Quantifying primary and secondary humic-like substances in urban aerosol based on emission source characterization and a source-oriented air quality model

Xinghua Li¹, Junzan Han¹, Philip K. Hopke², Jingnan Hu³, Qi Shu¹, Qing Chang¹, and Qi Ying⁴

¹School of Space and Environment, Beihang University, Beijing, 100191, China
²Center for Air Resources Engineering and Science, Clarkson University, Potsdam, NY, USA
³State Environmental Protection Key Laboratory of Vehicle Emission Control and Simulation, Chinese Research Academy of Environmental Sciences, Beijing 100012, China
⁴Zachry Department of Civil Engineering, Texas A&M University, College Station, TX 77843, USA

Correspondence: Xinghua Li (lixinghua@buaa.edu.cn) and Qi Ying (qying@civil.tamu.edu)

Received: 30 July 2018 – Discussion started: 2 October 2018
Revised: 21 December 2018 – Accepted: 3 February 2019 – Published: 22 February 2019

Abstract. Humic-like substances (HULIS) are a mixture of high-molecular-weight, water-soluble organic compounds that are widely distributed in atmospheric aerosol. Their sources are rarely studied quantitatively. Biomass burning is generally accepted as a major primary source of ambient humic-like substances (HULIS) with additional secondary material formed in the atmosphere. However, the present study provides direct evidence that residential coal burning is also a significant source of ambient HULIS, especially in the heating season in northern China based on source measurements, ambient sampling and analysis, and apportionment with source-oriented CMAQ modeling. Emission tests show that residential coal combustion produces 5 % to 24 % of the emitted organic carbon (OC) as HULIS carbon (HULISc). Estimation of primary emissions of HULIS in Beijing indicated that residential biofuel and coal burning contribute about 70 % and 25 % of annual primary HULIS, respectively. Vehicle exhaust, industry, and power plant contributions are negligible. The average concentration of ambient HULIS in PM₂.₅ was 7.5 µg m⁻³ in urban Beijing and HULIS exhibited obvious seasonal variations with the highest concentrations in winter. HULISc accounts for 7.2 % of PM₂.₅ mass, 24.5 % of OC, and 59.5 % of water-soluble organic carbon. HULIS are found to correlate well with K⁺, Cl⁻, sulfate, and secondary organic aerosol, suggesting its sources include biomass burning, coal combustion, and secondary aerosol formation. Source apportionment based on CMAQ modeling shows residential biofuel and coal burning and secondary formation are important sources of ambient HULIS, contributing 47.1 %, 15.1 %, and 38.9 %, respectively.

1 Introduction

Humic-like substances (HULIS) are a mixture of high-molecular-weight organic compounds that resemble terrestrial and aquatic humic and fulvic acids with similar structures and properties (Grabber and Rudich, 2006). HULIS are widely distributed in atmospheric aerosol, rain, and cloud and fog droplets and account for a significant proportion of organic carbon and water-soluble organic carbon (WSOC). For example, Zheng et al. (2013) reported that 9 % to 72 % of WSOC is HULIS. Because of their water-soluble and strong surface-active properties, HULIS may play an important role in the formation of clouds as condensation nuclei and ice nuclei and through aerosol hygroscopic growth (Dinar et al., 2006; Wang and Knopf, 2011; Gysel et al., 2004). Due to their strong light absorption in the UV range, HULIS can play an active role as brown carbon in radiative transfer and photochemical processes (Hoffer et al., 2006). HULIS deposition can also lead to a decrease in the albedo of ice and snow surfaces (Beine et al., 2011; France et al., 2011, 2012). Owing to their redox-active characteristics, HULIS were also suggested to induce adverse health effects (Lin and Yu, 2011; Ghio et al., 1996; Verma et al., 2012).

Published by Copernicus Publications on behalf of the European Geosciences Union.
In recent years, studies focusing on the spatial and temporal variations, sources, and formation of HULIS have been reported. A summary of these studies is provided in Table S1 in the Supplement. Previous studies have identified primary emissions and atmospheric secondary formation as important sources of HULIS. Among the primary emission sources, biomass burning is generally accepted as a major source of HULIS, with evidence from ambient and source sampling as well as source apportionment studies (Lin et al., 2010a, b; Kuang et al., 2015; Park and Yu, 2016; Schmidl et al., 2008a, b; Goncalves et al., 2010). Recently, residential coal burning was suggested as an important primary HULIS source during winter (Tan et al., 2016; Voliotis et al., 2017). However, direct evidence of HULIS emissions from coal combustion is limited. Only one study on HULIS emitted from residential coal combustion was reported and the results showed that HULIS accounted for 5.3 % of smoke PM$_{2.5}$ (Fan et al., 2016). Unfortunately, only a light coal in the shape of honeycomb briquette that did not reflect the variety of coal types and forms actually being used for residential heating and cooking in China was tested. Another possible primary HULIS source is vehicle exhaust, although there is uncertainty in the importance of this source (El Haddad et al., 2009; Salma et al., 2007; Lin et al., 2010b; Kuang et al., 2015). No direct evidence of primary HULIS in vehicle exhaust has been reported. Secondary processes, including formation in cloud droplets and heterogeneous or aerosol-phase reactions, can be important sources of HULIS (Lin et al., 2010b; Zheng et al., 2013).

Previous studies of HULIS source identification were generally based on the relationship between HULIS and the tracers for specific sources (such as K, levoglucosan, Cl$^-$, etc.) (Voliotis et al., 2017; Tan et al., 2016; Lin et al., 2010a; Park and Son, 2016; Baduel et al., 2010). Those correlation analyses between HULIS and some species may provide some information regarding possible sources and formation of HULIS. However, they do not provide quantitative source apportionments. To date, studies that quantitatively identify HULIS sources are limited (Kuang et al., 2015; Srivastava et al., 2018). Kuang et al. (2015) applied positive matrix factorization (PMF) to apportion sources of ambient HULIS in the Pearl River Delta (PRD) in southern China. Their study showed that secondary formation was the most important source of HULIS throughout the year with an annual average contribution of 69 % at an urban site. Biomass burning also contributed significantly to ambient HULIS.

