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The measurement of aerosol optical properties at a rural site in Northern China

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Abstract. Atmospheric aerosols constitute one of the largest sources of uncertainty in the estimation of radiative forcing for climate. From April 2003 to January 2005, in situ measurements of aerosol optical properties were conducted at a rural site in Northern China, Shangdianzi Global Atmosphere Watch (GAW) regional station (SDZ), about 150 km from Beijing. Mean values (standard deviation, S.D.) of scattering and absorption coefficients for the entire period are $174.6 \,\mathrm{Mm^{-1}} \ (189.1 \,\mathrm{Mm^{-1}}) \ \text{and} \ 17.5 \,\mathrm{Mm^{-1}} \ (13.4 \,\mathrm{Mm^{-1}}),$ respectively. These values are approximately one third of the reported values for scattering coefficients and one fifth of those for absorption coefficients obtained in the Beijing urban area. The mean single scattering albedo (SSA) for the entire period was estimated as 0.88 (0.05), which is about 0.07 higher than the values reported for the Beijing urban area, and also higher than the values (0.85) used in a reported climate simulation for China and India. Both the absorption and scattering coefficients showed a seasonal cycle with the lowest values in winter, while the highest values occurred in summer for absorption coefficients and in fall for scattering coefficients. The mean SSA values were lowest in spring and highest in winter. The daily variations of aerosol absorption and scattering coefficients were strongly influenced by synoptic changes throughout the observation period. A trajectory cluster analysis was applied to discern the source characteristics of aerosol optical properties for different air masses. The cluster-mean aerosol scattering coefficients, absorption coefficients and SSA were all high when the air masses moved from SW and SE-E directions to the site and aerosols were influenced with heavy pollution



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from the dense population centers and industrial areas. The cluster-mean SSA for air masses coming from the polluted areas was not only higher than those with trajectories from the "clean" directions, but also higher than the reported values for the regions with high pollution emissions (such as the Beijing urban area). This fact might reflect the substantial secondary aerosol production during transport. The characteristics of aerosol optical properties measured at this rural site suggest significant impacts of human activities on the regional aerosol.

1 Introduction

Aerosols perturb the radiation balance of the Earth directly through scattering and absorbing solar radiation, and indirectly by acting as condensation nuclei in cloud formation, thus affecting the optical properties and lifetimes of clouds (Rosenfeld, 1999, 2000; Twomey, 1977). The optical properties of aerosols are related to both their chemical composition and particle size distributions. Atmospheric aerosols in different regions consist of varying composition and size modes, which lead to different radiative impacts on regional climate. Black carbon (BC) aerosol is particularly important due to its characteristics of strong light absorption, because the absorbing aerosols heat the air, which can alter regional atmospheric stability and vertical motions and affect the large scale circulation and precipitation with significant regional climate effects (Jacobson, 2002; Menon et al., 2002; Ramanathan et al., 2001). As opposed to the well mixed greenhouse gases, aerosol has a highly inhomogeneous geographic distribution. The effect of aerosols is so far very poorly taken into account in climate models (Anderson et

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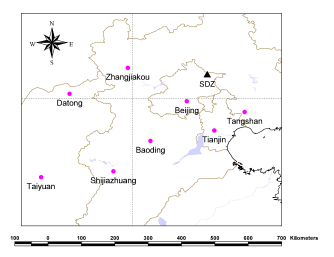


Fig. 1. Map of SDZ station and the major cities in the region.

al., 2003). The spatial and temporal variations of radiative forcing by aerosols are strongly influenced by variability of aerosol concentration distribution and optical properties on local to regional scales. For this reason, the characterization of aerosol properties in various regions around the globe is essential to understand the role of aerosols on climate.

In China, as a result of the rapid population and economic growth, emissions of anthropogenic pollutants have increased dramatically in recent years. Emissions of SO2 and NO_x in China are similar in magnitude with those of Europe and eastern North America (Akimoto and Narita, 1994), but aerosol sources in China have their unique features. The mixture of heavy air pollution from urbanization and industrial activities and desert dust particles from the frequent wind storms result in a rather complex nature of aerosol optical properties in China (Holler et al., 2003). The high rate of coal and biofuels usage and lower combustion efficiency has made China a significant source of BC. It was estimated that BC emitted in China accounted for roughly one-fourth of global anthropogenic emissions, although a significant uncertainty exists in estimations of the submicron BC emission inventory (Cooke et al., 1999; Street et al., 2001). Uncertainties in the measurement of optical properties of aerosols in China and other Asian countries present one of the largest sources of uncertainty in the estimation of aerosol radiative forcing (Holler et al., 2003). In the past 20 years, many studies have focused on the mass concentration and physical and chemical properties of aerosols in China, but few measurements were conducted focusing on aerosol radiative properties (Xu et al., 2002, 2004).

