Radiocarbon-based impact assessment of open biomass burning on regional carbonaceous aerosols in North China

Zheng Zong, Yingjun Chen, Chongguo Tian, Yin Fang, Xiaoping Wang, Guopei Huang, Fan Zhang, Jun Li, Gan Zhang

Key Laboratory of Coastal Environmental Processes and Ecological Remediation, Yantai Institute of Coastal Zone Research, Chinese Academy of Sciences, Yantai 264003, China
State Key Laboratory of Organic Geochemistry, Guangzhou Institute of Geochemistry, Chinese Academy of Sciences, Guangzhou 510640, China
Graduate University of Chinese Academy of Sciences, Beijing 100039, China

HIGHLIGHTS

• PM2.5 and TSP samples collected at Yellow River Delta were analyzed for OC and EC.
• OC, EC, TSP and PM2.5 concentrations were higher in daytime than in nighttime.
• Radiocarbon (14C) tracer, backward trajectories, and fire counts were used for the analysis.
• Agricultural waste open burning was a main contributor to summer PM2.5, OC and EC.

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ABSTRACT

Samples of total suspended particulates (TSPs) and fine particulate matter (PM2.5) were collected from 29th May to 1st July, 2013 at a regional background site in Bohai Rim, North China. Mass concentrations of particulate matter and carbonaceous species showed a total of 50% and 97% of the measured TSP and PM2.5 levels exceeded the first grade national standard of China, respectively. Daily concentrations of organic carbon (OC) and elemental carbon (EC) were detected 7.3 and 2.5 μg m−3 in TSP and 5.2 and 2.0 μg m−3 in PM2.5, which accounted 5.8% and 2.0% of TSP while 5.6% and 2.2% for PM2.5, respectively. The concentrations of OC, EC, TSP and PM2.5 were observed higher in the day time than those in the night time. The observations were associated with the emission variations from anthropogenic activities. Two merged samples representing from south and north source areas were selected for radiocarbon analysis. The radiocarbon measurements showed 74% of water-insoluble OC (WINSOC) and 59% of EC in PM2.5 derived from biomass burning and biogenic sources when the air masses were from south region, and 63% and 48% for the air masses from north, respectively. Combined with backward trajectories and daily burned area, open burning of agricultural wastes was found to be predominating, which was confirmed by the potential source contribution function (PSCF).

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1. Introduction

Carbonaceous aerosols are now of worldwide concern for their direct and indirect impacts on global and regional climate change, visibility degradation, air quality, and public health (Anenberg et al., 2012; Chung et al., 2012; Mahowald, 2011). Elemental carbon (EC) and organic carbon (OC) are two main fractions of carbonaceous species in the air. Particulate EC derives from the incomplete combustion of fossil fuels such as coal, gasoline, diesel, and biomass (i.e., vegetation and wood). Particulate OC originates from anthropogenic and biogenic sources, including primary organic carbon (POC) emitted from the incomplete combustion of carbon-contained material and primary biogenic source. While secondary organic carbon (SOC) formed through both photochemical oxidation of volatile precursors and subsequent gas-to-particle conversion processes (Fang et al., 2008). It is a critical job to quantify these sources for the development of effective and efficient air pollution control measures, especially in the areas with rapid economic growth and high population density where large amounts of those species are released into the air.

The Bohai Rim has become the third largest economic zone after the Yangtze River Delta and the Pearl River Delta in China (Xin et al., 2011). It is located in the northern China around Bohai Sea and comprises Beijing and Tianjin municipalities as well as part of Hebei, Liaoning and Shandong provinces. This region has been facing serious air pollution problems (including carbonaceous aerosols) (Boynard et al., 2014), due to dramatic increase in urbanization, industrialization and...
large-scale farming activities in the recent years. Some statistical methods, such as positive matrix factorization, eigenvector model and chemical mass balance model were used to identify the sources of carbonaceous aerosols in the region. However, significant difference was found from previous source apportionment assessments, such as about quarter \((Q. \text{ Wang et al., 2009})\) and half \((Cheng et al., 2013)\) of the carbon fractions in Beijing were associated with biomass burning processes. The uncertainty of the source apportionment results strongly depend on the temporal and spatial variations of emission sources such as source profiles, source strength, and non-linear environmental processes (atmospheric advection/diffusion and chemical reactions).

