	0.33	0.25	0.24	0.40	0.17	0.21	$0.25\ \pm 0.09$
Henan	0.20	0.17	0.39	0.35	0.33	0.23	0.18	0.22	$0.20\ \pm 0.02$
Hubei	0.20	0.11	0.42	0.30	0.41	0.21	0.08	0.30	$0.18\ \pm 0.08$
Hunan	0.20	0.33	0.47	0.31	0.43	0.22	0.24	0.35	$0.27\ \pm 0.06$
Guangdong	0.30	0.33	0.19	0.10	0.13	0.44	0.25	0.18	$0.30\ \pm 0.09$
Guangxi	0.20	0.33	0.40	0.25	0.33	0.25	0.25	0.35	$0.28\ \pm 0.06$
Hainan	0.30	0.33	0.35	0.16	0.21	0.51	0.25	0.56	$0.39\ \pm 0.12$
Chongqing	0.20	0.11	0.35	0.23	0.30	0.24	0.08	0.45	$0.22\ \pm 0.13$
Sichuan	0.20	0.11	0.37	0.22	0.28	0.26	0.09	0.30	$0.19\ \pm 0.08$
Guizhou	0.20	0.11	0.38	0.23	0.25	0.31	0.10	0.43	$0.23\ \pm 0.13$
Yunnan	0.20	0.11	0.36	0.26	0.31	0.24	0.09	0.28	$0.18\ \pm 0.07$
Tibet	0.00	0.16	0.15	0.09	0.05	0.00	0.30	0.16	$0.12\ \pm 0.11$
Shannxi	0.20	0.17	0.33	0.27	0.26	0.25	0.18	0.28	$0.22\ \pm 0.04$
Gansu	0.10	0.16	0.25	0.20	0.28	0.09	0.11	0.33	$0.16\ \pm 0.09$
Qinghai	0.00	0.16	0.23	0.10	0.08	0.00	0.20	0.28	$0.13\ \pm 0.11$
Ningxia	0.10	0.16	0.42	0.38	0.45	0.09	0.13	0.16	$0.13\ \pm 0.03$
Xinjiang	0.10	0.16	0.43	0.61	0.73	0.06	0.13	0.30	$0.15\ \pm 0.08$
Nationwide	0.21	0.16	0.34	0.27	0.31	0.26	0.15	0.27	0.21 ± 0.05

Table 4. Summary of field burning rates and economic data in China.

a. Zhao et al., 2012; Cao et al., 2006; Cao et al., 2011

b. Wang and Zhang., 2008

c. Calculated based on data from China Yearbook 2001~2013 (NBSC, 2001-2013), China Rural Statistic Yearbook 2001~2013, data available at http://www.grain.gov.cn/Grain/

d. Data from the National Development and Reform Commission report ([2014]No.516): http://www.sdpc.gov.cn/

	BAU-I		BAU-II		EM-1		EM-2		NDRC			Average						
Unit: Gg	Total	Summer	Autumn	Total	Summer	Autumn	Total	Summer	Autumn	Total	Summer	Autumn	Total	Summer	Autumn	Total	Summer	Autumn
PM _{2.5}	1001.05	218.99	782.06	835.42	209.29	626.13	1211.92	258.58	953.34	738.36	182.34	556.02	1241.69	258.24	983.46	1007.650	226.007	781.646
$PM_{1.0}$	897.52	198.93	698.59	748.57	189.92	558.65	1087.05	234.85	852.20	661.81	165.61	496.20	1111.90	234.44	877.46	903.125	205.217	697.911
OC	429.51	102.87	326.64	360.99	97.67	263.32	519.26	121.33	397.94	318.84	85.55	233.29	533.19	120.86	412.33	433.184	105.885	327.300
EC	133.61	27.37	106.24	111.40	26.52	84.88	162.71	32.39	130.32	98.06	22.85	75.21	164.97	32.53	132.45	134.414	28.404	106.010
SO_4^{2-}	30.22	3.96	26.26	24.97	3.94	21.04	36.39	4.71	31.68	22.09	3.32	18.76	38.21	4.78	33.44	30.440	4.155	26.285
NO ₃ ⁻	4.35	0.84	3.51	3.55	0.80	2.75	5.24	0.99	4.25	3.17	0.70	2.47	5.40	0.99	4.41	4.350	0.864	3.486
NH_4^+	32.08	6.37	25.71	26.65	6.21	20.44	39.09	7.54	31.55	23.43	5.32	18.11	39.46	7.59	31.87	32.202	6.623	25.580
\mathbf{K}^{+}	67.49	13.12	54.38	54.75	12.38	42.37	81.40	15.45	65.95	49.10	10.90	38.20	83.62	15.36	68.26	67.412	13.469	53.943
WSOA	24.44	6.55	17.89	21.94	6.39	15.55	29.69	7.76	21.93	18.77	5.48	13.30	30.82	7.81	23.01	25.174	6.815	18.360
WSA	5.75	0.95	4.80	4.85	0.95	3.90	6.99	1.13	5.86	4.23	0.80	3.43	7.19	1.15	6.04	5.815	1.000	4.815
PAHs	0.48	0.11	0.37	0.40	0.10	0.30	0.58	0.12	0.45	0.35	0.09	0.26	0.59	0.13	0.47	0.480	0.109	0.371
Phenols	2.71	0.85	1.87	2.25	0.78	1.47	3.25	0.99	2.26	2.02	0.70	1.323	3.40	0.98	2.36	2.721	0.861	1.861
THM	8.68	2.01	6.67	7.19	1.92	5.27	10.56	2.37	8.19	6.36	1.67	4.69	10.64	2.37	8.27	8.702	2.073	6.628
WSI	249.96	47.46	202.50	204.46	45.24	159.22	301.75	56.01	245.74	182.31	39.50	142.82	310.31	55.88	254.43	250.269	48.927	201.342