In this paper, a 16–21 month study of in situ measurements of aerosol properties at a rural site in northern China is presented, and the characteristics and variation of aerosol optical properties are reported. Filter analysis was performed to provide insight on the chemical composition of aerosols,

and the influence of air mass transport on the optical properties of aerosol was determined through trajectory clustering analysis.

2 Experiment

2.1 Site descriptions

Shangdianzi station (SDZ, 117°07′E, 40°39′N, 293.3 m a.s.l.) is located in Miyun County, a suburb of the greater Beijing metropolis. The site is on the gentle slope of a small hill. The geography surrounding the station is characterized by rolling hills with farmland, orchards and forests. On the foot of the hills about 2 km south of the station is Shangdianzi village with about 1200 inhabitants. The major local economical activities within Miyun County are farming and fruit growing. Since Miyun reservoir is the major water supply to the Beijing metropolitan area, Miyun County has been designated as a "Preservation Area", which means that there are only minimal natural and anthropogenic pollution sources within a 30 km range surrounding the SDZ site.

Figure 1 shows the map of the SDZ station and some cities and regions with dense population and high industrial activities. For example, Beijing and Tianjin are the two most populated cities in Northern China; Shijiazhuang and Taiyuan are the provincial capital cities of Hebei province and Shanxi province, respectively; Datong (in Shanxi province) is one of the largest coal production regions in China; and Baoding, Zhangjiakou and Tangshan in Hebei province are all large industrial cities in northern China. They are all regions with high pollution emissions (Cao et al., 2006; Streets and Carmichael, 2003). It can be seen from the map that the major cities and industrial areas with high pollution emissions in this region are located predominantly in the sectors from the west to the southeast of the site. Areas in the northern sector are much less inhabited, comprised of the vast grassland of Inner Mongolia (or Nei Mongol) and mountainous rural regions of Hebei province, where the population is relatively sparse and industrial activities less prevalent.

2.2 Measurements and instrumentation

During the period from April 2003 to January 2005, an intensive campaign of aerosol measurement was conducted at the SDZ GAW station in Northern China. The experiment included the measurement of aerosol scattering and absorption coefficients, size-resolved aerosol mass, chemical compositions, and meteorological observations.

Table 1 lists the measured parameters and instruments used in the experiment. Aerosol scattering coefficient (σ_{sca}) was measured with an integrating Nephelometer (Model M9003, EcoTech, Australia). This instrument used LEDs as the light source at a wavelength of 525 nm. The scattering integration angle is from 10° to 170°. The truncation error correction was not applied to the data in the study since there

Table 1. The measurements and instrumentation.

Measurements	Time period	Instrumentation
Scattering coefficient $\sigma_{ m sca}$	10 September 2003 to 15 January 2005	Integrating Nephelometer (Model M9003, ECOTech, Australia), with wavelength of 525 nm
Absorption coefficient σ_{abs}	15 April 2003 to 15 January 2005	Aethalometer (Model AE31, Magee Scientific, USA), with seven wavelengths at 370, 470,
Filter sampling for aerosol composition (Ions and OC, EC)	Summer, 17 July to 2 August 2004	520, 590, 660, 880, 950 nm Andersen KA200 multi-stage impactor, with Quartz and Teflon sampling membranes

was no dust storm during the entire observation period and the truncation errors are small for submicrometer aerosols. A theoretical estimation based on Mie-theory indicated that the truncation error was less than 10% for fine particles. A background (zero) check was done automatically by pumping in particle-free air once each day and a weekly span check was performed manually by the operator using particle-free HFC-R134a gas recommended by the manufacturer. The results indicated that the bias was less than 2 Mm⁻¹ for zero checks, and less than 10% for span checks. The relative humidity (RH) in the cell of the instrument was controlled below 60% by an automatic heating inlet provided by the manufacturer. This heating inlet could cause evaporation of volatile inorganic species (such as nitrate) and volatile organic matters. Bergin et al. (1997) studied the decrease of the scattering coefficient caused by the evaporative loss of aerosols in a nephelometer, and it was generally less than 20% for pure nitrate aerosol. Therefore, the reduction of the measured aerosol scattering coefficient due to the loss of nitrate by the heating of our inlet was estimated to be small because nitrate represents only a small fraction of the total mass in fine particles in northern China (Zhang et al., 2003). A precise estimation on the loss of organic matters due to the inlet heating was difficult, however, because of the lack of information on organic species in our measurements.