Radiocarbon \(^{14}\text{C}\) measurements provide a chance to differentiate between fossil and non-fossil sources of carbonaceous particles because the carbon isotopic composition in the ambient aerosol remains stable during atmospheric transport and chemical transformations \((\text{Sun et al., 2012})\). Recently, as a powerful tool, this method has been used to distinguish fossil and non-fossil sources in the Chinese cities \((\text{Chen et al., 2013; Huang et al., 2014; Liu et al., 2014})\) or rural sites \((\text{Liu et al., 2012})\). Recently, as a powerful tool, this method has been used to distinguish fossil and non-fossil sources in the Chinese cities \((\text{Chen et al., 2013; Huang et al., 2014; Liu et al., 2014})\) or rural sites \((\text{Liu et al., 2012})\). However, available knowledge of emission sources of carbonaceous aerosols on regional scale in the Bohai Rim is still poorly understood.

Previously monitoring data from a background area which has not been contaminated by certain preponderant source emissions are often used to assess the source apportionment of the regional pollution \((\text{Puxbaum et al., 2007; Wang et al., 2014; Zhang et al., 2010})\). In this work, total suspended particulate (TSP) and fine particulate matter (PM\(_{2.5}\)) samples were collected in summer, 2013 at the Yellow River Delta, North China. This is a regional background receptor site in the Bohai Rim, which is influenced by seasonally air pollution outflows from Beijing, Tianjin and Hebei (BTH) and Shandong, Henan and Jiangsu province (SHJ) under the East winter and summer monsoon. The objectives of this study are 1) to assess the concentration levels and intraday variations of particulate matter and carbonaceous species at a background area in Bohai Rim; 2) to discriminate and quantify carbon originating from fossil fuel and non-fossil sources by using radiocarbon \(^{14}\text{C}\) measurements in different carbonaceous sub-fractions; and 3) to assess the potential source area by the application of potential source contribution function (PSCF) and backward trajectories.

2. Methodology

2.1. Sample location

The sampling campaign was conducted in the Yellow River Delta Ecological Research Station of Coastal Wetland (37°45′N, 118°59′E), Chinese Academy of Sciences as shown in Fig. 1. The station is located at about 3 km south to Yellow River channel and about 20 km southwest to the mouth of the Yellow River \((\text{Han et al., 2014})\). The sampling period was from 29 May to 1 July, 2013. During the sampling time, prevailing wind was from the south and air parcels arriving in the Yellow River Delta were mainly from the SHJ region dominated by the East Asian summer monsoon, which was illustrated by the backward trajectories as shown in supporting information (SI) Fig. S1.

2.2. Sampling procedure

PM\(_{2.5}\) and TSP samples were collected by using a Tisch high volume sampler at a flow rate of 1.13 m\(^3\) min\(^{-1}\) and a high volume sampler at a flow rate of 0.3 m\(^3\) min\(^{-1}\), respectively. The PM\(_{2.5}\) and TSP samples were collected on separate quartz filters by two time-resolutions (12 h & 6 h) in the sampling campaign. The 12 hour sample was collected from 6:00 to 18:00 (day time) and from 18:00 to 6:00 (night time) the next day. The 6 hourly sampling was from 6:00 to 12:00, 12:00 to 18:00, 18:00 to 24:00, and 0:00 to 6:00, respectively. Two 6-hourly interval sampling activities were performed from 1st (18:00) to 4th (18:00) and from 18th (18:00) to 21st (18:00) in June, 2013, and 24 pairs of PM\(_{2.5}\) and TSP samples were collected. The other samples were collected by the 12-hourly sampling and 52 samples were collected. A total of 76 PM\(_{2.5}\) and 76 TSP samples were collected during the sampling period. All quartz filters were preheated at 450 °C for 6 h before sampling. The filters were wrapped in aluminum foil and sealed in polyethylene bags before and after the sample collection.

