Decreasing concentrations of carbonaceous aerosols in China from 2003 to 2013

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Carbonaceous aerosols were characterized in 19 Chinese cities during winter and summer of 2013. Measurements of organic carbon (OC) and elemental carbon (EC) levels were compared with those from 14 corresponding cities sampled in 2003 to evaluate effects of emission changes over a decade. Average winter and summer OC and EC decreased by 32% and 17%, respectively, from 2003 to 2013, corresponding to nationwide emission control policies implemented since 2006. The extent of carbon reduction varied by season and by location. Larger reductions were found for secondary organic carbon (SOC, 49%) than primary organic carbon (POC, 25%). PM2.5 mass and total carbon concentrations were three to four times higher during winter than summer especially in the northern cities that use coal combustion for heating.

Carbonaceous aerosols, including organic carbon (OC) and elemental carbon (EC), are important for their adverse effects on visibility, climate, and human health1–5. OC is one of the largest constituents of suspended particulate matter (PM) mass and originates mostly from combustion processes. While EC derives from incomplete combustion, OC results from both direct emissions and atmospheric reactions of organic gases that form condensable compounds6,7. Carbonaceous aerosols influence the climate directly by scattering and absorbing incoming solar radiation, and indirectly by acting as cloud condensation nuclei and/or ice nuclei, which modify the microphysics, radiative properties, as well as lifetime and extent of clouds8. Light scattering by OC cools both the Earth’s surface and the atmosphere8, while light absorption by EC heats the atmosphere, possibly reducing cloud cover (termed the “semidirect effect”)10. Carbonaceous aerosol in the 0.1 to 1.0 µm size range can be transported and deposited in the human respiratory tract and is associated with cardiopulmonary mortality4,5,11,12. Positive associations of OC and EC with cardiovascular and respiratory mortality are found in urban environments13.

Concerns over climate and health effects have prompted efforts to reduce carbonaceous aerosols, including improvements for vehicle engines and exhaust-treatment technologies, better fuel quality, traffic management optimization, and implementation of stringent emission standards. Environmental protection policies prior to 2005 were not effective for reducing source emissions14,15. However, the 11th Five-Year Plan (2006–2010) set quantitative milestones for environmental and energy targets, including the “total emission control on SO2 [sulfur dioxide]” and an “energy saving” policy. These measures were maintained and extended in the 12th Five-Year Plan (2011–2015), which resulted in decreased ambient SO2 and PM10 emissions and concentrations14. Although a 10% reduction of SO2 and a 20% reduction of energy consumption per unit of gross domestic product (GDP) were reported by the end of the 11th Five-Year Plan (2006–2010)14, these reductions are somewhat offset by population and economic growth. From 2003 to 2013, fossil fuel consumption increased by 96.8%, with a 141% increase in vehicle-kilometers travelled (VKT)16. VKT increased faster in large cities due to rapid urbanization and motorization. The PM chemical composition has also changed over time due to changes in the emissions mixture. Increased flue gas desulfurization (FGD) applied to power and industrial sectors and the shift from gasoline to natural gas in some vehicle engine exhausts might increase emissions of volatile organic compounds (VOCs) and nitrogen oxides (NOx)17–19. Enhanced emissions from gasoline and liquefied petroleum gas (LPG) vehicles in Hong Kong were related to increases in VOC and ozone (O3) concentrations exceeding the ambient air quality objectives20.
Biases of 10–20% have been reported for different OC and EC measurements due to variations in sampler configuration, sampling sites, monitoring periods, and carbon analysis protocols. To obtain long-term trends, comparable sampling and analysis methods are needed.

Nationwide measurements of PM$_{2.5}$ mass (particles with aerodynamic diameter less than 2.5 µm), OC, and EC started in 14 Chinese cities during the summer and winter of 2003 to establish a baseline. This paper presents results from a follow-up study in 2013 for 19 cities, including the original 14. Comparison between 2003 and 2013 measurements are made to evaluate carbonaceous aerosol changes over the intervening decade.