The BC aerosol concentration was measured with an Aethalometer (Model AE31, Magee Scientific, USA). The AE31 Aethalometer measured the optical attenuation of light from LED lamps with seven different wavelengths (370, 470, 520, 590, 660, 880, and 950 nm) transmitted through the aerosols deposited continuously on a quartz fiber filter (Hansen et al., 1984). The difference in light transmission through the particle-laden sample spot and a particle free reference spot of the filter is attributed to the absorption caused by aerosol. The attenuation of light is converted to the BC mass concentration using wavelength dependent calibration factors as recommended by the manufacturer (Aethalometer Manual, Magee Scientific). As discussed below, the Aethalometer light attenuation measurement was used to calculate the aerosol light absorption coefficient (σ_{abs}).

The inlets of the instruments were located on the roof of the monitoring room, at about 6 m above the ground and 3 meters above the roof. The sample air passed through 0.95 cm i.d. black conductive tubing into the room without any aerosol size cut-off.

All of the scattering and absorption coefficient data and relevant ancillary parameters were recorded at 5-min intervals. The datasets have been manually edited to remove invalid data resulting from instrumental or sampling problems.

The size-resolved aerosol filter sampling was taken only for some specific time periods during the experiment. The analysis included mass concentrations and some of the major chemical species, such as ions, organic carbon (OC) and elemental carbon (EC) of aerosols. The procedure for mass gravimetric and chemical analysis was identical to those described in previous work (Bergin et al., 2001; Yan et al., 2006).

2.3 Method for aerosol absorption coefficient (σ_{abs}) calculation

Aerosol light absorption coefficient (σ_{abs}) can be directly calculated from the attenuation measured by the Aethalometer or indirectly calculated based on the BC concentrations recorded by the instrument. The direct calculation of aerosol light absorption at any wavelength uses the following formula (Bodhaine, 1995; Weingartner et al., 2003):

$$\sigma_{\rm abs} = \frac{A}{Q} \times \frac{\Delta ATN}{\Delta t} \times \frac{1}{CR} \tag{1}$$

where, A is the filter spot area, Q the volumetric flow rate and $\triangle ATN$ is the change in attenuation during the time interval $\triangle t$, and C is a wavelength-independent empirical correction factor, which corrects for the enhancement of the optical path in the filter due to multiple scattering of the light beam at the filter fiber, R is the correction factor describing the changes in instrumental response with increased particle loading on the filter (shadowing effect). The C factor was found to vary with different aerosol types and mixing state, and R is equated to unity for aged particles (Weingartner et al., 2003).

The indirect way to obtain the light absorption coefficient from the recorded BC concentration is based on the following equation:

$$\sigma_{\text{abs}} = \alpha \times [BC] \tag{2}$$

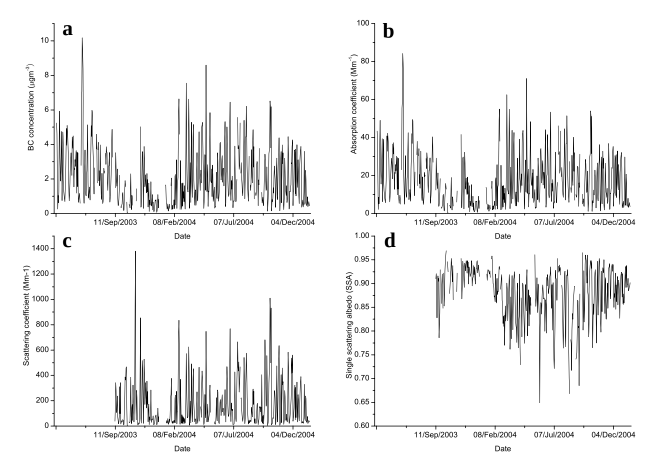


Fig. 2. The time series of daily averaged BC mass concentration, aerosol absorption and scattering coefficient, and single scattering albedo at SDZ (a) BC concentration; (b) Absorption coefficient; (c) Scattering coefficient; (d) Single scattering albedo (SSA).

where, α is the conversion factor or the BC absorption efficiency, which can be determined theoretically from Mie-theory or empirically from linear regression of the Aethalometer BC concentration data against the aerosol absorption coefficient measured from a reference method (Arnott et al., 2003, 2005; Clarke and Charlson, 1985).

In this work, the indirect method (based on the recorded BC concentration by the Aethalometer) was employed to calculate the aerosol light absorption coefficient. The conversion factor α was obtained based on the result of the inter-comparison experiment conducted in southern China (O. Schmid, D. Chand, and M. O. Andreae, Aerosol Optical Properties in Urban Guangzhou; PRD workshop, Beijing, 13-14 January 2005). In that experiment, the conversion factor $\alpha = 8.28 \,\mathrm{m}^2/\mathrm{g}$ (with correlation coefficient $R^2 = 0.92$) was derived by a linear regression of the BC concentrations at 880 nm (in the unit of $\mu g/m^3$) from the Aethalometer against the light absorption coefficients at 532 nm (in units of Mm⁻¹) simultaneously measured with a photoaccoustic spectrometer (PAS). This conversion factor is comparable to $\sim 8-10 \,\mathrm{m}^2/\mathrm{g}$ (at 550 nm) obtained in Mexico city (Barnard et al., 2005), but a little lower than the historical value of $10 \,\mathrm{m}^2/\mathrm{g}$ often used for urban aerosols (Moosmuller et al., 1998), and higher than the value of $7 \text{ m}^2/\text{g}$ for diesel soot suggested by Fuller (Fuller et al., 1999). It was very close, however, to the value of $8.0 \text{ m}^2/\text{g}$ obtained in Beijing urban area by Bergin (Bergin et al., 2001) and that of $8.5 \text{ m}^2/\text{g}$ reported by Arnott from BRAVO (Big Bend Regional Aerosol and Visibility Observation Study) (Arnott et al., 2003).