To obtain the mass concentrations of TSP and PM\(_{2.5}\) quartz filter fibers were analyzed gravimetrically using a Sartorius MC5 electronic microbalance with a ±1 μg sensitivity. These filters were weighted after 24-h equilibration at 25 °C and 39% relative humidity. Each filter was weighed for at least three times before and after sampling. The difference among the three repeated weighing was less than 10 μg for a blank filter and less than 20 μg for a sampled filter. After weighing, loaded filters were stored in a refrigerator at −20 °C before chemical
2.3. Thermal-optical carbon analysis

After measurement of PM$_{2.5}$ mass concentrations, carbonaceous components were analyzed by a Desert Research Institute (DRI) Model 2001 Carbon analyzer (Atmoslytic Inc., Calabasas, CA) following the Interagency Monitoring of Protected Visual Environment (IMPROVE) thermal/optical reflectance (TOR) protocol. The quartz filter collected particles was punched for specific size (0.544 cm$^2$) then placed onto the sample load position of the analyzer. The sample was heated to produce four OC fractions (OC1, OC2, OC3, and OC4) in four temperature steps (140, 280, 480, and 580 °C) under a non-oxidizing helium atmosphere, as well as three EC fractions (EC1, EC2, and EC3) in three temperature steps (580, 740, and 840 °C) under an oxidizing atmosphere of 2% O$^2$ / 98% He. At the same time, pyrolyzed organic carbon (OP) was produced in the inert atmosphere, which decreased the reflected light to correct for charred OC. Total OC is defined as the sum of the four OC fractions (OC1, OC2, OC3, and OC4) plus OP, and total EC is defined as the sum of the three EC fractions (EC1, EC2, and EC3) minus OP. The detection limits of the method for OC and EC were 0.82 and 0.20 μg cm$^{-2}$, respectively. 6 or 12 hour monitored data were averaged mathematically for the daily analysis.

Quality assurance and control (QA/QC) procedures were also performed during sample analysis. A sucrose solution of known concentration was tested to make sure that the error is within 5% every time before analyzing the samples. Blank filters and replicate samples were examined simultaneously in order to obtain their inherent concentrations on the filter and to evaluate measurement accuracy by using the same methods, respectively. Blank filters and replicate analyses were performed after analyzing a batch of 10 samples. The contribution of OC and EC from blank filters was < 3.5 and 0.6% of their respective average concentrations in aerosol samples for both TSP and PM$_{2.5}$, respectively. Comparison with average values from the replicate analyses showed a good precision with relative deviations of 5.7% and 4.9% for OC and EC in TSP, and 6.3% and 7.9% in PM$_{2.5}$.

2.4. $^{14}$C Analysis of the carbonaceous fractions

The measurement of the $^{14}$C in the OC and EC was performed using the OC/EC separation system described in the study of Liu et al. (2013). Briefly, with respect to OC, the filters were all extracted with ultrapure water to remove any interfering water-soluble inorganic and organic compounds. Therefore, the $^{14}$C values in the OC obtained in this work only represent the water-insoluble OC (WINSOC) and do not include the water-soluble OC. OC was isolated at 340 °C for 15 min after a flash heating of 650 °C for 45 s, which can minimize the char that is primarily produced by incomplete combustion of the OC. After the OC separation, the filters were removed from the system, placed into a muffle furnace at 375 °C, and combusted for 4 h in order to remove the char. Afterwards the filters were quickly put back into the system and oxidized under a stream of pure oxygen at 650 °C for 10 min to analyze the EC fraction. The carbon dioxide from WINSOC and EC was cryogenically trapped and sealed in a quartz tube for AMS target preparation. The preparation of graphite targets for accelerator mass spectrometry (AMS) analysis was performed using the graphitization line at the Guangzhou Institute of Geochemistry, CAS. The $^{14}$C/$^{12}$C ratios in the graphite samples were determined through NEC compact AMS at Peking University.

In this study, samples collected in two typical days prevailing north and south wind respectively were selected to analyze $^{14}$C of WINSOC and EC for studying the sources of carbonaceous aerosols.

2.5. Air trajectory generation and burned area

Backward trajectories and daily burned area were used to assess potential sources of observed carbonaceous aerosols in different air masses encountered during the sampling period. Backward trajectories were generated by the hybrid single-particle Lagrangian integrated trajectory (HYSPLIT) model, which is available on the National Oceanic and Atmospheric Administration Air Resource Laboratory website (www.arl.noaa.gov/hysplit4.html) (Draxler and Rolph, 2003). Backward trajectories (72-h) were generated with 6 h time intervals. The trajectories were calculated for air masses starting from the sampling site at 10 m above ground level. The fourth generation of the Global Fire Emissions Database (GFED4) burned area was used to assess agricultural waste burning (Giglio et al., 2013). The GFED4 gridded (0.25°) daily burned area data sets are available through the web site http://www.globalfiredata.org.