Method

Sampling sites and descriptions. Nineteen cities (Supplemental Table S1) including five megacities (i.e., Beijing [BJ], Tianjin [TJ], Shanghai [SH], Chengdu [CD], and Chongqing [CQ]) were selected to represent economically developed and developing urban regions, as shown in Fig. 1. The sampling network is documented in Table 1. In addition to those from the 2003 study, three northern cities were added that represent deve-
| City name                      | Code | Location   | Region description                                                                 | Site type                                                                 | Sampling site description                                                                 |
|-------------------------------|------|------------|------------------------------------------------------------------------------------|----------------------------------------------------------------------------|--------------------------------------------------------------------------------------------|
| Northern cities (n = 10)      |      |            |                                                                                   |                                                                            |                                                                                             |
| Changchun, Jilin Province     | CC   | 43.9° N, 125.3° E | Northeastern old industrial bases, An urban-commercial site in a continental and developed industrial city | Roof (6 m) of a building at Jilin University                               |                                                                                             |
| Urumqi, Xinjiang Uygur Auto- | UR†  | 43.5° N, 87.4° E | Northwest China, An urban-commercial site in a continental developing city, far from the ocean | Roof (10 m) of a building at the Institute of Desert Meteorology           |                                                                                             |
| Beijing Municipality          | BJ   | 39.9° N, 116.4° E | Beijing–Tianjin–Hebei region, An urban-commercial site in the capital of China, megacity | Roof (14 m) of a building at Institute of Atmospheric Physics, Chinese Academy of Sciences (CAS) |                                                                                             |
| Tianjin, Hebei Province       | TJ   | 39.1° N, 117.2° E | Beijing–Tianjin–Hebei region, An urban site in a developed industrial city         | Roof (20 m) of a building at Nankai University                              |                                                                                             |
| Jinchang, Gansu Province      | JC   | 38° N, 101.1° E | Surrounded by the Gobi Desert, An urban site in the Asian dust source region, non-urban desert | Roof (10 m) of a building at Jinchang Meteorological Bureau                |                                                                                             |
| Yulin, Shaanxi Province       | YL   | 38.3° N, 109.8° E | The juncture of loess plateau and Maowusu desert, An suburban site in a continental developing city, close to a desert | A observation tower (10 m) in Shaanxi Desert Institute                   |                                                                                             |
| Taiyuan, Shanxi Province      | TP†  | 37.5° N, 112.3° E | Coal base, An urban-commercial site in a continental and developing industrial city | Roof (20 m) of a building at Taiyuan University of Technology            |                                                                                             |
| Qingdao, Shandong Province    | QD   | 36° N, 120.3° E | Coastal region, An urban site in a developing coastal city                       | Roof (10 m) of a building at Chinese Ocean University                     |                                                                                             |
| Xining, Qinghai Province      | XN†  | 36.4° N, 101.5° E | The Northeastern Tibetan Plateau, A roadside-commercial site in a continental developing city, high altitude | Roof (18 m) of a building at Xining Environmental Monitoring Center       |                                                                                             |
| Xi’an, Shaanxi Province       | XA   | 34.2° N, 108.9° E | Fen Wei Plain, An urban site in a continental and developing industrial city      | Roof (10 m) of a building at Institute of Earth Environment, CAS          |                                                                                             |
| Southern cities (n=9)         |      |            |                                                                                   |                                                                            |                                                                                             |
| Nanjing, Jiangsu Province     | NJ   | 32° N, 118.5° E | Yangtze River Delta region, An urban-commercial site in a developing city         | Roof (80 m) of a building at Nanjing University                           |                                                                                             |
| Shanghai Municipality         | SH   | 31.2° N, 121.4° E | Yangtze River Delta region, An urban site in an industrial and commercial megacity | Roof (8 m) of a building at Donghua University                            |                                                                                             |
| Hangzhou, Zhejiang Province   | HZ   | 30.2° N, 120.1° E | Yangtze River Delta region, An urban site in an developing continental city       | A substation (20 m) at Hangzhou Environmental Monitoring Station         |                                                                                             |
| Wuhan, Hubei Province         | WH   | 30.5° N, 114.2° E | Jianghan Plain, An urban site in an industrial and commercial city               | Roof (8 m) of a building at Chinese University of Geosciences             |                                                                                             |
| Chengdu, Sichuan Province     | CD†  | 30.4° N, 104° E  | Sichuan Basin, An urban-commercial site in a continental developing city         | Roof (18 m) of a building at Chengdu Branch, Chinese Academy of Sciences  |                                                                                             |
| Chongqing Municipality        | CQ   | 29.5° N, 106.5° E | Sichuan Basin, An urban-commercial site in a continental and developing industrial city | Roof (10 m) of the Chongqing Academy of Environmental Sciences          |                                                                                             |
| Xiamen, Fujian Province       | XM   | 24.4° N, 118.1° E | Fen Wei Plain, An urban site in a coastal and commercial city, developing city    | Roof (8 m) of a building at Xiamen University                             |                                                                                             |
| Guangzhou, Guangdong Province | GZ   | 23.1° N, 113.2° E | Pearl River Delta, An urban site in an industrial and commercial megacity       | Roof (10 m) of a building at Guangzhou University                        |                                                                                             |
| Hong Kong Special Administrative Region | HK | 22.2° N, 114.1° E | Pearl River Delta/Coastal region, southern China, A roadside-commercial site in a coastal and commercial city, developed city | A monitoring site (10 m) at Hong Kong Polytechnic University              |                                                                                             |

Table 1. Location and description of sampling sites in 19 cities for 2013 and 14 cities for 2003. *The five cities that are not included in the 2003 study24.

opining industries (Taiyuan, TY), high altitudes (Xining, XN), and arid western (Urumqi, UR) environments, along with two southern cities that represent developing urban environments (Chengdu, CD and Nanjing, NJ). Sampling sites represent urban-scale exposures, with most located at university campuses or research centers (> 100 m from local sources such as major roadways).

