Emission and Transport of Carbonaceous Aerosols in Urbanized Coastal Areas in China

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ABSTRACT

Elemental and organic carbon (EC and OC), the principal short-lived climate forcers, were measured in fine particulate matter (PM2.5) collected at urban and rural sites in continental edge in Southern China. The carbonaceous matter (CM) contributed an average of 28.5 ± 7.2% (1 SD) of the mass of PM2.5 in urban areas and 30.3 ± 8.2% in rural areas. The annual average OC concentrations in PM2.5 in urban and rural areas were 7.6 ± 4.3 and 5.7 ± 3.1 μg/m3, respectively; and the annual average EC concentrations were 2.4 ± 0.8 and 1.3 ± 0.7 μg/m3, respectively. The higher EC concentration in urban area than in rural area showed significant anthropogenic emissions to the urban atmosphere. EC and OC concentrations displayed good correlation in samples collected in urban area during winter monsoon season, suggesting a dominant emission source (mostly traffic-related) in urban area. The carbonaceous aerosol pollution in the rural coastal receptor area can be attributed to the local emission and transport of air pollutants from urbanized areas in the eastern part of China. The surface observation together with backward trajectory analysis, satellite imaging, and meteorological simulation indicate that air pollutants transported from emission hotspots in the urbanized Eastern China area had caused an increase in the concentration of carbonaceous aerosols in the rural continental edge by a factor of 2–3. This significant aerosol forcing of the Chinese outflow plume should be paid attention in the study of air quality and climate changes in Eastern/Southern Asia.

Keywords: Elemental carbon; Organic carbon; PM2.5; Urban emission; Dispersion.

INTRODUCTION

Elemental carbon (EC) has a graphite-like structure, which intensively absorbs solar radiation, and is probably the most significant agent of global warming after CO2 with a climate forcing of 0.4–0.9 W/m2, which is roughly one quarter to one half of the climate warming effect of CO2 (Ramanathan and Carmichael, 2008; UNEP and WMO, 2011). The atmospheric EC causes a net global warming up to 0.8°C (Chung and Seinfeld, 2005; Chung et al., 2005; Ramanathan and Carmichael, 2008). EC is emitted directly from combustion sources and undergoes little chemical transformation, which makes it a good indicator for primary anthropogenic air pollution (Xiao et al., 2011; Kim et al., 2011). Organic carbon (OC) is a mixture of hydrocarbons and oxygenate compounds, which constitutes the rest of the carbonaceous aerosols (Gu et al., 2010). OC can be emitted directly in particulate form (primary) or formed in the atmosphere from semi- and low-volatile products of chemical reactions involving reactive organic gases (secondary). Moreover, carbonaceous aerosols play important roles in the reproduction of biological organisms and can cause or enhance diseases (Pöschl, 2005).

The atmospheric lifetimes of EC and OC are about one week (UNEP and WMO, 2011). These rapid atmospheric turnovers mean that reductions in emissions of carbonaceous aerosols would result in a quicker effect on the climate than from reduction strategies just focusing on CO2, the key long-lived climate forcer (atmospheric lifetime of ~100 years). The major source categories of combustion-derived carbonaceous aerosols are fossil fuel (e.g., diesel traffic and coal for energy), and biomass combustion (e.g., wood and manure as fuel and agricultural crop residue burning) (Cao et al., 2006; Ho et al., 2006; Zhang et al., 2009b). Emission inventory models suggest that the anthropogenic activities in China might account for 25% of global EC emissions (Zhang et al., 2009a; Chow et al., 2010). The Chinese EC emission exerts a particularly strong influence on the climate in Asia where it rivals CO2. Secondary effects include worsen atmospheric brown clouds (UNEP and WMO, 2011), weakened Asian monsoons (Menon et al., 2002),
and melting of the Himalayan glaciers (Xu et al., 2009). Mitigation of Chinese emissions offers a possibility to slow down and delay the onset of dangerous anthropogenic interference with the air and climate system. However, emissions of carbonaceous aerosols increased by 14% during the last five years (Zhang et al., 2009a), and mitigation efforts are hampered by poor knowledge of emission sources and transport pattern.