2.6. Potential source contribution function

The potential source contribution function (PSCF) was also used to assess potential source areas contributing high concentrations of PM$_{2.5}$, OC and EC in the PM$_{2.5}$ samples (Y.Q. Wang et al., 2009). PSCF can be briefly described as conditional possibility and characterizing the spatial distribution of possible geophysical source locations made out by using trajectories reaching to the sampling site. The ijth component of a PSCF field can be defined as given in Eq. (1) (Jeong et al., 2011):

$$ PSCF_{ij} = \frac{m_i}{n_{ij}} $$

where $n_i$ is the total number of end points that fall in the $i$th cell and $m_i$ is the number of endpoints for the $i$th cell with arrival times at the sampling site that correspond to each type of aerosol concentrations higher than an arbitrarily set criterion. The 75th percentile of the monitored concentrations of species of interest (OC: 6.82, EC: 2.96, PM$_{2.5}$: 110.90 μg m$^{-3}$) were utilized for PSCF calculations to identify the potential source areas.

In the PSCF method, cells with few endpoints may result in high uncertainty. Thus, to avoid uncertainties that can occur due to a limited amount of available data, an arbitrary weight function $W(n_{ij})$ as shown in Eq. (2) is multiplied with the PSCF value (Jeong et al., 2011).

$$ W(n_{ij}) = \begin{cases} 
1.00 & n \geq 24 \\
0.75 & 6 \leq n < 24 \\
0.42 & 3 \leq n < 6 \\
0.05 & n < 3 
\end{cases} $$

3. Results and discussion

3.1. Mass concentrations of TSP, PM$_{2.5}$ and carbonaceous species

Table 1 lists statistical values of daily mean concentrations for TSP, PM$_{2.5}$, and carbonaceous aerosols. During the sampling period, the mean concentration of TSP was 125.4 μg m$^{-3}$ and ranged between 46.1 and 198.5 μg m$^{-3}$, while the mean concentration of PM$_{2.5}$ was 92.3 μg m$^{-3}$ and ranged between 33.3 and 194.3 μg m$^{-3}$. Both monitored daily TSP and PM$_{2.5}$ concentrations exceeded the First Grade National Ambient Air Quality Standard (80 μg m$^{-3}$ for TSP and 35 μg m$^{-3}$ for PM$_{2.5}$) of China (Ministry of Environmental Protection of China: GB 3095-2012, www.zhb.gov.cn, 2012-02-29). The largest ratios of monitored concentrations to the first grade ambient air quality standard are 1.6 for TSP and 5.6 for PM$_{2.5}$. Total number of sampling days exceeding the standard values accounts for 50% for TSP and 97% for PM$_{2.5}$ during the whole sampling period. The ratio of PM$_{2.5}$ to TSP
was 0.75, ranging from 0.45 to 0.99, which indicates that fine particulate matter is a primary pollutant in the region. Daily mean concentrations of OC and EC were 7.3 and 2.5 μg m⁻³ in TSP, while 5.2 and 2.0 μg m⁻³ in PM₂.₅, respectively. OC and EC concentrations accounted for 5.8% and 2.0% of the TSP mass concentration, while 5.6% and 2.2% for PM₂.₅. The contributions of carbonaceous species to particulate matter were notably lower than in Chinese cities (Cao et al., 2007), and comparable with that at coastal background regions, such as at Tuqi Island (Wang et al., 2014). The contribution of OC in PM₂.₅ was slightly lower than that in TSP, while the contribution of EC was slightly higher. Our findings are in agreement with the Qinghai Lake region (Zhang et al., 2014a). The ratios of OC to EC were 3.1 (2.0–6.2) for TSP and 2.7 (2.0–4.0) for PM₂.₅, respectively. The signals consist with the finding that EC abundances in fine particulate matter (Duarte et al., 2008). In addition, the results of the Kolmogorov–Smirnov test shows that the particulate matter and carbonaceous aerosol concentrations were normally distributed, implying that they were supported by a relative stable source emission and/or environmental condition (Wang et al., 2014). The aspect was also characterized by respective smaller coefficient of variation as shown in Table 1.