3 Results and discussion

3.1 Aerosol optical properties and BC mass concentrations

Figure 2 and Table 2 show the time series and the statistics of daily-average BC mass concentration and aerosol optical properties measured during the experiment. The mean (standard deviation) concentration of BC, aerosol absorption and scattering coefficient for the entire period were $2.12\,\mu\text{g/m}^3$ ($1.62\,\mu\text{g/m}^3$), $17.54\,\text{Mm}^{-1}$ ($13.44\,\text{Mm}^{-1}$), and $174.6\,\text{Mm}^{-1}$ ($189.1\,\text{Mm}^{-1}$), and the median values were $1.78\,\mu\text{g/m}^3$, $14.71\,\text{Mm}^{-1}$, and $101.6\,\text{Mm}^{-1}$, respectively. It should be noted that the large standard deviation for each variable

reflected the strong fluctuation of the measured aerosol properties

On average, the aerosol absorption and scattering coefficients at SDZ were about one-fifth and one-third lower than those measured in the Beijing urban area in summer 1999 (Bergin et al., 2001). Table 3 lists a few other published results for the purpose of comparison. The mean aerosol absorption coefficient at SDZ was comparable to that obtained at Lin'an regional background station in a rural area of the Yangtze Delta Region in eastern China (Xu et al., 2002) and at the Cape D'Aguilar station of Hong Kong in eastern and southeastern China (Man and Shih, 2001), but higher than that measured at Yulin in the Gobi Desert (Xu et al., 2004). The aerosol scattering coefficient at SDZ was only about half of that at Lin'an station, but much higher than that of Cape D'Aguilar (Man and Shih, 2001; Xu et al., 2002). It is similar to that at Yulin in the Gobi Desert (Xu et al., 2004).

The single scattering albedo (SSA) is defined as the ratio of the aerosol scattering coefficient to the extinction coefficient (sum of the absorption and scattering coefficient). This parameter is especially important in the estimation of direct aerosol radiative forcing, since even a small error in its estimation might change the sign of aerosol radiative forcing (Takemura et al., 2002). To calculate SSA, a wavelength correction for aerosol absorption was applied to the measured absorption coefficients using an empirical approach for BC aerosol, $\sigma_{abs} \infty \lambda^{\beta}$, with an absorption exponent $\beta = -1$ (wavelength λ in the interval 0.45 μ m–0.70 μ m) (Bodhaine, 1995). The results showed that the mean and the median value of SSA (at 525 nm) obtained at SDZ were 0.88 and 0.90, respectively (see Table 2). This mean SSA was within the range of 0.85-0.95 retrieved through AERONET for the northern hemisphere (Dubovik et al., 2002), but lower than the value of 0.93 obtained at Lin'an in November, 1999 (Xu et al., 2002) and 0.95 at Yulin (Xu et al., 2004). It was about 0.07 higher than that (0.81) reported for the Beijing urban site (Bergin et al., 2001), and also a little higher than the value (0.85) used in climate modeling simulations for China and India (Menon et al., 2002).

Compared to the results reported for North America (see Table 3), mean aerosol absorption coefficient at SDZ is about three to four times greater than the mean value measured at Bondville, Illinois (BND), a rural location in the east central region of the United States, and forty five times greater than that measured at Barrow, Alaska (BRW), a clean region in North America. The mean aerosol scattering coefficient is three times greater than that at BND and eighteen times greater than that measured at BRW, and the mean SSA is about 0.04 and 0.08 lower than the value (at 550 nm) of BND and BRW, respectively (Delene and Ogren, 2002).