Thus, information on the quantitative apportionment of HULIS sources is scarce in the more polluted regions in northern China, especially in the winter when large quantities of coal are consumed for indoor heating. Moreover, a considerable proportion of coal is burned in residential household stoves in rural, suburban, and even some urban areas under poor combustion conditions and without any emission controls. This coal combustion results in high air pollutant emissions and high ambient concentrations. Wang et al. (2016) estimated that more than 30 million tons of coal are burned per year in households in just the Beijing, Tianjin, and Hebei (BTH) region in northern China. Residential sources in the BTH region contributed to 32 % and 50 % of primary PM$_{2.5}$ emissions over the whole year and in winter, respectively (Liu et al., 2016).

The primary goals of this study are to determine whether residential coal combustion is a significant source of ambient HULIS and quantify its contributions to HULIS in Beijing. Given the large vehicle population in Beijing (up to 5.2 million in 2012), this study also provides a chance to examine the vehicular emissions’ contribution to ambient HULIS. Studies included (1) characterization of the HULIS emitted from residential coal stoves, vehicle exhaust, and residential biofuel burning; (2) estimation of anthropogenic primary emissions of HULIS based on these source measurements; (3) Measurement of HULIS concentrations and other major species in the ambient urban Beijing PM$_{2.5}$ from June 2012 to April 2013; and (4) application of the source-oriented Community Multiscale Air Quality (CMAQ) model to quantitatively determine the source contributions to HULIS. The information obtained in this study improves our understanding of the characteristics and sources of primary HULIS and the impact of those sources on HULIS in ambient PM$_{2.5}$.

2 Materials and methods

2.1 Ambient sampling

Beijing is surrounded by mountains to the west, north, and northeast and is adjacent to the northwest portion of the North China Plain. It has a warm and semi-humid continental monsoon climate with four distinctive seasons, characterized by strong windy and dusty weather in spring, high temperatures and humidity with extensive rain in summer, cool and pleasant weather in autumn, and cold and dry weather in winter. The annual average wind speed is 2.5 m s$^{-1}$ with mostly northerly or northwesterly winds in winter and southerly or southeasterly winds in summer.

From 14 June 2012 to 2 April 2013, 24 h ambient PM$_{2.5}$ samples were collected noncontinuously on the campus of Beihang University (BHU, 39°59′ N, 116°21′ E) (Fig. S1 in the Supplement). The sampling period covered four seasons with 132 samples being collected for HULIS analysis. The site is surrounded by educational and residential districts without major industrial sources. Major nearby roads are the North Fourth Ring Road about 900 m to the north, North Third Ring Road about 1.2 km to the south, and Xueyuan Road about 350 m to the east. Ambient PM$_{2.5}$ sampling instruments were installed on the roof of a building approximately 25 m above ground level at Beihang University. A high-volume aerosol sampler (RFPS-1287-063, Thermo Fisher Scientific, USA) was operated at a flow rate of 1.13 m$^3$ min$^{-1}$ to collect PM$_{2.5}$ samples on prebaked quartz
filters (with an area of 417.6 cm$^2$) for the determination of WSOC and HULIS. PM$_{2.5}$ samples were also collected using a five-channel spiral ambient speciation sampler (SASS, Met One Inc., USA) with a flow rate of 6.7 L min$^{-1}$. Wang et al. (2015) provided the details of the sample collection.

Meteorological data including wind speed (WS), temperature, relative humidity (RH), and precipitation were obtained from the China Meteorological Data Sharing Service System (http://cdc.cma.gov.cn/home.do, last access: 17 June 2017).

### 2.2 Source sampling

Residential biofuel and coal combustion emissions and vehicle exhaust, which are representative of typical emission sources around Beijing, were sampled in this study.

#### 2.2.1 Residential biofuel and coal combustion

Three typical types of biofuel, i.e., wheat straw, corn stover, and wood, were burned in an improved stove, which has an enclosed combustion chamber and a bottom grate and a chimney. The sampling procedures are described by Li et al. (2007, 2009) and are briefly summarized here. The water boiling test was used to simulate a common cooking procedure. The burning cycle included heating a specific amount of water from room temperature to its boiling point and then letting it simmer for a few minutes. Both the high-power and low-power phases were included in the burn cycle to simulate cooking practices of a typical household. The sampling period covered the entire cycle and lasted 20–30 min.

Five coal types were selected for source testing covering a wide range of maturity with volatile matter content varying from 9.6% to 32.4%. Two coal stoves were tested, including a high-efficiency heating stove that employs under-fire combustion technology and a traditional cooking and heating stove that employs over-fire combustion technology (Li et al., 2016). Four chunk coals and one briquette coal were burned in the high-efficiency stove and three chunk coals were burned in the traditional stove. Coal–stove combinations are presented in Table 2. To reduce the interference from igniting the fire, coal was ignited with a propane gas flame from a torch. Emission sampling covered from fire start to fire extinction and lasted 2 to 3 h.