Supporting information Table S1 showed a comparison of PM₂.₅, OC, EC concentration levels and OC/EC ratios in the present study with those previously reported from other coastal areas in China. It can be seen clearly that the summer PM₂.₅ concentration in the Yellow River Delta was higher than those in other coastal areas in China, whereas OC and EC concentrations were found lower, such as 10.2 μg m⁻³ and 5.5 μg m⁻³ in Tianjin (Gu et al., 2010), 9.3 μg m⁻³ and 2.3 μg m⁻³ in Shanghai (Feng et al., 2009), 9.6 μg m⁻³ and 2.2 μg m⁻³ in Xiamen (Zhang et al., 2011) and comparable with those in coastal background sites, such as 3.5 μg m⁻³ and 1.1 μg m⁻³ in Changdao Island (Feng et al., 2007), 5.7 μg m⁻³ and 1.7 μg m⁻³ in Tuqi Island (Wang et al., 2014), and 2.5 μg m⁻³ and 0.8 μg m⁻³ in Ningbo (Liu et al., 2013). It suggests the weak association of combustion sources to the levels of carbonaceous species in the Yellow River Delta and/or Chinese coastal background areas as compared to those in the coastal cities in China. On the other hand, the OC/EC ratio (2.7 ± 0.5) in the Yellow River Delta was higher than that in Tianjin (1.8), and was lower than that in the other southern sites listed in SI Table S1. It reflects that low temperature combustion emission, such as biomass combustion primary sources and/or the formation of secondary organic carbon contributed more carbonaceous aerosols in the Yellow River Delta than in Tianjin (Li et al., 2014).

3.2. Intraday fluctuation of TSP, PM₂.₅ and carbonaceous species

Table 1 also lists those statistical values of OC, EC, TSP and PM₂.₅ concentrations in the day time and night time. In general, these concentrations were higher in daytime than in nighttime. The average day-to-night ratios (D/N) for TSP and PM₂.₅ were 1.28 and 1.32, respectively, suggesting the relative strong emission of fine particulates in the day time, which is largely attributed to the anthropogenic activities. Similarly, the D/N ratios for OC and EC were 1.45 and 1.18 in TSP, and 2.54 and 2.09 in PM₂.₅, respectively. The difference between these D/N ratios is an indication that OC and EC originated from different sources. The slightly higher D/N ratio for OC than for EC in both TSP and PM₂.₅ may be ascribed to the formation of SOC under intense photochemical conditions in daytime, which is similar to those (1.4 and 1.1 for OC and EC) found in Shanghai (Cao et al., 2013). The D/N ratios for both OC and EC in PM₂.₅ were near twice that in TSP, indicating more notable intraday fluctuation of carbonaceous species in fine particles. To identify the co-emission and/or transport of these atmospheric species, daytime and nighttime concentration series were correlated and listed in SI Table S2. Generally, the high correlations of the atmospheric species suggested their strong co-emission and/or transport patterns. TSP and PM₂.₅ showed higher correlation with each other and also with carbonaceous species in the night time as compared to the day time. It suggests weak co-emission signal of particulate matter and carbonaceous species in daytime, which can be attributed to the contribution of emission unassociated with combustion (such as build activities, dust) on the particulate matter in the Yellow River Delta. The correlations between OC and EC in TSP were the same in the day and night, while the correlations of carbonaceous species in PM₂.₅ were lower in day time than in night time. The lower correlation in the day time is indicating a stronger formation of SOC in the day time than in the night time.

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<th>Table 1</th>
<th>Statistical values of TSP, PM₂.₅, OC and EC concentrations (μg m⁻³).</th>
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<td><strong>TSP</strong></td>
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<td><strong>CV</strong></td>
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*CV: Coefficient of variation.*
3.3. Source apportionment of carbonaceous species in PM$_{2.5}$

In order to compensate the excess $^{14}$C produced by nuclear bomb testing in 1950 and 1960s, the fraction of modern carbon ($f_{m}$) given by AMS was further converted into the fraction of contemporary carbon ($f_{c}$) by normalizing with a conversion factor of 1.06 and 1.10 for EC and OC (Liu et al., 2014). The $f_{c}$ values in the samples were defined as $f_{c} = f_{m}/1.10$ for EC, $f_{c} = f_{m}/1.06$ for OC, and the fraction of fossil ($f_{f}$) was defined as $f_{f} = 1 - f_{c}$. Two combined samples representing different sources were selected to analyze $^{14}$C of WINSOC and EC. They were collected on the 3rd, and 11th June, showing prevailing south and north wind, respectively. The first sample was merged by four 6-hourly samples, and the other sample was merged by two 12-hourly samples. The values of $^{14}$C in the WINSOC and EC for the samples with different wind directions are listed in Table 2.