It is suspected that urbanized areas of China significantly account for the emissions of carbonaceous aerosols (Avino et al., 2011; Shi et al., 2010). More than 70% of major cities and 40% of total population of China are located in coastal areas (Cao and Wong, 2007). Coastal development in China plays a leading role in the national economy, accounting for 60% of its industrial production and 55% of gross domestic product (GDP) (Wang, 1992; Cao and Wong, 2007). The industrialization and urbanization in coastal areas of China should be a great contribution to anthropogenic emissions. In this study, we measured EC and OC concentrations in fine particulate matter (PM$_{2.5}$, particles smaller than 2.5 μm in aerodynamic diameter) at urban and rural sites in Southern China continental edge. The aim is to characterize the emission and transport of carbonaceous aerosols in Eastern and Southeastern Asia.

**METHODS**

**Research Sites**

Xiamen, which is located in Southern China coastal area, is an ideal sampling site for intercepting Chinese continental outflow in southeastern pathway (Liu et al., 2003, Fig. 1(a)). The whole region has a southern subtropical monsoon climate, with an annual mean temperature of 20.8°C. During the winter monsoon season (winter and autumn), the northern airflow from inland China brings cold and dry air to Xiamen, whereas during the summer monsoon season (March to September), the southern airflow brings brisk wind and marine air mass. Annual precipitation is 1143.5 mm, 80% of which occurs from March to September.

The field sampling for PM$_{2.5}$ occurred at three urban sites (24°26’N, 118°5’E; 24°29’N, 118°6’E; 24°29’N, 118°2’E; 20 m high above ground level (AGL)) and two rural sites (24°37’N, 118°3’E, 20 m AGL; 24°49’N, 118°9’E, 10 m AGL) in Xiamen (Fig. 1(b)). Urban Xiamen situates on an island connecting East China Sea and South China Sea. The three urban sites were selected to cover the main roads/streets, downtown, and nearby residential areas. Meanwhile, west of the city is an industrial area including a coal-fired power plant. Potential emission sources nearby the sampling sites in the urban area thus included motor vehicles, industrial emissions, and residential homes. Rural areas in Xiamen stretch out along the Southern China coast, and mountain forest spreads over the north to west. The two rural sites (~20 and 40 km from urban Xiamen) have similar surrounding of water body, woods and a few farmlands. The rural sampling sites are considered to be under the influence from biogenic emission, biomass burning in a few local residences, soil dust particles from farmlands and local country roads.

**Samples and Measurements**

PM$_{2.5}$ was collected on quartz filters (90 mm, Whatman, Quartz Microfibre Filters, Kent, UK) mounted on a mid-volume atmospheric sampler at the flow-rate of 100 L/min for 12–48 h. The total air volumes through the sampler for each sample were between 72–288 m$^3$. A constant flow-rate within an uncertainty of ±3% was obtained during sampling periods. The atmospheric sampler (TH-150C-III, Wuhan Tianhong Instrument Co., Ltd, China) consists of a size selective inlet followed by impactors to obtain a cut efficiency of 2.5 ± 0.2 μm. Quartz filters were pre-combusted at 450°C for 4 h before sampling. The filters were weighed (0.1 mg accuracy) before and after sampling to determine the total mass of collected PM$_{2.5}$. Before weighting, the filters were dried in an electronic desiccator at 25°C and relative humidity of 50% for 24 h. Then the sample filters were stored in a refrigerator before analysis.

A total of 64 samples were collected at three urban and two rural sampling sites. A 2.0 cm$^2$ piece of filter was cut off from each sample and analyzed with an automated semi-continuous thermal-optical transmittance (TOT) carbon...
were reported as mass in a unit volume of air (μg/m³). Carbonaceous matter (CM) was estimated from

\[ CM = k \times OC + EC \]  

where “k” applied the values of 1.6 and 2.1 for urban and rural aerosols, respectively (Turpin and Lim, 2001).