Source testing of residential biofuel and coal combustion was performed at Beihang University. The test fuels were air-dried, and the results of their proximate and ultimate analyses are listed in Table S2 in the Supplement. An outline of the sampling system is shown in Fig. S2. The stove was placed into a chamber. Purified air was introduced into the chamber with a fan to provide dilution air. Emissions were extracted from the chimney with an exhaust hood and were diluted with purified air, cooled to no more than 5°C at ambient temperature, and then drawn through a sampling duct and exhausted from the laboratory. Both air flows were adjusted using frequency modulators to change fan speeds. The gas flow velocity in the sampling duct was measured by a pitot tube to be over 5 m s$^{-1}$. Flow was isokinetically withdrawn from the sampling duct with a probe and directed into the residence chamber. PM$_{2.5}$ samples were collected from the end of the residence chamber onto prebaked quartz filters with a diameter of 47 mm through PM$_{2.5}$ cyclones at a flow rate of 16.7 L min$^{-1}$.

---

**Table 1.** Summary of the concentrations of PM$_{2.5}$, carbon species, water-soluble ions, and ratios between compounds.

<table>
<thead>
<tr>
<th>Species</th>
<th>Average ± SD</th>
<th>Summer ± SD</th>
<th>Autumn ± SD</th>
<th>Winter ± SD</th>
<th>Spring ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>PM$_{2.5}$ (µg m$^{-3}$)</td>
<td>106 ± 89</td>
<td>98 ± 60</td>
<td>58 ± 48</td>
<td>150 ± 121</td>
<td>120 ± 76</td>
</tr>
<tr>
<td>OC (µg m$^{-3}$)</td>
<td>16.0 ± 15.8</td>
<td>8.5 ± 5.2</td>
<td>10.3 ± 7.4</td>
<td>28.9 ± 22.0</td>
<td>14.6 ± 10.8</td>
</tr>
<tr>
<td>EC (µg m$^{-3}$)</td>
<td>5.0 ± 4.8</td>
<td>3.3 ± 1.8</td>
<td>3.5 ± 2.9</td>
<td>7.8 ± 6.6</td>
<td>5.3 ± 4.7</td>
</tr>
<tr>
<td>OC/EC</td>
<td>3.6 ± 1.4</td>
<td>2.8 ± 0.8</td>
<td>3.8 ± 1.9</td>
<td>4.3 ± 1.2</td>
<td>3.3 ± 0.9</td>
</tr>
<tr>
<td>WSOC (µg m$^{-3}$)</td>
<td>6.5 ± 6.5</td>
<td>4.4 ± 3.6</td>
<td>5.2 ± 4.0</td>
<td>10.3 ± 9.8</td>
<td>5.9 ± 4.9</td>
</tr>
<tr>
<td>HULIS (µg m$^{-3}$)</td>
<td>7.5 ± 7.8</td>
<td>5.5 ± 4.4</td>
<td>5.6 ± 4.7</td>
<td>12.3 ± 11.7</td>
<td>6.5 ± 5.5</td>
</tr>
<tr>
<td>HULIS / PM$_{2.5}$ (%)</td>
<td>7.2 ± 3.3</td>
<td>5.9 ± 3.5</td>
<td>9.4 ± 3.1</td>
<td>7.9 ± 2.5</td>
<td>4.8 ± 1.7</td>
</tr>
<tr>
<td>HULIS$_C$ / OC (%)</td>
<td>24.5 ± 8.3</td>
<td>29.2 ± 6.2</td>
<td>26.2 ± 9.6</td>
<td>21.0 ± 7.1</td>
<td>22.0 ± 6.9</td>
</tr>
<tr>
<td>HULIS$_C$ / WSOC (%)</td>
<td>59.5 ± 9.24</td>
<td>66.7 ± 5.4</td>
<td>54.1 ± 11.2</td>
<td>62.3 ± 5.7</td>
<td>56.6 ± 6.3</td>
</tr>
<tr>
<td>SO$_4^{2-}$ (µg m$^{-3}$)</td>
<td>22.3 ± 24.1</td>
<td>22.6 ± 17.0</td>
<td>10.9 ± 13.2</td>
<td>32.7 ± 35.1</td>
<td>22.5 ± 16.5</td>
</tr>
<tr>
<td>NO$_3^-$ (µg m$^{-3}$)</td>
<td>18.6 ± 18.0</td>
<td>17.2 ± 13.4</td>
<td>10.8 ± 13.2</td>
<td>20.1 ± 17.8</td>
<td>29.0 ± 23.8</td>
</tr>
<tr>
<td>Cl$^-$ (µg m$^{-3}$)</td>
<td>4.2 ± 4.9</td>
<td>1.8 ± 1.9</td>
<td>1.3 ± 1.6</td>
<td>6.5 ± 5.7</td>
<td>7.9 ± 5.2</td>
</tr>
<tr>
<td>Na$^+$ (µg m$^{-3}$)</td>
<td>0.60 ± 0.51</td>
<td>0.40 ± 0.30</td>
<td>0.33 ± 0.41</td>
<td>0.89 ± 0.61</td>
<td>0.79 ± 0.36</td>
</tr>
<tr>
<td>K$^+$ (µg m$^{-3}$)</td>
<td>2.2 ± 2.6</td>
<td>2.2 ± 2.9</td>
<td>1.3 ± 1.0</td>
<td>3.2 ± 3.6</td>
<td>2.2 ± 1.3</td>
</tr>
<tr>
<td>Mg$^{2+}$ (µg m$^{-3}$)</td>
<td>0.18 ± 0.19</td>
<td>0.15 ± 0.07</td>
<td>0.18 ± 0.08</td>
<td>0.24 ± 0.32</td>
<td>0.10 ± 0.07</td>
</tr>
<tr>
<td>Ca$^{2+}$ (µg m$^{-3}$)</td>
<td>0.97 ± 0.57</td>
<td>0.99 ± 0.52</td>
<td>1.14 ± 0.48</td>
<td>0.83 ± 0.70</td>
<td>0.89 ± 0.46</td>
</tr>
<tr>
<td>NH$_4^+$ (µg m$^{-3}$)</td>
<td>14.1 ± 13.0</td>
<td>13.2 ± 9.8</td>
<td>6.6 ± 7.0</td>
<td>19.1 ± 16.9</td>
<td>18.4 ± 11.8</td>
</tr>
</tbody>
</table>
2.2.2 Vehicle exhaust