3.2 Seasonal variations of aerosol optical properties

In this section, we use the data measured from December 2003 to November 2004 to discuss the seasonal variations

of aerosol optical properties and the influencing factors at the SDZ site. The calculated seasonal average (based on daily averages) of aerosol absorption, scattering coefficient and SSA are presented in Table 4, where the winter time is the period from December 2003 to February 2004, the spring from March to May, the summer from June to July, and the fall from September to November of 2004. It was found that the aerosol absorption coefficient was relatively higher in summer and spring and lower in fall and winter with the highest value in summer and the lowest in winter. The aerosol scattering coefficient showed a slightly different seasonal distribution. The aerosol scattering coefficients were elevated in fall and summer with the highest levels observed in fall, and lower in winter and spring with the lowest levels in winter. Calculated SSA values were higher in winter and fall, but relatively low in spring and summer. Sheridan et al. (2001) reported similar seasonal variations of aerosol absorption coefficient at the Southern Great Plains Cloud and Radiation Testbed site (SGP CART site), where the light absorption coefficients were higher in the late spring (April) through fall, and also generally lower SSA values were found over the same time period. However, the seasonal variations of aerosol properties observed during the selected period at SDZ were obviously different from those measured at Cape D'Aguilar in Hong Kong (see Table 3), where the highest values of absorption and scattering coefficients were found in winter and fall, and the lowest in summer (Man and Shih, 2001). Man and Shih (2001) attributed such seasonal variations mainly to the large-scale changes in transport of pollutants, the northerly wind in winter brought polluted continental air masses while the southern flows in summer contained clean oceanic air masses from the tropics. The reasons for the seasonal variations at SDZ site are complicated, and include the seasonal changes in source emissions, atmospheric transport, and chemical transformations. The prevailing wind direction shifting over different seasons is probably one of the important factors, as indicated from the wind rose plot of the site (Fig. 3).

In winter (from December 2003 to February 2004), the prevailing winds were from ENE and NE directions, where there were no strong anthropogenic or combustion aerosol sources, and cleaner air masses resulted in the lowest levels of measured aerosols at the site. In spring (from March to May, 2004), the dominant wind directions at the SDZ site were from WSW and SW, and as a consequence, the advection of the air masses from highly populated and polluted regions caused the increase of aerosol absorption and scattering coefficients at the site compared to those in winter. In summer, although the frequency of winds from NE and ENE direction increased, an equal fraction of the winds still came from the WSW and SW. Crop residue burning frequently occurred after the harvest in this region, and the combination of transport and increased biomass combustion emissions caused the highest absorption coefficient in summer at the site. In fall, the frequency of the WSW wind

Table 2. The statistical results of 24 h average BC mass concentration, absorption and scattering coefficient, and single scattering albedo (SSA) of aerosol during the measurements at SDZ from April 2003 to January 2005 for BC (or aerosol absorption coefficient) and from September 2003 to January 2005 for aerosol scattering coefficient.

	BC Concentration (µg/m ³)	$\sigma_{\rm abs} {\rm Mm}^{-1})$	$\sigma_{\rm sca}{\rm Mm}^{-1}$	SSA (at 525 nm)
Mean	2.12	17.54	174.6	0.88
S.D.	1.62	13.44	189.1	0.05
Maximum	10.17	84.23	1381.8	0.97
Minimum	0.035	0.29	4.3	0.65
Median	1.78	14.71	101.6	0.90
Number of days	575	575	461	426

Table 3. Aerosol optical properties at some different sites.

Site	Period	$\sigma_{abs}(\mathrm{Mm}^{-1})$	$\sigma_{sca}(\mathrm{Mm}^{-1})$	SSA	Instrumentation	References
SDZ, China	2003.9-2005.1	17.54	174.6	0.88	AE31, Magee Scientific, M9003	This work
(rural)					Nephelmeter, EcoTech	
Lin'an, China	1999.11	23	353	0.93	PSAP, Radiance Research, M903	(Xu et al., 2002)
(rural)					Nephelometer, Radiance Research	
Yulin, China	2001.4	6	158	0.95	PSAP, Radiance Research, M903	(Xu et al., 2004)
(Gobi desert)					Nephelometer, Radiance Research	
Cape D'Aguilar,	1997.1-1998.2	25.72 15.79 6.03	64.77 38.65 8.71		PSAP, Radiance Research, M903	(Man and Shih, 2001)
Hongkong					Nephelometer, Radiance Research	
	1998.3-1998.4	18.98 31.22	70.91 96.75			
	1998.11-1999.2					
	1998.5-1998.8					
	1998.9-1998.10					
Beijing, China (urban)	1999.6	83	488	0.81	PSAP, Radiance Research, M903	(Bergin et al., 2001)
					Nephelometer, Radiance Research	
Beijing, China (urban)	2003.6-12			0.793	AE14, Magee Scientific	(Mao and Li, 2005)
Bondville, Illinois	1996.9.19-2000.9.26	4.66 ± 2.27 ,	57.7 ± 17.7	0.924 ± 0.0028 ,	PSAP, Radiance Research	(Delene and Ogren, 2002)
Great Plains, US		$<10\mu\mathrm{m}$	$<10 \mu\mathrm{m}$	$<10\mu\mathrm{m}$	3563 Nephelometer, TSI, Inc.	
Barrow, Alaska, US	1997.10.6-2000.9.26	0.39 ± 0.41 ,	$9.76\pm5.20,$	0.99 ± 0.48 ,	PSAP, Radiance Research	
		$<10\mu\mathrm{m}$	$<10 \mu\mathrm{m}$	$<10\mu\mathrm{m}$	3563 Nephelometer, TSI, Inc.	
Southern Great Plain,	1997.4.6-2000.9.26	2.46 ± 1.09 ,	46.9 ± 16.9 ,	0.947 ± 0.025 ,	PSAP, Radiance Research	
Oklahoma, US		$<10\mu\mathrm{m}$	$<$ 10 μ m	$<$ 10 μ m	3563 Nephelometer, TSI, Inc.	