Sample collection. Integrated daily, 24 h PM$_{2.5}$ sampling (0900 to 0900 local standard time) was conducted during winter (5–26 January) and summer (1–31 July) of 2013 (Table S2). PM$_{2.5}$ samples were collected on 836 prefired (900 °C for 3 h) 47 mm Whatman QMA quartz-fiber filters, using mini-vol air samplers (Airmetrics, Eugene, OR, USA) at a flow rate of 5 L min$^{-1}$. These samplers were located on rooftops at varying heights (~ 6–20 m) above the ground level (Table 1). Quartz-fiber filters were analyzed gravimetrically for mass concentrations$^{25}$ using a Sartorius MC5 electronic microbalance with a ± 1 μg sensitivity (Sartorius, Gottingen, Germany).

To minimize particle volatilization and aerosol liquid water biases, these filters were weighed after 24-h equilibration at a constant (within ± 2 °C) temperature between 20 and 23 °C and within ± 5% relative humidity (RH) between 30 and 40% following U.S. EPA guidelines$^{26}$. Nominal values of 20 °C and 30% RH best conserve particle deposits during sample weighing$^{26}$. Each filter was weighed at least twice before and after sampling to...
ensure reproducibility. The differences between the replicated weights were < 10 μg per filter for laboratory blanks and < 20 μg for exposed samples.

**Thermal/optical carbon analysis.** A 0.5 cm² punch from each quartz-fiber filters was analyzed for eight carbon fractions following the IMPROVE_A (Interagency Monitoring of Protected Visual Environments) thermal/optical reflectance (TOR) protocol³²⁻³⁵ using a DRI Model 2001 Carbon Analyzer (Atmoslytic Inc., Calabasas, CA). This protocol reports four OC fractions (OC1 to OC4 at 140 °C, 280 °C, 480 °C, and 580 °C, respectively, in a 2% oxygen [O₂]/98% He atmosphere); and three EC fractions (EC1 to EC3 at 580 °C, 740 °C, and 840 °C, respectively, in a 100% helium [He] atmosphere); and three EC fractions (EC1 to EC3 at 580 °C, 740 °C, and 840 °C, respectively, in a 2% oxygen [O₂]/98% He atmosphere). A laser beam at 633 nm monitored changes in optical reflectance and estimated pyrolyzed carbon (OP). IMPROVE_TOR OC is the sum of four OC fractions plus OP, whereas EC is the sum of three EC fractions minus OP³⁶, with total carbon being the sum of OC and EC. Average field blanks were low, 0.97 and 0.09 μg cm⁻² for OC and EC, respectively.

**Estimation of primary and secondary OC (POC and SOC).** While particulate EC is directly emitted from primary sources, particulate OC originates from both direct particulate emissions (primary OC, POC) and conversion of directly emitted gases to particles (secondary OC, SOC). POC includes directly emitted particulate OC and vapors condensed onto the particulate phase as the emissions cool³⁰. SOC starts its atmospheric life in the gas phase as VOCs which undergo chemical transformations to less volatile compounds that shift to the particulate phase³⁰. Although POC tends to dominate in most polluted areas, SOC can exceed POCs contribution during haze episodes. Turpin et al.³¹ reported ~ 80% of OC as SOC during photochemical smog event in southern California, USA. Huang et al.³² found 44–71% SOC in organic aerosols in four Chinese cities (i.e., Beijing, Shanghai, Xi’an, and Guangzhou).

Multiple methods have been reported to estimate SOC. Wu and Yu³³ propose the use of the minimum R squared (MRS) method, as the MRS method presents a quantitative criterion to determine the primary OC and EC ratio (i[(OC/EC)ₚ]*i, which minimizes the uncertainties in estimating SOC. In comparing five linear regression techniques, Wu and Yu³³ further recommended the use of Deming regression, orthogonal distance regression, and York regression to estimate SOC.

As formation of SOC, often referred to secondary organic aerosol (SOA), involves complicated atmospheric processes of photochemical oxidation, gas/particle partitioning and nucleation/condensation, it presents a large challenge to accurately estimate SOC. There are pros and cons of applying different methods to estimate SOC with uncertainties associated with each. The underlying principle of the tracer approach³¹ is based on the assumption of a fixed relationship between primary OC and EC, while in reality, the OC and EC ratios and background concentrations vary over time. To ensure adequate validity of the ([OC/EC]ₚ)*i data from extreme weather conditions (e.g., rain or storms) and from combustion dominated environment (with high OC/EC ratios) are excluded, and OC/EC measurements with the lowest potential in photochemical production (i.e., lowest 20% of OC/EC ratios) are retained to estimate (OC/EC)ₚ*³³⁻³⁶. The EC tracer approach used in this study intends to maintain consistency in SOC comparison with the 2003 study.

Table S3 summarizes linear regressions between OC and EC, with similar findings for both years. During winter, good OC/EC correlations (R²: 0.66–0.95) were found for all but two northern industrial cities (i.e., Changchun and Xi’an), consistent with contributions from a mixture of sources (e.g., residential and commercial coal combustion, and motor vehicle exhaust). Correlations between OC and EC (R²: 0.16–0.92) were variable in summer, implying different source mixtures.

The OC/EC slopes were 2.34 and 2.08 in winter for the northern and southern cities, respectively, similar to those of 2.81 and 2.10 in 2003. As shown in Table S4, the summer OC/EC ratios of 0.97 and 0.51 during 2013 in the northern and southern cities are lower than the corresponding 2003 values of 1.99 and 1.29.