Four light-duty gasoline vehicles certified as meeting the China 4 emission regulations were tested for their emissions on a chassis dynamometer. The tests were conducted using the New European Driving Cycle (Marotta et al., 2015) and lasted 1180 s, including four repeated urban driving cycles and one extra-urban driving cycle. The emission testing and sampling systems are described in detail by Li et al. (2016) and are briefly summarized here. The vehicle exhaust was directed into a critical flow venturi constant volume sampler in a full flow dilution tunnel. The PM$_{2.5}$ samples were collected on prebaked quartz filters with a diameter of 47 mm through PM$_{2.5}$ cyclones at a flow rate of 80 L min$^{-1}$.

Three heavy-duty diesel trucks were selected to perform on-road emission tests. The tests were conducted on both highway and city roads. The emission testing and sampling systems are described in detail elsewhere (He et al., 2015) and are briefly summarized here. A micro proportional sampling system (SEMTECH-MPS; Sensors Inc., MI, USA) was used to draw a constant ratio of sample flow to exhaust flow and dilute the sample flow. PM$_{2.5}$ samples were collected onto prebaked quartz filters with a diameter of 47 mm through PM$_{2.5}$ cyclones at a flow rate of 10 L min$^{-1}$.

Tunnel measurements were also conducted to collect vehicle exhaust at the Badaling Tunnel in Beijing. The length of the tunnel is 1085 m. The ventilation in the tunnel was achieved by the flow of traffic into the tunnel during the sampling period. PM$_{2.5}$ samplers with prebaked 47 mm quartz filters were operated at a flow rate of 16.7 L min$^{-1}$ at the inlet and the outlet of the tunnel simultaneously. The sampling period was 2 h and the samples represent the mixed exhaust of gasoline-fueled vehicles and diesel-fueled vehicles.

All source samples collected on the quartz filters were analyzed for HULIS, WSOC, and OC/EC (organic carbon and elemental carbon) according the methods described in Sect. 2.3.

2.3 Chemical characterization

HULIS isolation was based on the extraction method developed by Varga et al. (2001) and used in many other studies (Nguyen et al., 2014; Lin et al., 2010b; Lin and Yu, 2011; Fan et al., 2012; Song et al., 2012; Salma et al., 2013; Feczko et al., 2007; Krivácsy et al., 2008). The separation procedure is provided in Text S1 of the Supplement. WSOC and HULIS$_C$ were determined using a total organic carbon (TOC) analyzer (Shimadzu TOC-VCPH, Japan) based on a combustion–oxidation, nondispersive infrared absorption method. The TOC was determined by subtracting inorganic carbonate (IC) from total carbon (TC): TOC = TC − IC. The reported data were the average results of three replicate measurements. Mass concentrations of HULIS were obtained from HULIS$_C$ by multiplying a scaling factor of 1.9 as
suggested by Lin et al. (2012), Kiss et al. (2002), and Zheng et al. (2013).

A 0.5 cm$^2$ punch from each quartz filter was analyzed for OC and EC using a DRI model 2001 thermal–optical carbon analyzer (Atmoslytic Inc., Calabasas, USA) following the IMPROVE-A thermal optical reflectance (TOR) protocol (Chow et al., 2007).

The PM$_{2.5}$ samples from SASS were also analyzed for mass and water-soluble inorganic ion analysis as described by Wang et al. (2015).

2.4 CMAQ modeling of primary HULISc

A source-oriented version of the Community Multiscale Air Quality (CMAQ) model (version 5.0.1) was used in this study to track primary PM$_{2.5}$ (PPM$_{2.5}$) from different emission sectors and determine the resulting concentrations of primary HULIS. The model was used in a previous study to determine source contributions to PPM$_{2.5}$ mass, EC, and primary OC (POC) in China. Details of the source apportionment technique can be found in Hu et al. (2015). In summary, source contributions to PPM$_{2.5}$ mass were directly determined using nonreactive source-specific tracers to track the emissions of PPM$_{2.5}$ from different sources. These nonreactive tracers were treated identically to the other PPM components when simulating their emissions, transport, and removal. A constant scaling factor (typically $10^{-4}$ or $10^{-5}$) was used to scale the actual emission rate of these tracers to ensure that their concentrations are sufficiently low that they do not alter the removal rates of other PM components. The concentrations and source contributions to EC and POC were determined during post-processing by using source-specific emission factors as well as the model-predicted source contributions to PPM$_{2.5}$ mass concentrations. This technique can be used to determine source contributions to primary HULIS.

For example, contributions of the $i$th emission source to primary HULISc concentration (HULISc$_{i}$) can be calculated using Eq. (1):

$$\text{HULISc}_{i} = \text{PPM}_{2.5,i} \cdot f_{\text{OC},i} \cdot f_{\text{HULIS},i},$$

where $f_{\text{HULIS},i}$ is the mass fraction of HULIS per unit of emission of POC from the $i$th source (see Sect. 3.3 below for estimation of HULIS primary emissions), $f_{\text{OC},i}$ is the mass fraction of POC per unit of emission of PPM$_{2.5,i}$ from the $i$th source, and PPM$_{2.5,i}$ is the calculated source contribution to PPM$_{2.5}$ from the $i$th source based on the nonreactive tracer. The total concentration of primary HULIS can be determined by adding the primary HULIS contributions from the different sources.