Table 4. The seasonal averages of aerosol absorption and scattering coefficient and SSA (from December 2003 to November 2004).

	$\sigma_{\rm abs} {\rm Mm}^{-1}$	$\sigma_{\rm sca}{\rm Mm}^{-1}$	SSA (at 525 nm)
Winter	8.89 (10.10)	115.76 (155.27)	0.91 (0.03)
Spring	18.27 (14.02)	154.09 (160.10)	0.85 (0.05)
Summer	21.21 (12.22)	190.30 (167.52)	0.86(0.06)
Fall	16.73 (11.62)	215.07 (222.91)	0.88 (0.06)

direction decreased and the ENE wind increased, and as a result mean aerosol absorption coefficients were lower in fall than in summer and spring but higher than in winter.

However, in contrast to the aerosol absorption coefficient, the higher levels of aerosol scattering coefficient in the fall could not be satisfactorily explained only by shifts in the wind direction. Another factor that can enhance aerosol scattering coefficients is the chemical production of secondary aerosols, either through photochemical reactions or in fogs (visual range < 1 km) and mists (1 km < visual range <10 km). The more frequent occurrence of fogs and mists in the fall period was documented through meteorological observations indicating more foggy (fog and/or mist) days and poorer visibility (visual range <10 km) at the site. Table 5 shows the number of days with fog/mist and poorer visual range in each month. There were 30 foggy days in the fall (September, October and November) of 2003, and 29 days in the fall of 2004, with 3 fog days in 2003 and 3 fog days in 2004. Under the fog/mist weather conditions the growth and/or heterogeneous production of aerosol was favored and could significantly enhance the observed aerosol scattering coefficients. This can be illustrated in the daily average dataset with the highest daily aerosol scattering coefficients often found on fog days in fall.

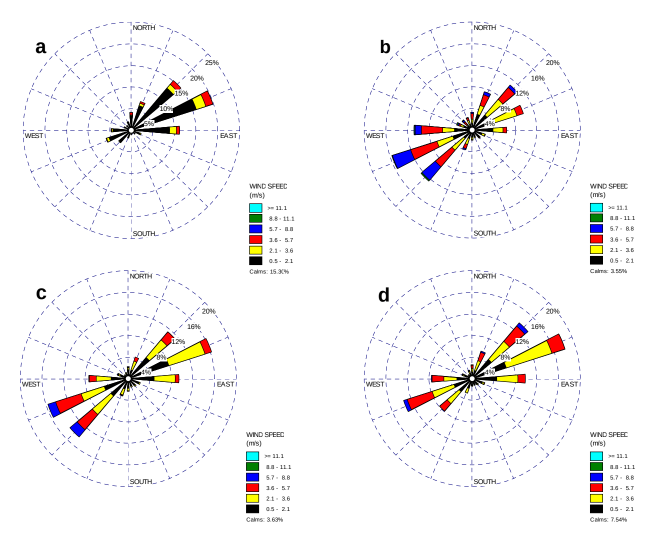


Fig. 3. Wind rose for surface winds at SDZ from December 2003 to November 2004 (plotted with WRPlot View software, http://www.weblakes.com/lakewrpl.html). (a) Winter (December 2003–February 2004); (b) Spring (March–May 2004); (c) Summer (June–August 2004); (d) Fall (September–November 2004)