**Meteorological measurements.** Table S5 summarizes average meteorological measurements by season for 2003 and 2013. Besides rainfall, cities in northern and southern China showed similar levels of atmospheric pressure, temperature, relative humidity (RH), and wind speed between 2003 and 2013 for both seasons. An increase in precipitation was found for both northern and southern cities during the summer of 2013. Summertime rainfall in Xiamen, Guangzhou, Chongqing, Beijing, and Yulin exceeded by twofold that in 2003. In contrast, precipitation decreased during the winter of 2013 in southern cities. There were over twofold higher rainfalls in 2003 for Xiamen, Guangzhou, Shanghai, Nanjing, and Chengdu. Average precipitation for northern cities were 6 and 4.5 mm for 2003 and 2013, respectively, generally lower than 10 mm per city.

**Results and discussion**

**Seasonal variations of OC and EC in 2013.** Table 2 summarizes average PM₁₅ mass and carbon concentrations by season for each city and region. Average wintertime PM₁₅ mass (167.6 ± 97.9 μg m⁻³) in the north were ~ 23% higher than in the south (135.7 ± 68.0 μg m⁻³). Wintertime PM₁₅ mass concentrations were ~ 2.8–3 times higher than summer levels (43.8–60.9 μg m⁻³). Carbonaceous aerosol (1.6 × OC plus EC, with a factor of 1.6 to compensate for unmeasured oxygen and hydrogen³⁰) accounted for 30–40% of the PM₁₅ mass. Average northern OC concentrations were 34.7 ± 21.4 μg m⁻³ in winter, ~ 50% higher than for the southern region (22.9 ± 11.6 μg m⁻³), and five times those of the summer OC (6.8 ± 3.6 μg m⁻³). EC concentrations were over twice as high in winter (8.5 ± 5.7 μg m⁻³) than in summer (3.2 ± 1.7 μg m⁻³) with less regional variability.

The 19-city average 24 h PM₁₅ mass concentration was 103.3 ± 82.3 μg m⁻³ (Table 3). Daily PM₁₅ varied by a factor of 78, ranging from 5.8 μg m⁻³ (Qingdao, summer) to 451.2 μg m⁻³ (Beijing, winter). Approximately ~ 81% and ~ 18% of PM₁₅ mass measurements exceeded China’s 24 h air quality standard of 75 μg m⁻³ during winter and summer, respectively. On average, carbonaceous aerosols constituted 36.4 ± 16.3% of PM₁₅.
Average northern POC and SOC concentrations were 22.2 ± 13.3 µg m⁻³ and 12.5 ± 15.2 µg m⁻³ in winter, accounting for ~ 52% and ~ 29% of TC, respectively (Fig. 2). Summer concentrations were lower (3.2–3.6 µg m⁻³), accounting for ~ 32% POC and ~ 36% SOC in TC. Winter is the most polluted season in northern China, with ~ 64% POC and ~ 36% SOC in OC; while summer is relatively clean, with ~ 47% POC and ~ 53% SOC.

In southern China, average wintertime POC concentrations were higher (19.6 ± 8.3 µg m⁻³, ~ 63% of TC) than those in summer (4.6 ± 1.5 µg m⁻³, ~ 40% of TC), with ~ 86% of OC as POC and ~ 14% as SOC. Minimal differences were found in summer SO2 between winter (3.3 ± 6.4 µg m⁻³, ~ 11% of TC) and summer (3.1 ± 3.2 µg m⁻³, ~ 27% of TC).

Table 2. Concentrations of OC and EC among 19 cities in 2013. *See site description in Table 1. bValues represent average ± standard deviation in µg m⁻³. cCM,carbonaceous matter = 1.6 × OC + EC. dNumbers of samples. Ave, average for Northern and Southern Cities. *The five sites that were not included in the 2003 study24.

Table 3. Average winter and summer concentrations for PM2.5 mass and carbonaceous aerosols in 2003 and 2013. (PM2.52003 − PM2.52013) × 100/PM2.52003. *Winter and summer average in Table 2 of the 2003 study24.

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Reduction in carbonaceous concentrations and implementation of control policies for 14 cities. For the 14 cities with 2003 results, Table 3 shows that average PM$_{2.5}$ mass decreased by 19% from 117.6 µg m$^{-3}$ in 2003 to 95.6 µg m$^{-3}$ in 2013. A similar 19% reduction was found for EC, from 6.8 to 5.6 µg m$^{-3}$; average OC decreased by 32% from 26.0 µg m$^{-3}$ to 17.5 µg m$^{-3}$. These reductions correspond to the emission control and energy policies of the 2006–2010 Five-Year Plan, despite increases of ~93% for coal consumption, ~77% for the crude oil usage, and ~470% for the number of motor vehicles.

Annual average concentrations were 11.7 ± 10.3 µg m$^{-3}$ for POC and 5.8 ± 10.3 µg m$^{-3}$ for SOC in 2013, yielding ~25% and ~49% reductions, respectively, since 2003 (Table 3). POC reductions were more pronounced in summer (~61% in the north and ~26% in the south) than the corresponding ~28% and ~22% in winter. Summer SOC decreased by ~44% and ~58% for northern and southern China, respectively. Wintertime SOC experienced a ~73% reduction in southern China, but a ~10% increase was observed for northern China from 2003 to 2013, possibly due to less stringent VOC and NOx controls for small boilers. Past studies found that VOCs and NOx emissions influence winter air quality in China.