The values for $f_{c}$ (WINSOC) ranged from 0.63 to 0.74 with a mean of 0.69, which indicates that contemporary carbon sources (biogenic and biomass burning) were the primary sources of WINSOC at the Yellow River Delta. The $f_{c}$ (EC) values ranged between 0.48 and 0.59 with a mean of 0.54, which were lower than the corresponding $f_{c}$ (WINSOC) values, demonstrating relative larger contribution of non-fossil source emission to WINSOC. Higher contributions of contemporary carbon sources to WINSOC than EC were also found at background sites in the Yangtze River Delta (56% and 48%) (Liu et al., 2013) and South China (83% and 62%) (Zhang et al., 2014b). The contributions of contemporary carbon sources to EC at these background sites were significantly lower than those in the Chinese cities, such as 17% in Beijing and Shanghai (Chen et al., 2013; Sun et al., 2012), 13% in Xiamen (Chen et al., 2013), and 29% in Guangzhou (Liu et al., 2014).

During the sampling period, the synoptic system at the sampling site was dominated by southerly or southeast wind under the East Asian summer monsoon as a back trajectory cluster map during the sampling period (Fig. 5). The $f_{c}$ (WINSOC) and $f_{c}$ (EC) were 0.74 and 0.59 on 3rd June, indicating 74% and 59% of the contribution of biogenic and biomass burning emission to the carbonaceous species, respectively, while the contribution of fossil source to WINSOC and EC only was 26% and 41%, respectively. On the day, the OC and EC concentrations reached a peak for both PM$_{2.5}$ (14.0 and 4.6 $\mu$g m$^{-3}$) and TSP (17.7 and 5.0 $\mu$g m$^{-3}$). The back trajectory showed that the air masses passed through the area with intensive open agricultural burning activities in the SHJ region under the south wind (Fig. 2). As the upwind area, the SHJ region has been estimated as a major emission area of biomass burning (Zhao et al., 2012). Open burning of agricultural wastes in summer in the SHJ region contributed largely to the concentration burden and atmospheric transport of carbonaceous aerosol in the Bohai Rim (Wang et al., 2014).

In comparison with WINSOC and EC on 3rd June, the concentrations of carbonaceous species were very low on 11th June, when the air masses come from the north and pass through the BTH region before reaching the sampling site as shown in Fig. 3. $^{14}$C results indicated that the contribution of biomass-burning and biogenic emissions to the WINSOC and EC accounted for 63% and 48%, respectively, which was 10% lower than the south wind. In recent years, the BTH region has been considered as the most severely polluted area in China and industrial emissions, coal burning and automobile emissions are considered as major contributors (Pui et al., 2014; Sun et al., 2014). The high contribution of biomass-burning emission suggests weak impact of carbonaceous aerosols emitted from the BTH region on the air quality at the sampling site.

The contemporary carbon concentrations of the WINSOC and EC were 3.7 and 2.7 $\mu$g m$^{-3}$, and the fossil fuel carbon concentrations of the WINSOC and EC were 1.3 and 1.9 $\mu$g m$^{-3}$ on June 3rd, prevailing south wind. While the contemporary carbon concentrations of the WINSOC and EC were 0.6 and 0.4 $\mu$g m$^{-3}$, and the fossil fuel carbon concentrations of the WINSOC and EC were 0.4 and 0.4 $\mu$g m$^{-3}$ on June 11th prevailing north wind. The huge difference of the concentrations can be attributed to the biogenic origin, such as biomass combustion in the SHJ region.