3.3 The diurnal variation of aerosol optical properties

Figure 4 shows values of aerosol absorption coefficient, scattering coefficient, and SSA averaged for each hour of a day during the period from September 2003 to January 2005 as a function of local time. There were clear diurnal variations for all the variables. For the absorption coefficient, low values occurred in the early afternoon (around 13:00), and high values were observed during the night, which reflects the evolution of the boundary layer and its effect on the concentration of pollutants at the site. During the night, as the height of the planetary boundary layer (PBL) decreased, pollutants were entrapped in the lower levels of the atmosphere and caused high concentrations of aerosols at the surface, while during the daytime, as the PBL developed, the turbulent dilution led to a lower concentration of pollutants at the surface in the early afternoon. A local maximum value of the absorption

coefficient appeared in the morning (around 08:00), which corresponds to local emissions of light-absorbing material from human activities surrounding the site. The decrease of aerosol absorption coefficient from the midnight to the early morning (from 00:00 to 05:00) suggests a removal of aerosol particles by dry deposition. A similar diurnal pattern for scattering coefficients was obtained, except that no significant peak around 08:00 corresponding to local human activities was found. Consequently, the diurnal variation of SSA showed the highest values in the afternoon with a maximum at around 14:00, and SSA decreased to a minimum observed at 07:00. This pattern of SSA diurnal variability indicated the enhancement of a light-scattering aerosol loading during the afternoon, which might reflect secondary aerosol production through photochemical processes at the site. Both aerosol absorption and scattering coefficients measured at the SDZ

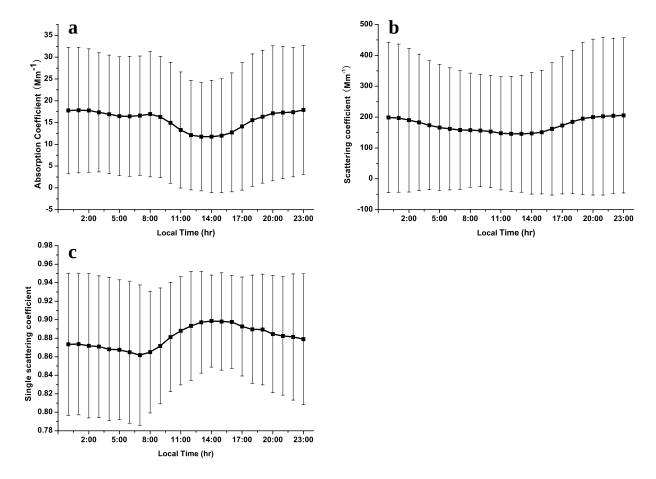


Fig. 4. Diurnal variability of aerosol optical properties at SDZ from September 2003 to January 2005, error bars represent S.D. (a) Absorption coefficient; (b) Scattering coefficient; (c) Single scattering albedo (SSA).

site showed similar diurnal patterns as those obtained at another GAW regional background station in the Yangtze delta region, the Lin'An station in the eastern part of China (Xu et al., 2002).

Further comparisons were conducted for summer and winter separately. The diurnal variations of both aerosol scattering and absorption coefficients remained similar to those illustrated in Fig. 4 for the entire experiment period. However the morning peak was more significant during the winter time, and it appeared later (about 09:00–10:00) in winter and earlier (~07:00) in summer compared with those calculated from the entire datasets.

The diurnal patterns during the summer time at the SDZ site were different from those of ozone and other trace gases measured at a mountainous site closer to Beijing during the summer of 2005 (Wang et al., 2006), who obtained higher concentrations of the gases in the afternoon (~16:00). We suspect that the distance of the measurement site from the heavily-polluted regions probably plays a very important role. In Wang's measurements, the site was relatively close to the urban area, about 50 km south to the center of Bei-

jing, and only about 15 km southwest to Changping (a satellite town of Beijing, with a population of \sim 110 000). However, the SDZ site was farther away from the urban area (about 150 km northeast of the Beijing urban area), and there are only limited natural and anthropogenic pollution sources within 30 km range surrounding the site (e.g., the town of Miyun County, with an area of \sim 24 km² and population of \sim 146 000 being about 55 km to the south). As our site is on a gentle slope of a small hill, the topography-induced local circulation may have some impact on the transport of the pollutants to the site, but the lack of an enhancement of aerosol extinction in the summer afternoon at the SDZ suggests that the upslope flow did not transport much urban pollutants to the measurement site.

3.4 The aerosol optical properties within different air masses

In order to further investigate the effects of different emission sources on aerosol optical properties, a trajectory clustering technique (Stunder, 1996) was applied in the analysis. In this work, we chose the period from December of 2003 to February of 2004 to represent the winter period and from June to August of 2004 to represent the summer period.

Backward air mass trajectories were calculated using an approach developed by Draxler (1997), and the meteorological data used were the NOAA global model final analysis archived dataset with a time resolution of 6 h. Backward air mass trajectories were calculated for 36 h back in time at 100 m above the ground.

The cluster-mean value of each parameter was calculated through the formula given below. It is composed of onehour average data centered at each trajectory start time for all trajectories within that cluster:

$$\overline{A}_i = \frac{1}{n} \sum_{j=1}^{n} A_{i,j}, i = 1, N$$
 (3)

where n represents the total number of trajectories in the ith cluster, $A_{i,j}$ represents the aerosol properties corresponding to the jth trajectory, and N represents the total number of clusters.