Table 5. National agricultural field burning emissions of BAU, EM, and NDRC scenarios in China in 2012.

									-			
Species		BAU-I		BAU-II		EM-I		EM-II		NDRC		Average
PM _{2.5}	1001.1	(-52.3% , 73.5%)	835.4	(-48.7% , 68.8%)	1211.9	(-63.6% , 84.3%)	738.4	(-55.9% , 74.3%)	1241.7	(-46.2% , 65.1%)	1005.7	(-24.6% , 33.7%)
$PM_{1.0}$	897.5	(-51.6% , 73.0%)	748.6	(-48.4% , 68.6%)	1087.1	(-62.9%, 83.8%)	661.8	(-55.5% ,74.1%)	1111.9	(-45.7% , 64.7%)	901.4	(-24.4% , 33.5%)
OC	429.5	(-50.5% ,71.5%)	361.0	(-48.9% , 69.2%)	519.3	(-61.4% , 81.8%)	318.8	(-55.6% , 74.1%)	533.2	(-47.1% , 66.7%)	432.4	(-24.2% , 33.3%)
EC	133.6	(-52.1%, 73.6%)	111.4	(-50.1% , 71.0%)	162.7	(-63.3% , 84.3%)	98.1	(-56.8% , 75.7%)	165.0	(-46.7% , 66.0%)	134.2	(-24.8% , 34.0%)
WSOA	24.4	(-68.5% , 86.2%)	21.9	(-75.7%, 95.2%)	29.7	(-78.7%, 96.2%)	18.8	(-77.8% , 95.4%)	30.8	(-67.5% , 85.1%)	25.1	(-33.3% ,41.4%)
WSA	5.8	(-62.8% , 82.1%)	4.9	(-65.9%, 84.1%)	7.0	(-73.9%, 93.2%)	4.2	(-69.3%, 86.3%)	7.2	(-58.7% , 75.9%)	5.8	(-30.1% , 38.5%)
WSI	250.0	(-54.4% , 77.2%)	204.5	(-47.5% , 67.4%)	301.8	(-66.9%, 89.3%)	182.3	(-56.1% , 74.8%)	310.3	(-46.9% , 66.4%)	249.8	(-25.4% , 34.9%)
THM	8.7	(-56.2% , 77.5%)	7.2	(-52.8% ,71.4%)	10.6	(-67.5%, 88.3%)	6.4	(-61.2% , 79.5%)	10.6	(-50.8% , 69.4%)	8.7	(-26.6% , 35.6%)
PAHs	0.5	(-55.2% , 75.7%)	0.4	(-52.4% , 72.2%)	0.6	(-66.5% , 86.8%)	0.4	(-58.8% , 76.9%)	0.6	(-49.3% , 67.8%)	0.5	(-26.0% , 34.9%)
Phenols	2.7	(-56.1% , 77.6%)	2.3	(-51.4% , 70.6%)	3.3	(-67.3%, 88.3%)	2.0	(-59.9% , 78.4%)	3.4	(-48.7% , 67.1%)	2.7	(-26.1% , 35.1%)

Table 6. Uncertainties for the national smoke aerosol emissions in 2012 (pollutant emission in unit of Gg/yr, 95% CI in percentage)

011111, 2012	-			
Emission version	Mortality Respiratory hospital admission		Cardiovascular hospital admission	Chronic bronchitis
BAU-I	7864 (3154, 12489)	31123 (21114, 40788)	29454 (12849, 45481)	7577067 (2952006, 11024705)
BAU-II	7187 (3056, 11260)	28711 (19443, 37693)	27156 (11825, 42007)	7132581 (2735111, 10523803)
EM-I	9435 (3817, 14933)	36950 (25151, 48269)	35116 (15373, 54042)	8712880 (3484325, 12430411)
EM-II	6175 (2554, 9751)	25166 (17004, 33112)	23745 (10316, 36816)	6383442 (2407643, 9526727)
NDRC	8523 (3581, 13377)	33957 (23015, 44542)	32131 (14003, 49664)	8332216 (3228351, 12148274)
Average	7836 (3232, 12362)	31181 (21145, 40881)	29520 (12873, 45602)	7267237 (2961487, 1130784)
CRC	538 (227, 850)	2191 (1462, 2920)	2038 (874, 3199)	636650 (214617, 1052153)

Table 7. Estimated number of cases (95% CI) attributable to agricultural fire smoke $PM_{2.5}$ exposure in China, 2012.