Average 14-city carbonaceous aerosols constituted ~38% and 42% of PM$_{2.5}$ in 2003 and 2013, respectively. From 2013 to 2003, SOC and POC decreased by ~49% and 25%, respectively, whereas EC decreased by ~17%.

2013/2003 ratios for PM$_{2.5}$ mass, OC, and EC. Figure 3 illustrates nationwide pollution reduction, with most of the average 2013/2003 ratios less than unity for PM$_{2.5}$ mass, OC, and EC. Concentration reductions are most apparent in the three northern (i.e., Jinchang, Xi'an, Yulin) and five southern (i.e., Chongqing, Guangzhou, Hangzhou, Shanghai, and Wuhan) cities. However, increased wintertime OC concentrations were found in Beijing and Changchun, whereas increased EC levels were found during winter in Tianjin and during summer in Qingdao and Xiamen. Hong Kong showed increases in OC and EC from 2003 to 2013 for both seasons.

Changes of OC and EC concentrations between 2013 and 2003. Figure 1 shows that wintertime carbon decreased for most cities since 2003. The most apparent wintertime OC reductions (~49–62%) were found for Shanghai (~62%), Guangzhou (~53%), and Chongqing (~49%); EC also decreased (34–58%) with ~34% for Chongqing, ~40% for Shanghai, and ~58% for Guangzhou. Reduction of wintertime carbonaceous aerosols for the remaining cities ranged from 13 to 37% for OC and 2–49% for EC. Shanghai, Guangzhou, and Chongqing represent the three most developed regions in southern China: the Yangtze River Delta (YRD) region, the Pearl River Delta (PRD) region, and the Chong-Yu (CY) region, respectively. Control measures such as industrial plant closures and traffic controls during the 2010 (May to October) World Exposition in Shanghai and the 2010 (November) Asian Games in Guangzhou demonstrated the effectiveness of regional air quality management strategies that were implemented on larger scales.

Increases in wintertime OC (~46%) and EC (~18%) in Beijing and the OC increase (~26%) in Changchun may be associated with severe haze episodes under sluggish weather systems that occurred over central and eastern China in 2013. During the two January episodes (9th–15th and 25th–31st), maximum hourly PM$_{2.5}$ mass
concentrations in Beijing were 680 and 530 µg m⁻³, respectively. During January 2013, PM₁₂ in Beijing ranged 106–451 µg m⁻³, consisting for 35–55% of carbonaceous aerosol. Ji et al. also noted elevated carbon concentrations during winter of 2016/2017 in the Beijing–Tianjin–Hebei region, which were attributed to intense coal combustion for residential heating and a high frequency of unfavorable meteorological conditions.

Compared to 2003, summertime OC decreased by 53–77% in seven cities including: Xi’an (~77%), Jinchang (~73%), Chongqing (~71%), Yulin (~69%), Hangzhou (~59%), Tianjin (~57%), and Wuhan (~53%). Summertime EC reductions varied by city, with 64% in Chongqing and 20–40% in Beijing, Yulin, Xin’an, Shanghai, Wuhan, and Guangzhou. In addition to emission reductions, improved atmospheric dispersion in summer favored lower concentrations. Air quality in Beijing improved before and during the 2008 summer Olympics due to the temporary implementation of pollution control measures. However, air pollution levels surged after the Olympics, signifying the importance of implementing long-term control measures.

Increases in summertime OC and EC since 2003 were found in coastal cities such as Qingdao, Xiamen, and Hong Kong, with low TC concentrations (7.8–19.4 µg m⁻³). Coastal cities usually experience lower air pollution levels because of clean marine air intrusions. However, increases of primary emissions in Hong Kong may have resulted in the increase of OC and EC.

Spatial distribution of carbonaceous aerosols in 2013 and its association with economic development. Winter carbon concentrations in both monitoring years were high in inland cities and low in coastal and background cities. Lower wintertime OC concentrations were found in the coastal cities of Shanghai (10.9 ± 4.6 µg m⁻³), Xiamen (12.3 ± 3.2 µg m⁻³), and Hong Kong (15.3 ± 5.1 µg m⁻³) and the non-urban desert city of Jinchang (15.9 ± 5.8 µg m⁻³). The eight highest inland OC cities were Xi’an (64.1 ± 25.5 µg m⁻³), Changchun (49.5 ± 20.4 µg m⁻³), Taiyuan (45.3 ± 18.7 µg m⁻³), Beijing (39.8 ± 21.1 µg m⁻³), Chongqing (38.9 ± 8.9 µg m⁻³), Xining (36.2 ± 14.5 µg m⁻³), Chengdu (36.0 ± 9.0 µg m⁻³), and Tianjin (30.0 ± 15.6 µg m⁻³), with concentrations exceeding the 19 cities winter average OC of 29.1 ± 18.5 µg m⁻³.