RESULTS AND DISCUSSION

**Daily Levels of EC and OC in PM2.5**

Fig. 2 shows the observed concentration of EC and OC in PM2.5 in urban and rural areas of Xiamen. The EC concentrations in urban area were in the range of 1.4 to 4.9 μg/m³, giving an annual average of 2.4 ± 0.8 μg/m³ (1 SD). The EC concentrations in rural area were in the range of 0.4 to 2.8 μg/m³ (average 1.3 ± 0.7 μg/m³), which were lower than those values in urban area by a factor of 2. OC concentrations for PM2.5 in urban area varied from 1.6 to 15.5 μg/m³. The OC levels in rural area varied from 1.7 to 13.6 μg/m³, which showed no significant difference from the values in urban area (student t-test at the level of significance of 5%).

It was noted that the observed mean-PM2.5 of 53.4 ± 25.0 μg/m³ in Xiamen, although not representative for the annual mean due to the limited coverage, exceeded the United States’ National Ambient Air Quality Standard (NAAQS, http://www.epa.gov/air/particlepollution/standards.html) of annual-mean 15 μg/m³ for PM2.5. Fig. 3 shows the relationships between PM2.5 and carbonaceous aerosol concentrations. CM made the contributions of 28.5 ± 7.2% and 30.3 ± 8.2% to PM2.5 (CM/PM2.5) in urban and rural areas, respectively. The relatively high value of CM/PM2.5 in rural area indicated probably significant contribution of biogenic organic aerosols (e.g., secondary organic carbon formation from biogenic gases, pollen) in coastal forest under strong solar radiation in Southern China. PM2.5 concentration was well correlated with OC in both urban (r = 0.85, k = 5.3, p < 0.001) and rural areas (r = 0.81, k = 3, p = 0.002). This suggests that higher loading of fine particulate matter was mostly due to intensified emission of carbonaceous aerosols. CM/PM2.5 in urban area showed better positive correlation with EC (r = 0.51, k = 4.4, p = 0.006) than with OC (r = 0.16, k = 0.3, p = 0.420), suggesting that primary emission may mostly account for particulate air pollution in urban area. CM/PM2.5 in rural area showed considerable relationship with both OC (r = 0.61, k = 1.3, p = 0.037) and EC (r = 0.37, k = 5.3, p = 0.233). Both secondary and primary sources could be important for particulate air pollution in rural area.

**Emission of Carbonaceous Aerosols from Urbanized Coastal Area**

Fig. 4 shows the relationship between EC and OC for PM2.5. During winter monsoon season, EC and OC showed good correlation (r = 0.91) for samples in urban area, but poor correlation (r = 0.59) for samples in rural area. This suggested a dominant emission source for carbonaceous aerosols and small amount of secondary organic aerosol (SOA) in rural area during winter monsoon season. The samples from urban area in summer showed very low OC/EC ratio (1.8 ± 0.9), suggesting the dominant contribution from motor vehicle emissions. Measurements of OC and EC concentrations at roadside and road tunnel showed lower OC/EC ratio (~1) than other nearby sources, such as coal combustion, residential wood burning, and forest fire (2–10, Jones and Harrison, 2005; Gelencsér et al., 2007; Shen et al., 2010). Mitigation of motor vehicle emissions may be efficient to control air pollution in summer. However, the high OC/EC ratio (4.4 ± 0.8) in winter monsoon season suggests that the high levels of carbonaceous aerosols (Fig. 2) should be contributed by multiple types of emission sources.

The large variability in OC/EC ratio in rural area indicates the combination of two or more sources of CM in rural atmosphere. The possible sources of CM in rural areas include rural emissions, carbonaceous aerosols transported from urban areas, and SOA generated from atmospheric chemistry processes (Han et al., 2008; Wang et al., 2011). OC/EC ratio was higher in rural sites (5.6 ± 5.0) than urban sites (3.6 ± 1.5), suggesting lower fraction of vehicle emissions in rural area than urban area.