			Economic	- T-4-1 4				
Emission version	Emission version	Mortality	Respiratory hospital admission	Cardiovascular hospital admission	Chronic bronchitis	(million US\$)	GDP ratio (‰)	
	BAU-1	1544.5 (730.7, 2430.0)	19.6 (13.3, 25.7)	36.0 (15.7, 55.6)	7187.6 (2800.3, 10458.3)	8787.8 (3560.0, 12969.4)	1.0 (0.4, 1.5)	
	BAU-2	1453.9 (719, 2252.2)	18.1 (12.2, 23.8)	33.2 (14.4, 51.3)	6766.0 (2594.5, 9982.9)	8271.2 (3340.3, 12310.3)	1.0 (0.4, 1.4)	
	EM-1	1855.2 (870.3, 2913.7)	23.3 (15.9, 30.5)	42.9 (18.8, 66.1)	8265.0 (3305.2, 11791.5)	10186.5 (4210.2, 14801.8)	1.2 (0.5, 1.7)	
	EM-2	1228.1 (600.6, 1917.6)	15.9 (10.7, 20.9)	29.0 (12.6, 450)	6055.3 (2283.9, 9037.1)	7328.4 (2907.9, 11020.7)	0.9 (0.3, 1.3)	
	NDRC	1573.4 (759.3, 2456.2)	21.4 (14.5, 28.1)	39.3 (17.1, 60.7)	7903.9 (3062.4, 11523.9)	9538.2 (3853.4, 14069.0)	1.1 (0.4, 1.6)	
	Average	1531.0 (736.0, 2393.9)	19.7 (13.3, 25.8)	36.1 (15.7, 55.7)	7235.6 (2809.3, 10558.7)	8822.4 (3574.4, 13034.2)	1.0 (0.4, 1.5)	
	CRC	100.0 (48.0, 157.1)	1.3 (0.9, 1.8)	2.4 (1.0, 3.9)	603.9 (203.6, 998.1)	707.8 (253.6, 1160.9)	0.1 (0.0, 0.1)	

 Table 8. Health-related economic loss (95% CI) from agricultural fire smoke PM2.5 exposure in China, 2012.







residue burnings. OM (organic matter = $1.3 \times OC$). OWSI, other water soluble ions including F^- , NO_2^- , Na^+ , Ca^{2+} , and Mg^{2+} .





Figure 3. (a) Emission factors of 16 USEPA priority PAHs in smoke PM_{2.5} and PM_{1.0};
(b) expulsion-accumulation of PAHs in OC-EC of smoke PM_{2.5} and PM_{1.0}.



Figure 4. Transmission electron microscope (TEM) images and EDX analysis of fresh agricultural residue burning particles. (a)-(c) Crystal and amorphous KCl particles internally mixed with sulfate, nitrate, and carbonaceous materials. (d)-(f) Heavy metal-bearing fractal-like fly ash particles. (e)-(g) Chain-like soot particles and tar ball.





Figure 5. Annual agricultural residue production of five major crops and allocated
into two harvest (summer and autumn harvest) based on agricultural yield in China,
2012. (Abbreviation, BJ: Beijing; TJ: Tianjin; HB: Hebei; SX: Shanxi; IM: Inner Mongolia; LN: Liaoning; JL:
Jilin; HLJ: Heilongjiang; SH: Shanghai; JS: Jiangsu; ZJ: Zhejiang; AH: Anhui; FJ: Fujian; JX: Jiangxi; SD:
Shandong; HN: Henan; HUB: Hubei; HUN: Hunan; GD: Guangdong; GX: Guangxi; HAN: Hainan; CQ:
Chongqing; SC: Sichuan; GZ: Guizhou; YN: Yunnan; SHX: Shannxi; GS: Gansu; QH: Qinghai; NX: Ningxia; XJ:
Xinjiang)



Figure 6. Statistical analysis of field burning rates from BAU, EM, and NDRC
versions. The North Plain (Anhui, Shandong, Hebei, Shanxi, Tianjin, Beijing), the
Central of China (Hunan, Henan, Hubei), the Yangtze River Delta (Zhejiang, Jiangsu,
Shanghai), the Northeast of China (Heilongjiang, Liaoning, Jilin), the Pan-Pearl River
Delta (Hainan, Guangdong, Fujian, Guangxi, Guizhou, Sichuan, Yunnan, Jiangxi), the
West of China (Shannxi, Chongqing, Xinjiang, Qinghai, Ningxia, Tibet, Inner
Mongolia, Gansu)



38

Figure 7. Spatial and temporal distribution of smoke $PM_{2.5}$ emissions and flux concentrations from agricultural field burning over China, 2012.



Figure 8. Nationwide PM_{2.5} emissions and flux concentrations based on different
burning versions. The inset pie-graphs are chemical compositions of integrated PM_{2.5}
from five major agricultural residue burning.