With the exception of the Chongqing and Chengdu (capital of Sichuan Province) megacities, elevated carbon concentrations were found in northern China, reflecting the impacts from industries and central heating systems during cold winters with poor dispersion. After its establishment as a Municipality in 1997, Chongqing’s population increased to >30 million by 2014. Both cities are located in the Sichuan basin surrounded by mountains and have warmer climates with less winter heating. They suffer from severe acid rain from coal combustion, with carbonaceous aerosols accounting for 34–43% of PM₁₂ mass during winter.

Most of the cities with elevated wintertime OC concentrations are associated with decades-long economic development. For example, Changchun, the capital of Jilin Province, is the largest industrial and transportation hub in northeast China, producing ~9% of the Chinese automobiles in 2009, and it experienced a ~26% increase in OC from 2003 to 2013; whereas Taiyuan, the capital of Shanxi Province, produces ~25% of China’s coal (designated as the Taiyuan Coal Transaction Center since 2012), and reported average winter TC levels of 64.0 ± 24.5 µg m⁻³.

The highest summer OC (10.6 ± 2.2 µg m⁻³) was found in Beijing, but it is less than the lowest winter OC concentrations for the 19 cities. The minimum summer OC average occurred in Jinchang (2.2 ± 1.5 µg m⁻³), a desert city in Gansu Province (bordering Inner Mongolia to the north with a population of 0.2 million) with few local industries.

There were eight inland cities that exceeded the 19-city EC averages of 8.5 ± 5.0 µg m⁻³ during winter, ranging 9.1 to 18.6 µg m⁻³, including: Hangzhou (9.1 ± 3.5 µg m⁻³), Tianjin (9.1 ± 4.0 µg m⁻³), Hong Kong (9.6 ± 2.5 µg m⁻³), Chongqing (11.0 ± 2.8 µg m⁻³), Xi’an (11.2 ± 4.5 µg m⁻³), Changchun (11.9 ± 4.6 µg m⁻³), Chengdu (14.8 ± 3.4 µg m⁻³), and Taiyuan (18.6 ± 6.2 µg m⁻³). With the exception of Hangzhou and Hong Kong, these are industrial cities with abundant coal combustion.

Hong Kong reported an increase of ~29% in OC and ~122% in EC since 2003. Vehicle engine exhaust, especially from diesel vehicles, is an important contributor, accounting for 20–51% of fine particles.\(^{39–42}\)
Variability of OC/EC ratios. Figure 4 shows variations in OC/EC ratios as a function of EC concentrations, ranging 1.2–11.6 during winter and 0.01–6.9 during summer of 2013. Similar variability was found for 2003 (red circles in Fig. 4) with lower OC/EC ratios in winter for concentrations < 20 µg m\(^{-3}\). Changes in PM carbon properties may influence the relative amounts of particle light scattering and absorption\(^{24}\) related to visibility impairment and climate change.

Higher winter OC/EC ratios (6.0 to 11.6) during 2013 (barely seen in 2003) were found in the northwestern cities of Urumqi, Yulin, Xining, and Xi’an. This may be due to changes in the source mixtures. Abundances of the thermal carbon fractions also changed. In wintertime Xi’an, high temperature carbon fractions of OC3 and OC4 (480 °C and 580 °C) were most abundant in 2013, whereas lower temperature OC2 (280 °C) and OC3 fractions dominated in 2003. These changes may reflect changes in the composition of organic compounds in the combustion emissions.

Ten northern cities (with the exception of Taiyuan) reported higher wintertime OC/EC ratios than southern cities. Taiyuan reported the lowest winter OC/EC ratio of 2.4, close to the ratio of 2.0 for primary fossil fuel combustion from the national emission inventory\(^{43}\). This is reasonable considering that Taiyuan is China’s coal capital, with coal energy, metallurgy, chemicals, and machinery industries\(^{44}\).

While the distribution pattern of OC/EC ratios during summer in 2013 was similar to that of 2003, the average ratio is 23% lower. This is attributed to the decrease in OC from 13.8 µg m\(^{-3}\) in 2003 to 7.1 µg m\(^{-3}\) in 2013 as the average EC concentrations (3.4–3.6 µg m\(^{-3}\)) remained similar. The summer OC/EC ratios were the highest in Beijing (3.0–4.8), followed by Guangzhou (1.9–6.9), with the lowest in Hong Kong (0.6–0.8). There are different vehicle fuel formulations among the cities. The higher proportion (~ 38%) of diesel vehicle emissions in Hong Kong resulted in lower OC/EC ratios. This is consistent with the average OC/EC ratio of 2.7 for gasoline and LPG vehicles and 0.5 for diesel vehicles\(^{45}\).

Conclusions

Decreasing OC and EC concentrations were apparent at a national scale from 2003 to 2013; but the extent of reduction varied by season and location. Most inland cities showed decreases in carbon concentrations with the exception of Beijing and Changchun (capital of Jilin in northern China). Shanghai showed an apparent decrease, with a less distinct reduction in Qingdao and Xiamen for OC and slight increases for EC. The exception was found in Hong Kong, with a 2003 to 2013 increase of ~29% for OC and ~122% for EC.