Fig. 5 shows average EC and OC concentration of PM2.5 in urban and rural areas of Xiamen (this study), in comparison with previously reported data for Beijing (Feng et al., 2006; Song et al., 2007), Guangzhou (Feng et al., 2006; Duan et al., 2007; Ding et al., 2011), Shanghai (Feng et al., 2006, 2009), Hong Kong (Duan et al., 2007), Europe and China (e.g., secondary organic carbon formation from biogenic gases, pollen) in coastal forest under strong solar radiation in Southern China.
Fig. 3. Relationships between EC and PM$_{2.5}$ (a), OC and PM$_{2.5}$ (b), EC and CM/PM$_{2.5}$ (c), and OC and CM/PM$_{2.5}$ (d) for samples in urban and rural areas of Xiamen.

Fig. 4. Correlation between EC and OC in PM$_{2.5}$ for urban and rural samples of Xiamen.

(Viidanoja et al., 2002; Lonati et al., 2007; Viana et al., 2007; Harrison and Yin, 2008), and North America (Na et al., 2004; Sawant et al., 2004; Park et al., 2005; Grover et al., 2008) in winter season. EC and OC concentrations in PM$_{2.5}$ in urban Xiamen showed higher margin value close to the concentrations in urban areas of Shanghai and Guangzhou, while lower margin value close to the concentrations in urban areas of Hong Kong. The EC and OC concentrations in the five cities in China were ranked in the following order: Beijing > Guangzhou and Shanghai > Xiamen and Hong Kong. It was noted that the similar linear distributions of

Fig. 5. Comparison of EC and OC concentration in PM$_{2.5}$ in Xiamen (this study) with those in other cities for urban and rural areas in winter season. Average EC and OC concentrations measured by TOT method for PM$_{2.5}$ in Beijing (Feng et al., 2006; Song et al., 2007), Guangzhou (Feng et al., 2006; Duan et al., 2007; Ding et al., 2011), Shanghai (Feng et al., 2006, 2009), Hong Kong (Duan et al., 2007), Europe (Viidanoja et al., 2002; Lonati et al., 2007; Viana et al., 2007; Harrison and Yin, 2008), and North America (Na et al., 2004; Sawant et al., 2004; Park et al., 2005; Grover et al., 2008) are shown for comparison. Each data point in this figure represents the average value of EC or OC concentrations measured in a specific site in available literature.
average EC-OC in China, Europe and Northern America in Fig. 5 can be probably explained by similar dominant emissions from traffic and industry sources in urban and surrounding rural areas in the three regions. Meanwhile, the EC and OC concentrations in both urban and rural areas in China were higher than those in Europe and North America. Locally emitted air pollutants may incorporate into the continental pollution plume from urbanized areas in China and transport to remote areas.

**Enhanced Outflow Plume from Urban Emission**

The 3-day backward trajectories during the sampling period in the rural area of Xiamen were shown in Fig. 6. Lower concentrations of EC (0.7 ± 0.3 μg/m³) and OC (4.0 ± 1.3 μg/m³) were observed when the air masses came from East China Sea, Japan, or South China Sea (Fig. 6(a–b)). Higher values (EC, 1.9 ± 0.4 μg/m³; OC, 8.0 ± 3.2 μg/m³) were observed when the air masses came from the eastern part of China (Fig. 6(c)) or urban Xiamen (Fig. 6 (d)). Intense emissions from industrial and urbanized hotspots in the eastern part of China (Beijing and surrounding urban clusters, and Shanghai and surrounding urban clusters) were highlighted in the annual EC inventory map of China (shaded area in Fig. 6(c)). The pollution of carbonaceous aerosols in the rural coastal area during winter monsoon season was mostly associated with the transport of pollutants from emission hotspots in the eastern part of China. PM$_{2.5}$ was considerably correlated with EC concentration ($r = 0.53$) for the high-trajectory-group of winter monsoon season (samples of Fig. 6(c)). This may be accounted by significant vehicle and industrial emissions in a regional atmosphere in the eastern part of China (Wang et al., 2011). Meanwhile, oceanic airflows occasionally made dispersion of pollutants from urban and industrial area of Xiamen to surrounding rural and coastal area during summer monsoon season. The pollution plumes transported from regional/local urbanized areas increased the concentration of carbonaceous aerosols in the rural coastal receptor area by a factor of 2–3 according to the comparison between the high- and low-trajectory-groups.