In this study, the model uses a $36 \times 36$ km horizontal resolution to cover a rectangular domain that includes all of China and neighboring countries. Source contributions to HULIS were calculated for the periods when observations of HULIS are available. Emissions from anthropogenic source sectors (residential sources, power plants, industries, and transportation) are based on the Multi-resolution Emission Inventory of China (MEIC) 2012 (http://www.meicmodel.org, last access: 7 May 2018). Open biomass burning was estimated using the FINN dataset from the National Center for Atmospheric Research (NCAR) (Wiedinmyer et al., 2011). Natural emissions from soil erosion and sea spray were modeled within the CMAQ model (Appel et al., 2013; Kelly et al., 2010). Biogenic emissions were estimated using the Model for Emissions of Gases and Aerosol from Nature (MEGAN) version 2.10. Meteorological fields were calculated using the Weather Research and Forecasting (WRF) model. Details of the model setup, input data preparation, and model evaluation for PPM$_{2.5}$ mass are documented by Hu et al. (2015). In this study, a comparison of predicted daily PPM$_{2.5}$ concentrations with observations was performed and only those predictions with fractional errors of less than 0.6 were included in the calculation of primary HULIS. The values of $f_{\text{OC}}$ for different source sectors used in the calculation are included in Table S4 of the Supplement. These values were used in Ying et al. (2018), and the predicted daily average POC and EC concentrations generally agree with predictions for both daily and annual average concentrations.

3 Results and discussion

3.1 General characteristics of ambient aerosol

The concentrations of PM$_{2.5}$, carbonaceous species (OC, EC, WSOC, and HULIS), and inorganic ions such as SO$_4^{2-}$, NO$_3^-$, NH$_4^+$, and K$^+$ are summarized in Table 1. The 24 h average PM$_{2.5}$ concentration for the sample set was $106 \pm 89 \mu$g m$^{-3}$, about 3 times the national annual air quality standard ($35 \mu$g m$^{-3}$). The highest concentration ($\sim 600 \mu$g m$^{-3}$) appeared on 12–13 January 2013 as reported in other studies (Quan et al., 2014; Tian et al., 2014; Zheng et al., 2015). The severe pollution events were always accompanied by high relative humidity and low wind speeds (Fig. 1). During the entire sampling period, severely polluted days (PM$_{2.5}$ concentrations $\geq 150 \mu$g m$^{-3}$) constituted about 22 %, while fair days (PM$_{2.5}$ concentrations $\leq 75 \mu$g m$^{-3}$) approached 50 %. The average PM$_{2.5}$ concentrations in summer, autumn, winter, and spring were $98 \pm 60, 58 \pm 48, 150 \pm 121$, and $120 \pm 76 \mu$g m$^{-3}$, respectively.

The average HULIS concentration for the study period was $7.5 \pm 7.8 \mu$g m$^{-3}$. This value is lower than the average value of $11.8 \mu$g m$^{-3}$ measured at a rural site in the PRD region that was heavily influenced by biomass burning (Lin et al., 2010b). However, it is higher than those measurements in the urban areas (about $5 \mu$g m$^{-3}$ in the PRD; Lin et al., 2010a; Kuang et al., 2015), urban Shanghai (about $4 \mu$g m$^{-3}$; Zhao et al., 2015), and urban Lanzhou (about $4.7 \mu$g m$^{-3}$; Tan et al., 2016). HULIS exhibited obvious seasonal variations as shown in Fig. 1 and Table 1. The seasonal average concentrations were $5.5 \pm 4.4, 5.6 \pm 4.7, 12.3 \pm 11.7$, and...
6.5 ± 5.5 µg m⁻³ in summer, autumn, winter, and spring, respectively. The winter mean was about twice the value in any other season, and the highest concentration (54.96 µg m⁻³) of HULIS was observed on the same day that the highest concentration of PM₂.₅ was observed. The mean HULIS concentrations were very similar between summer and autumn in contrast with PM₂.₅, which has much higher concentrations in the summer (Table 1). These seasonal variations were similar to those observed in Aveiro and K-puszta (Feckzo et al., 2007), but those annual average concentrations (about 2.4 and 3.2 µg m⁻³, respectively) were much lower than in Beijing. The concentrations of HULIS in previously reported studies are summarized in Table S1 of the Supplement.

HULIS and PM₂.₅ had a significant correlation with the annual \( r^2 = 0.90 \) (\( r^2 = 0.77, 0.96, 0.96, \) and 0.94 in summer, autumn, winter, and spring, respectively) (Fig. S4a). The seasonal average of HULIS / PM₂.₅ was 5.9 %, 9.4 %, 7.9 %, and 4.8 % in summer, autumn, winter, and spring, respectively. The annual average of HULIS / PM₂.₅ was 7.2 % ± 3.3 % and was approximately 10 % lower than that in the PRD region (Lin et al., 2010a).

Strong correlations of HULIS_C with OC were observed, with the annual \( r^2 = 0.87 \) (\( r^2 = 0.94, 0.82, 0.89, \) and 0.84 in summer, autumn, winter, and spring, respectively) (Fig. S4c). The percentages of HULIS_C in OC for summer, autumn, winter, and spring, respectively, were 29.2 % ± 6.2 %, 26.2 % ± 9.6 %, 21.0 % ± 7.1 %, and 22.0 % ± 6.9 % with an annual average of 24.5 % ± 8.3 %.