Carbonaceous aerosols constituted ~37.5% and ~41.5% of PM\(_{2.5}\) mass for 2013 and 2003, respectively. Majority of the TC is in OC, with average OC/EC ratios of 3.1 and 4.0 for 2013 and 2003, respectively.

Summertime PM\(_{2.5}\) mass and OC concentrations were ~25–35% of those in winter with less variability in EC. Wintertime carbon concentrations were higher in northern than southern China, elevated in inland cities and low in coastal and non-urban desert cities. Winter in northern China is the most polluted season with ~64% primary organic carbon (POC) and ~36% of secondary organic carbon (SOC) in OC.

Wintertime OC/EC ratios were higher in most northern than southern cities, indicating the impact of haze episodes and fossil fuel combustion from heating. Elevated 2013 OC/EC ratios (6.0 to 11.6) in northwestern cities such as Urumqi, Yulin, Xining, and Xi’an, were barely seen in 2003. Changes in abundances among eight
thermal carbon fractions suggest changes in emission properties over the past decade. Average summer 2013 OC/EC ratio of 2.5 was much lower than corresponding 2003 ratio of 4.2, with less variability in OC/EC ratios (3.7–3.8) during winter.

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References
1. Charlson, R. J. et al. Climate forcing by anthropogenic aerosols. Science 255, 423–430 (1992).
2. Intergovernmental Panel on Climate Change (IPCC) Climate change 2007: The physical science basic. In Changes in Atmospheric Constituents and in Radiative Forcing (eds Forster, P. et al.) 129–234 (Cambridge University Press, Cambridge, 2007).
3. Ramana, M. V. et al. Warming influenced by the ratio of black carbon to sulphate and the black-carbon source. Nat. Geosci. 3, 542–545 (2010).
4. Shi, T. S. et al. Elemental and organic carbon exposure in highway tollbooths: A study of Taiwanese toll station workers. Sci. Total Environ. 402, 163–170 (2008).
5. Klotz, S. V. et al. Elemental carbon exposure at residence and survival after acute myocardial infarction. Epidemiology 20, 547–554 (2009).
6. Gentner, D. R. et al. Elucidating secondary organic aerosol from diesel and gasoline vehicles through detailed characterization of organic carbon emissions. P Natl. Acad. Sci. U.S.A. 109, 18338–18333 (2012).
7. Robinson, A. L. et al. Revisiting organic aerosol: Semivolatile emissions and photochemical aging. Science 315, 1259–1262 (2007).
8. Huang, Y., Dickinson, R. E. & Chameides, W. L. Impact of aerosol indirect effect on surface temperature over East Asia. PNAS 103(12), 4371–4376 (2006).
9. Haywood, J. & Boucher, O. Estimates of the direct and indirect radiative forcing due to tropospheric aerosols: A review. Rev. Geophys. 38, 513–543 (2000).
10. Hansen, J., Sato, M. & Ruedy, R. Radiative forcing and climate response. J. Geophys. Res. Atmos. 102, 6831 (1997).
11. Lanki, T. et al. Can we identify sources of fine particles responsible for exercise-induced ischemia on days with elevated air pollution? The ULTRA study. Environ. Health Perspect. 114, 655–660 (2006).
12. Lewene, M., Plato, N. & Gustavsson, P. Exposure to particles, elemental carbon and nitrogen dioxide in workers exposed to motor exhaust. Ann. Occup. Hyg. 51, 693–701 (2007).
13. Cao, J. J., Xu, H. M., Xu, Q., Chen, B. H. & Kan, H. D. Fine particulate matter constituents and cardiopulmonary mortality in a heavily polluted Chinese city. Environ. Health Persp. 3, 373–378 (2012).
14. Jin, Y., Henriek, A. & Zhang, S. Air pollution control policies in china: A retrospective and prospects. Int. J. Environ. Res. Public Health. 13, 1219 (2016).
15. Song, X. Research on total emission control policies in China. Ph.D. Thesis. Tsinghua University, Beijing, China (2015).
16. National Bureau of Statistics. China Statistical Yearbook (2003 and 2014 Editions) (China Statistics Press, Beijing, 2003–2014).
17. Tian, L. et al. Increasing trend of primary NO2 exhaust emission fraction in Hong Kong. Environ. Geochim. Health 33, 623–630 (2011).
18. Wild, R. et al. On-road measurements of vehicle NO2/NOx emission ratios in Denver, Colorado, USA. Atmos. Environ. 148, 182–189 (2017).
19. Zheng, C. H. et al. Quantitative assessment of industrial VOC emissions in China: Historical trend, spatial distribution, uncertainties, and projection. Atmos. Environ. 150, 116–125 (2017).
20. Ai, Z. T., Mak, C. M. & Lee, H. C. Roadside air quality and implications for control measures: A case study of Hong Kong. Atmos. Environ. 137, 6–16 (2016).
21. Watson, J. G., Chow, J. C. & Chen, L. W. A summary of organic and elemental carbon/black carbon analysis methods and intercomparisons. Aerosol Air Qual. Res. 5, 65–102 (2005).