To confirm the contribution of open burning of agricultural wastes on carbonaceous aerosol, a scatter plot showing daily OC and EC concentrations against burned area during the sampling period was obtained and presented in Fig. 4. When the burned area is lesser, the linear of OC and EC is not obvious, though the linear grows with the burned area increasing. Supporting information Figs. S4–6 showed 72-h back trajectories and burned area on 4th, 8th and 13th June, 2013, respectively. High OC and EC concentrations were found when air masses passing through the sampling site along with agricultural waste burning, such as on 3rd (see Fig. 2), 4th (see SI Fig. S4) and 13th June (SI Fig. S5). Similarly, the relative low concentrations of OC and EC were presented when air masses via the sampling site were not experienced with burned areas, such as on 8th June (see SI Fig. S5). It shows that the significant contribution of open agricultural burning on the concentration levels of carbonaceous aerosols under a favorable meteorological condition.

3.4. Source area identification

The domain of interest extended from 104° E to 130° E and 30° N to 46° N, which was divided into 1836 grid cells of 0.5° × 0.5° latitude and longitude according to the geographical locations of 72-h back trajectories during the sampling period. Fig. 5 shows the spatial patterns of PSCF values of PM$_{2.5}$, and OC and EC in the PM$_{2.5}$ samples. In
general, the potential source areas with high PSCF values for PM$_{2.5}$, OC and EC were in the Southern side. Although the northern BTH region has been identified as a main source area in the Bohai Rim (Zhao et al., 2012), the prevailing southerly winds during sampling period deterred the atmospheric transport of pollutants emitted from the BTH region to the Yellow River Delta. Those areas bordering Shandong, Jiangsu, Anhui and Henan province were mainly identified as potential source areas for PM$_{2.5}$, OC and EC, in particular for EC. It indicates that the combustion emission in the area affected largely concentration levels of EC in the Yellow River Delta.

Fig. 6 shows accumulated burned area during the sampling period. Most intensive agricultural waste open burning activities were found in the contiguous area, which is in agreement with the finding as mentioned above. The spatial patterns of the potential source areas were slightly different for PM$_{2.5}$, OC and EC, which are attributed to their different origins. For example, Dongying city was considered as a potential source area with high probability for both PM$_{2.5}$ and OC as shown in Fig. 5. However, the city was not highlighted as a significant source area for EC. Dongying is located about 60 km southwest of the sampling site, and is the nearest city from the sampling site. It is a medium city, and the industry is relatively underdeveloped, which contributed to that the EC source, such as vehicle emission and combustion, was less, compared with other big cities. As a result, non-combustion emissions (such as buildings) and high OC emission (such as biomass burning)
in the city contributed partly fine particulate matter and OC burden in the Yellow River Delta.

4. Conclusions

In this study, carbonaceous aerosols in TSP and PM$_{2.5}$ samples collected at the Yellow River Delta, China in summer, 2013, were analyzed to assess the temporal variation of particulate matter and carbonaceous aerosol concentrations associated with source emissions and environmental factors. The TSP and PM$_{2.5}$ concentrations exceeded the corresponding national standards, and were considerably higher compared with other Chinese coastal areas. OC and EC concentrations were contributed to 5.8% and 2.0% of the TSP mass concentration and 5.6% and 2.2% for PM$_{2.5}$, respectively. The contribution was comparable to that at other background regions in Bohai Rim. The concentrations of particulate matter and carbonaceous species in the Yellow River Delta were higher in daytime than in nighttime, which was associated with emission of anthropogenic activities. Due to the costly measurement of $^{14}$C, two merged samples were carefully selected based on predetermined concentration levels of carbonaceous species and meteorological conditions. Demonstrated by the $^{14}$C and PSCF results, agricultural waste open burning in the contiguous area of the SHJ region was concluded as a main contributor of WISOC and EC concentrations in the Yellow River Delta in summer.

Controlling fossil-fuel combustion is highlighted as a major action to improve air quality in the cities of Bohai Rim, China (Chen et al., 2013; Sun et al., 2012). Our radiocarbon analysis indicates more pronounced signal contemporary carbon sources and the need for more stringent control of agricultural waste open burning emission on the regional scale. These data are very helpful to better constitute and implement relevant regulation policy associated with emissions to improve air quality efficiently in the Bohai Rim, China.

Acknowledgments

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Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version, at http://dx.doi.1016/j.scitotenv.2015.01.113.

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