Strong correlations of HULIS_C with WSOC were also observed, with the annual \( r^2 = 0.98 \) (\( r^2 = 0.99, 0.96, 0.99, \) and 0.98 in summer, autumn, winter, and spring, respectively) (Fig. S4b). The percentages of HULIS_C in WSOC for summer, autumn, winter, and spring, respectively, were 66.7 % ± 5.4 %, 54.1 % ± 11.2 %, 62.3 % ± 5.7 %, and 56.6 % ± 6.3 %, with an annual average of 59.5 % ± 9.2 %, suggesting that HULIS_C was the major constituent of WSOC. This value is comparable to the results (about 60 %) at urban sites in the PRD region (Lin et al., 2010b; Fan et al., 2012), Shanghai (Zhao et al., 2015), South Korea (Park et al., 2012), Budapest (Salma et al., 2007, 2008, 2010), and high-alpine areas of Jungfraujoch, Switzerland (Krivácsy et al., 2001). However, it is higher than the rural areas in K-puszta, Hungary (Salma et al., 2010), and the northeastern US (Pavlovic and Hopke, 2012). The ratios of HULIS_C / WSOC reported by previous studies are listed in Table S1 of the Supplement.
3.2 HULIS emission characteristics from various sources

The measured HULISC/OC (i.e., $f_{\text{HULIS},i}$) and HULISC/WSOC ratios from the source samples are presented in Table 2. Biomass combustion produces a significant fraction of HULIS in OC (0.41–0.50) whether burning wood or crop straw. Those values are high compared to previous studies (see Table S3 of the Supplement). The HULISC/OC values obtained by Lin et al. (2010a, b) were 0.14 to 0.34 from rice straw and sugarcane open burning in the PRD region in south China. Park and Yu (2016) found the ratios from open-burning rice straw, pine needles, and sesame stems in South Korea were in the range of 0.15 to 0.29. Schmidl et al. (2018a, b) reported ratios of 0.01–0.12 for wood burned in the stove and 0.33–0.35 for leaves burning in the open in the mid-European Alpine region. Goncalves et al. (2010) obtained ratios of 0.04 to 0.11 from wood burned in the stove in Portugal. HULIS are an important component of water-soluble organic matter (WSOM). High HULISC/WSOC ratios (0.62 to 0.65) were observed for three types of biomass burning in this study. These results are comparable with two previous studies. Fan et al. (2016) reported the ratios from open-burning rice straw, corn straw, and pine branch were in the range of 0.57 to 0.66. Park and Yu (2016) obtained ratios in the range of 0.36 to 0.63 from open-burning three types of biomass. However, Lin et al. (2010a) reported relatively low values ranging from 0.30 to 0.33 from open-burning rice straw and sugarcane. Possible factors influencing HULISC/OC ratios were summarized in Table S3 of the Supplement. Combustion conditions have much influence on the HULIS-to-OC ratios. For biomass open burning, HULIS-to-OC ratios varied less (from 0.14 to 0.35), while for biomass burned in the stove, ratios varied a lot (from 0.01–0.50). For those advanced stoves used in Europe (with secondary air), combustion is relatively complete; thus HULIS were generated less often (0.01–0.12). However, for the stoves used in Chinese rural households, combustion is relatively inadequate; thus HULIS were generated more often (0.41–0.50). Dilution ratio (DR) and residence time (RT) could affect gas–particle partitioning and thus have an effect on the HULIS-to-OC ratios (Lipsky and Robinson, 2006; May et al., 2013). Moisture content of fuels was not found to be correlated with HULIS-to-OC ratios.

Residential coal combustion produces 5 % to 24 % of the OC as HULIS for all the coal–stove combinations in this study. Only one prior study measured HULIS emitted from residential honeycomb coal briquette combustion (Fan et al., 2016). However, the HULIS-to-OC ratio was not reported in that study. HULIS/WSOM ratios (0.46) in that study are comparable with our HULISC/WSOC data (0.41–0.62).

Light-duty gasoline and heavy-duty diesel vehicles also produced primary HULIS on the order of 5 % to 16 % of the emitted OC. The HULIS content detected in the vehicle exhaust samples was generally less than the detection limit for these measurements. Thus, these reported ratios of HULISC to OC for vehicle emissions have high uncertainties. Ratios of HULISC to OC for vehicle emissions obtained in this study are much higher than prior tunnel measurements (2 %–5 %) (El Haddad et al., 2009). However, they are comparable with those from residential coal combustion. Due to more complete combustion or more advance emission controls in vehicles than residential solid fuel combustion, OC emission factors for vehicles are normally around 2 orders of magnitude less than those for residential coal combustion (MEP of China, 2014), so HULIS emissions from vehicles can be neglected as described in Sect. 3.3.

3.3 Estimation of HULIS primary emissions

The average values of $f_{\text{HULIS},i}$ for residential biofuel and coal combustion and vehicle exhaust obtained from our measurements were used to assess the extent of primary emissions. Due to the lack of information on $f_{\text{HULIS},i}$ for the other sectors, such as power plants and industries, considering combustion/production technology and emission control technology, we assumed values for these two sectors to be 0.01 and 0.05, respectively.

Based on OC emissions for different sources in the MEIC inventory and the $f_{\text{HULIS},i}$ for the various sources described above, the annual anthropogenic primary emissions of HULIS in Beijing are estimated to be approximately 6.3 Gg with over 60 % of this primary HULIS being emitted during the heating season. Residential biomass and coal burning contribute about 70 % and 25 % of the annual primary HULIS emissions, respectively. Vehicle exhaust contributions to annual primary HULIS emissions are negligible (less than 2 %). Meanwhile the industry sector and power plants contribute about 3 % and close to zero of the annual primary HULIS emissions, respectively. In winter, residential biomass and coal burning contribute close to 98 % of primary HULIS (Table S5 of the Supplement).

Terrestrial and marine emissions were not included in these estimations of primary HULIS emissions since they were considered to be negligible for inland cities, such as Beijing (Graber and Rudich, 2006; Zheng et al., 2013). Cooking contributes about 20 % of ambient fine organic aerosols in Beijing (Wang et al., 2009; Zhang et al., 2018; Sun et al., 2016). Since cooking emissions were not included in MEIC, and no HULIS emission information about cooking is available, cooking is not considered in the current model. Cooking emissions might contribute to ambient HULIS and need to be explored in the future.
3.4 Possible primary sources of HULIS

Ambient HULIS sources include primary sources and atmospheric secondary processes that convert gaseous precursors to HULIS. The correlation between HULIS and other measured constituents provides information regarding possible sources and formation mechanisms of HULIS.