22. Chow, J. C., Watson, J. G., Crow, D., Lowenthal, D. H. & Merrill, T. M. Comparison of IMPROVE and NIOSH carbon measurements. Aerosol Sci. Technol. 34, 23–34. https://doi.org/10.1080/07367480600623711 (2001).
23. Ji, D. et al. The carbonaceous aerosol levels still remain a challenge in the Beijing–Tianjin–Hebei region of China: Insights from continuous high temporal resolution measurements in multiple cities. Environ. Int. 126, 171–183 (2019).
24. Cao, J. J. et al. Spatial and seasonal distributions of carbonaceous aerosols over China. J. Geophys. Res. Atmos. 112, 11–22 (2007).
25. Watson, J. G., Tropp, R. J., Kohl, S. D., Wang, X. L. & Chow, J. C. Filler processing and gravimetric analysis for suspended particulate matter samples. Aerosol Sci. Eng. 1, 193–205 (2017).
26. Chow, J. C. & Watson, J. G. Guideline on Speciated Particulate Monitoring. Report 32–34 (United States Environmental Protection Agency, Research Triangle Park, 1998).
27. Chow, J. C. et al. The IMPROVE A temperature protocol for thermal/optical carbon analysis: Maintaining consistency with a long-term database. J. Air Waste Manage. Assoc. 57(9), 1014–1023 (2007).
28. Chow, J. C. et al. The DRI thermal reflectance carbon analysis system: Description, evaluation and applications in U.S. Air quality studies. Atmos. Environ. A 27(8), 1185–1201 (1993).
29. Chow, J. C. et al. Equivalence of elemental carbon by thermal/optical reflectance and transmittance with different temperature protocols. Environ. Sci. Technol. 38(16), 4414–4422 (2004).
30. Seinfeld, J. H. & Pandis, S. N. Atmospheric Chemistry and Physics: From Air Pollution to Climate Change (Wiley, Hoboken, 2006).
31. Turpin, B. J. & Huntzicker, J. J. Identification of secondary organic aerosol episodes and quantification of primary and secondary organic aerosol concentrations during SCAQS. Atmos. Environ. 29, 3527–3544 (1995).
32. Huang, R. J. et al. High secondary aerosol contribution to particulate pollution during haze events in China. Nature 514, 218–222 (2014).
33. Wu, C. & Yu, J. Z. Determination of primary combustion source carbon organic to elemental carbon (OC/EC) ratio using ambient OC and EC measurements: Secondary OC-EC correlation minimization method. Atmos. Chem. Phys. 16, 5453–5465. https://doi.org/10.5194/acp-16-5453-2016 (2016).
34. Wu, C. & Yu, J. Z. Evaluation of linear regression techniques for atmospheric applications: The importance of appropriate weight. Atmos. Meas. Tech. 11, 1233–1250. https://doi.org/10.5194/amt-11-1233-2018 (2018).
35. Turpin, B. J. & Huntzicker, J. J. Secondary formation of organic aerosol in the Los Angeles basin: A descriptive analysis of organic and elemental carbon concentrations. Atmos. Environ. A Gen. Top. 25(2), 207–215 (1991).
36. Park, S. S. et al. Evaluation of the TM0 and TOT methods for OC and EC measurements and their characteristics in PM2.5 at an urban site of Korea during ACE-Asia. Atmos. Environ. 39(28), 5101–5112 (2005).
37. Chow, J. C., Lowenthal, D. H., Chen, L.-W., Wang, X. L. & Watson, J. G. Mass reconstruction methods for PM2.5: A review. Air Qual. Atmos. Health 8, 243–263. https://doi.org/10.1007/s11869-015-0338-3#page-1 (2015).
38. Yuan, B. et al. VOC emissions, evolutions and contributions to SOA formation at a receptor site in eastern China. Atmos. Chem. Phys. 13, 8815–8832 (2013).
39. Wang, Y. S. et al. Mechanism for the formation of the January 2013 heavy haze pollution episode over central and eastern China. Sci. China Earth Sci. 57, 14–25 (2014).
40. Cheng, Y. et al. Characteristics and source apportionment of PM$_1$ emissions at a roadside station. J. Hazard. Mater. 195, 82–91 (2011).
41. Louie, P. K. K. et al. Seasonal characteristics and regional transport of PM$_{2.5}$ in Hong Kong. Atmos. Environ. 39(9), 1695–1710 (2005).
42. Wu, X. et al. Characterization and source apportionment of carbonaceous PM$_{1.5}$ particles in China—A review. Atmos. Environ. 189, 187–212 (2018).
43. Cao, G. L., Zhang, X. & Zheng, F. Inventory of black carbon and organic carbon emissions from China. Atmos. Environ. 40, 6516–6527 (2006).
44. Zhang, F., Zhao, J., Chen, J. & Xu, L. Pollution characteristics of organic and elemental carbon in PM$_{2.5}$ in Xiamen. China. J. Environ. Sci. China 23(8), 1342–1349 (2011).
45. Cheng, Y. et al. Chemically-speciated on-road PM$_{2.5}$ motor vehicle emission factors in Hong Kong. Sci. Total Environ. 408, 1621–1627 (2010).

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Y.C. and J.J.C. prepared the main manuscript; J.M.Z. and S.X.L. provided experimental data; J.C.C. and J.G.W. participated in data interpretation and English editing. All co-authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

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