During 9–10 January 2011, the EC concentrations measured in rural Xiamen reached the highest (2.8 μg/m³), and aerosol optical depth (AOD) were recognized hotspots (AOD > 1) in Shanghai and surrounding urban clusters (Fig. 7(a)). The high AOD (> 0.3) observed in Southern China coastal area, South China Sea and East China Sea can be justified by the AOD hotspots in Shanghai and the southward wind, which was supposed to drive the air mass from Shanghai and surrounding areas to Southern China coastal area. This is supported by the observation of high

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**Fig. 6.** Three-day backward trajectories for the samples of rural area of Xiamen. Trajectories were grouped for lower level of EC (0.7 ± 0.3 μg/m³) and OC (4.0 ± 1.3 μg/m³) for winter (a) and summer (b) monsoon seasons, and higher values (EC, 1.9 ± 0.4 μg/m³; OC, 8.0 ± 3.2 μg/m³) for winter (c) and summer (d) monsoon seasons. Backward trajectories were produced by HYSPLIT model (Draxler and Rolph 2011). Shaded area in (c) indicates high emission rate of EC (2756–10063 tone carbon per year in 1° longitude and latitude grid in the annual emission inventory map at [http://www.cger.uiowa.edu/EMISSION_DATA_new/] in the eastern part of China.
EC concentrations at rural Xiamen during the 9–10 January 2011. The Mongolia high pressure system provided the driving force for the air mass traveling from the industrial and urbanized pollution hotspots in the eastern part of China to south and east. This explained the high AOD belt along coastal areas, South China Sea, and East China Sea for the high-trajectory-group (Fig. 7(b)). The average AOD map and our observation of carbonaceous aerosols intercepted at the Southern China continental edge showed significant aerosol forcing of Chinese outflow plume through the incorporation of urban emissions.

CONCLUSION

Carbonaceous aerosols emitted from urbanized coastal area not only contribute to urban air pollution, but also transport to the surrounding rural and ocean areas and significantly influence the air quality remotely. The transport of air pollutants from local and regional urbanized hotspots in the eastern part of China caused an average increase in EC concentration from 0.7 ± 0.3 to 1.9 ± 0.4 μg/m³, and an average increase in OC concentration from 4.0 ± 1.3 to 8.0 ± 3.2 μg/m³ at the rural area in Southern China continental edge. Both urban and rural areas of China showed higher concentrations of carbonaceous aerosols than those in Europe and North America, suggesting the regional dispersion of air pollution plume over Chinese continent and the surrounding region. Although much more study is necessary, the surface observation at the rural coastal receptor area, backward trajectories analysis and satellite imaging indicate 2–3 times of aerosol forcing of Chinese outflow plume by industrial and urbanized emissions. The atmospheric brown clouds over Southeastern Asia could be controlled through the effective management of anthropogenic emissions in urbanized coastal areas.

ACKNOWLEDGEMENTS

We thank Dr. Shun-cheng Lee, Dr. Òrjan Gustafsson for valuable discussion of this work. Mr. Wenquan Wu and Jieru Zhang helped on field work. This research was financially supported by National Natural Science Foundation of China (no. 40905065), the Knowledge Innovation Program of the Chinese Academy of Sciences (no. KZCX2-YW-422-4), and Public Interest Program of Chinese Ministry of Environmental Protection, Grant No. 201009004. The authors gratefully acknowledge the NOAA Air Resources Laboratory (ARL) for the provision of the HYSPLIT transport and dispersion model used in this publication. NCEP reanalysis data was provided by the NOAA/OAR/ESRL PSD, Boulder, Colorado, USA, from their Web site at http://www.esrl.noaa.gov/psd/. MODIS AOD map used in this paper were produced with the Giovanni online data system, developed and maintained by the NASA GES DISC.

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