Correlations between HULIS and primary species in PM$_{2.5}$ are shown in Fig. 2. POC and secondary organic carbon (SOC) were estimated using the EC tracer method (Lim and Turpin, 2002; Turpin and Huntzicker, 1995). The details of the method and evaluation are provided in Text S2. Figure 2a and b show that there are strong correlations between HULIS and POC and HULIS and EC throughout the year, indicating that HULIS have sources and/or transport processes similar to those of POC and EC. Both POC and EC are emitted by the incomplete combustion of carbon-containing fuels. Thus, the correlation of HULISc with POC would be expected given its correlation with EC and the correlation between EC and POC. According to the 2010 MEIC data for Beijing 2010, biomass and residential coal burning contribute more than 80% of the POC emissions, the industrial sector contributes over 10%, and vehicular exhaust contributions are negligible. For EC emissions, residential coal burning contributes more than 50%; biomass burning, industry, and vehicles contribute the rest.

K$^+$ generally originates from biomass burning with lesser contributions from coal burning and dust. However, biomass burning is regarded as the most important source for K$^+$ and it is often used as an indicator of biomass burning (Kuang et al., 2015; Zhang et al., 2013; Park et al., 2015; Pio et al., 2008; Wang et al., 2011, 2012; Cheng et al., 2013). In northern China, biomass burning including residential cooking, heating, and open biomass burning occurred in all seasons (Cheng et al., 2013; Zheng et al., 2015). High K$^+$ concentrations in this study were observed with mean values of 2.2 ± 2.9, 1.3 ± 1.0, 3.2 ± 3.6, and 2.2 ± 1.3 µg m$^{-3}$ in summer, autumn, winter, and spring, respectively, and an annual average of 2.2 ± 2.6 µg m$^{-3}$. As shown in Fig. 3c, HULIS and K$^+$ exhibited a strong correlation with $r^2 = 0.76$, 0.73, and 0.61 in summer, autumn, and spring, respectively, suggesting the contribution of biomass burning to HULIS. During the winter sampling period, a low correlation was initially obtained ($r^2 = 0.21$). However, two extreme values of K$^+$ were observed on New Year’s Eve (9 February 2013, 14.6 µg m$^{-3}$) and during the Lantern Festival (24 February 2013, 17.6 µg m$^{-3}$). Prior studies had suggested that fireworks during the Spring Festival and Lantern Festival produce very high K$^+$ concentrations (Shen et al., 2009; Jing et al., 2014; Jiang et al., 2015). Excluding these two days (red points in Fig. 2c), the correlation between HULIS and K$^+$ increased to $r^2 = 0.73$, indicating the contribution of biomass burning to HULIS in winter. The strong correlation coefficient between HULIS and K$^+$ across all the seasons also confirmed that biomass burning was a significant primary HULIS emission source as presented in Sect. 3.3.

Cl$^-$ is usually believed to be associated with coal combustion and biomass burning (Yu et al., 2013; Gao et al., 2015; Yao et al., 2002; Li et al., 2007, 2009). A significant contribution from sea-salt particles for Cl$^-$ in PM$_{2.5}$ can be excluded since the average mole ratios of Cl$^-$ to Na$^+$ across four seasons in this study is 5.0, much higher than the ratio in seawater of 1.17. Moreover, the sampling site in Beijing is about 200 km from the sea. The correlation of HULIS and Cl$^-$ is shown in Fig. 2d. In winter and spring, HULIS are moderately correlated with Cl$^-$ with $r^2 = 0.56$ and $r^2 = 0.64$, respectively. Weaker correlations between HULIS and Cl$^-$ were observed in summer and autumn with $r^2 = 0.40$ and $r^2 = 0.43$, respectively. This result reflects the different amounts of coal burned in specific seasons. In winter and spring in northern China, coal combustion for heating was quite prevalent and more coal was burned compared to the other two seasons, resulting in the substantial emissions of gaseous and particulate pollutants, including HULIS and Cl$^-$: The correlation coefficient between HULIS and Cl$^-$ in winter and spring provides additional support for coal burning being an important primary HULIS emission source as discussed in Sect. 3.3. The strong correlation between HULIS and Cl$^-$ in winter ($r^2 = 0.89$) and weak correlation in summer ($r^2 = 0.17$) were also revealed in Lanzhou, another city in northern China (Tan et al., 2016). Significant correlations between HULIS and Cl$^-$ in wintertime urban aerosols from central and southern Europe were also found (Voliotis et al., 2017). The authors suggest the high concentration of HULIS during winter was probably related to residential coal burning (Tan et al., 2016; Voliotis et al., 2017).

Ca$^{2+}$ would more likely originate from the re-suspended road dust and long-range-transported dust (Gao et al., 2014). The poor correlation between HULIS and Ca$^{2+}$ (as shown in Fig. 2e) indicated dust was not likely to be an important source of HULIS.

3.5 HULIS source apportionment based on CMAQ modeling

CMAQ-predicted concentrations of PPM$_{2.5}$ from different sources were used to calculate HULISc from these sources using Eq. (1). The total concentration of primary HULIS can be determined by adding up primary HULIS from different sources. Figure 3 shows the predicted primary HULISc and observed HULISc concentrations with the prediction uncertainty. Only days with acceptable PPM$_{2.5}$ performance were shown in Fig. 3. Primary HULISc in January and March 2013 accounts for almost all observed HULISc in these two months. In summer and autumn 2012, predicted primary HULISc concentrations are approximately 1–2 µg m$^{-3}$. There were days when the observed HULISc concentrations were much higher than the predicted primary...
Figure 2. Correlations between HULIS and POC (a), HULIS and EC (b), HULIS and K$^+$ (c), HULIS and Cl$^-$ (d), and HULIS and Ca$^{2+}$ (e). Concentrations in four seasons are represented by different shapes with different colors. Linear regressions are also given with corresponding equations.
HULISC concentrations, indicating potential contributions of secondary HULISC.

Table 3 shows the seasonal contributions for each source as well as average source contributions for the whole sampling period to ambient HULISC in Beijing based on the observed total HULISC and CMAQ-predicted primary HULISC on the days with acceptable PPM$_{2.5}$ performance. Contributions of HULISC from secondary processes were determined by subtracting predicted primary HULISC from observed HULISC. For those days when the predicted primary HULISC concentrations are greater than the observed HULISC, the predicted primary HULISC concentrations were set to equal the observed HULISC and the secondary HULISC was set to zero. Based on the HULISC emissions from residential biofuel and coal burning described in Sect. 3.3, the contributions of biofuel and coal burning in the residential sector to ambient HULISC were estimated separately.

Overall, residential biofuel burning was the most important source of ambient HULIS, contributing nearly half of the ambient HULISC concentrations, much higher than those results from the PRD in southern China (less than 20 %) (Kuang et al., 2015). This difference is likely with greater biofuel burning during the heating seasons in the Beijing area. Residential coal burning contributes 15.1 ± 2.9 % to ambient HULIS and is also a significant source of ambient HULIS. Great contribution from the residential sector to ambient HULIS is consistent with the estimation of HULIS primary emissions and the correlations between HULIS and primary species previously presented. Vehicle emissions and other primary sources, such as industries, contribute negligible amounts to the ambient HULIS. Contributions from the residential sector display strong seasonal variations. In winter and spring, residential biofuel and coal burning accounted for about 80 % of the total HULISC while their contributions were reduced to approximately 40 % in summer and autumn. The seasonal variations were a reflection of the seasonal pattern of those activities in this region.

Secondary formation is estimated to have contributed an average of 38.9 ± 9.1 % to the HULISC concentrations and was another major source of ambient HULIS. However, our result is much lower than those results from the PRD in southern China (55 % to 69 %) (Kuang et al., 2015). The dif-
Concentration is driven by the differences in sources and climatological patterns between these two sites. There is much greater combustion for space heating in the colder north and atmospheric reaction rates will be higher in the warmer south. The contribution from secondary processes also shows obvious seasonal variations. In winter and spring, secondary processes accounted for 25% to 30% of the total HULISc with large uncertainties while their contributions were increased to 50.2% ± 19.3% and 63.2% ± 18.3% in summer and autumn. Higher secondary contributions were also found during warm seasons in the PRD region (Kuang et al., 2015). In addition to the proposed heterogeneous secondary formation pathways for HULISc, oxidation reactions initiated by chlorine (Cl) radicals can form SOA (Wang and Ruiz, 2017; Riva et al., 2015). Thus, Cl release by coal combustion may have the potential to contribute to HULISc, especially during winter when OH radical concentrations are much lower (monthly average 5.5 × 10⁻³ ppt for winter vs. 1.25 × 10⁻¹ ppt for summer based on CMAQ calculations for Beijing). However, the concentrations of secondary HULISc for winter estimated in this study are uncertain (1.8 ± 2.2 µg m⁻³) compared to the summertime average concentration (1.0 ± 0.4 µg m⁻³). Therefore, the role of Cl-initiated reactions producing HULISc cannot be definitively determined.

Figure 4 shows the scatter plot of predicted primary HULISc vs. observed HULISc concentrations. Moderate to strong correlations between predicted primary HULISc and observed HULISc were observed in winter and spring, while relatively weaker correlations were found in autumn. Moreover, low correlations were observed in summer. The variation of correlations between predicted primary HULISc and observed HULISc in different seasons also provides additional support for the relative importance of primary and secondary HULIS as shown in Table 3.

Data availability. Meteorological data are downloaded from http://cdc.cma.gov.cn/home.do (last access: 17 June 2017). Emissions from anthropogenic source sectors are downloaded from http://www.meicmodel.org (last access: 7 May 2018).

Supplement. The supporting information file for this paper provides the details of HULIS analytical procedures and prior literature regarding HULIS in the ambient aerosol. It also provides some additional figures and descriptions that help to support the analyses and discussion presented in the paper. The supplement related to this article is available online at: https://doi.org/10.5194/acp-19-2327-2019-supplement.

Author contributions. XL designed the experiments, analyzed the data, and wrote the paper. JH collected and analyzed the samples. PKH provided advice, supervision, and feedback throughout the drafting and submission process. JH provided samples from vehicle emissions. QS analyzed the samples. QC collected the samples. QY conducted the CMAQ modeling and wrote the paper.

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

Acknowledgements. This work was supported by the National Nature Science Foundation of China (grant nos. 41575119, 41275121) and the National Key Research and Development Program of China (no. 2017YFC0211404). The authors also want to acknowledge the Texas A&M Supercomputing Facility (http://sc.tamu.edu, last access: 10 May 2018) for providing computing resources useful in conducting the CMAQ simulations reported in this paper.

Edited by: Urs Baltensperger
Reviewed by: two anonymous referees

References


Krivácsy, Z., Kiss, G., Ceburnis, D., Jennings, G., Maenhaut, W., Salma, I., and Shooter, D.: Study of water-soluble atmospheric


X. Li et al.: Quantify HULIS sources using emission characterization and CMAQ 2339

www.atmos-chem-phys.net/19/2327/2019/ Atmos. Chem. Phys., 19, 2327–2341, 2019