Figure 5 shows the mean cluster trajectories to the site for both the summer (June, July and August, 2004) and winter (December, 2003, January and February, 2004) periods. Tables 6 and 7 present the cluster-mean values of aerosol properties for each period respectively. For the summer period, the cluster analysis had all trajectories grouped into 7 clusters (Fig. 5a). Trajectory clusters labeled #1, #2, and #6 were from the southern sector of the site, and these air masses originated from the more heavily polluted regions. As Table 6 illustrates, cluster #1 was the most frequently observed flow pattern to the site in the summer, and the absorption and scattering coefficients of cluster #1 were quite high, with the mean values of 30.2 Mm⁻¹ for aerosol absorption coefficient and 313.2 Mm⁻¹ for aerosol scattering coefficient (see Table 6). A similar result was found for cluster #2, for which the cluster-mean absorption coefficient and scattering coefficients were 32.7 Mm⁻¹ and 308.7 Mm⁻¹ respectively. The SSA for both clusters had the same mean value of 0.89, which was the highest among all the clusters, and also was higher than the average SSA (0.81 and 0.793 respectively, see Table 3) reported for the Beijing urban areas (Bergin et al., 2001; Mao and Li, 2005). The average 36-hour backward trajectory of cluster #1 and cluster #2 indicated that the air masses for these two clusters were mainly from the heavily-polluted regions. For instance, the average 36-h backward trajectory for cluster #1 could be traced back to Beijing and its southern outlying areas, and the average trajectory for cluster #2 extended a longer distance (indicating more rapid transport) with the air masses passing through Beijing, Baoding, and Shijiazhuang, which are all regions with large pollution emissions in northern China.

Cluster #6 represented air masses coming from southeasterly directions. Along the trajectory pathway, big industrial cities, such as Tianjin and Tangshan, are located within a 200 km range from the site. The 36-h backward trajectory also suggested that the air masses passed over the Shandong

Table 5. The number of days with mist (1 km < visual range < 10 km) and fog (visual range < 1 km) and poorer visibility in each month from September 2003 to January 2005 at the SDZ site.

Month	Mist/fog (Day)	Visual range <10 km (Day)
September 2003	12	12
October 2003	4/1	5
November 2003	11/2	13
December 2003	2	2
Jan., 2004	1	1
February 2004	2	3
March 2004	4	6
Apr., 2004	5	6
May 2004	4	4
June 2004	4	4
July 2004	12	12
August 2004	8/1	9
September 2004	12/1	13
October 2004	8/2	10
November 2004	6	7
Dec., 2004	5/2	9
January 2005	3	3

Peninsula (one of the rapidly developing regions in China) and could have come across the Bohai sea to influence the site. As a consequence, the mean absorption and scattering coefficients for cluster #6 were the third highest among all the clusters, with a cluster-mean aerosol absorption, scattering coefficient, and SSA of 19.1 Mm⁻¹, 233.4 Mm⁻¹, and 0.89, respectively.

Cluster #3 was the second most frequently observed flow pattern to the site in the summer. The average 36-h backward trajectory of the cluster indicated that the wind speed was weak and the air masses for this cluster originated in regions close to the site because the pathway of the mean trajectory for this cluster was very short. Weak winds were often observed during periods of stable atmospheric conditions, and in this case the site was influenced more by local sources and the accumulation of pollutants. The mean absorption and scattering coefficients were 19.0 Mm⁻¹ and 117.5 Mm⁻¹, respectively, and the SSA was 0.84. The aerosol loading in the atmosphere for this cluster was lower than those with air masses coming from the heavily-polluted regions, such as cluster #1, #2 and #6, but was still rather high compared with those from relatively cleaner clusters, such as clusters #4, #5 and #7.

The clusters labeled #4, #5 and #7 represented the cases influenced by relatively clean air masses at the site according to mean trajectory pathways (Fig. 5a) and the corresponding values of aerosol properties (see Table 6). The absorption and scattering coefficients were significantly lower than the previously discussed clusters, and the SSA for these three clusters ranged from 0.78 to 0.82, which were also significantly lower than the previously discussed clusters.

Cluster No.	SSA (at 525 nm)	Number of trajectories	$\sigma_{\rm abs} {\rm Mm}^{-1}$	Number of trajectories	$\sigma_{\rm sca}{ m Mm}^{-1}$	Number of trajectories
1	0.89(0.04)	106	30.24(15.93)	106	313.22(220.56)	116
2	0.89(0.05)	29	32.67(11.59)	29	308.66(159.49)	31
3	0.84(0.07)	80	18.97(10.69)	82	117.53(98.27)	90
4	0.78(0.08)	36	9.30(8.73)	39	32.56(33.38)	40
5	0.82(0.09)	29	10.11(6.28)	29	49.72(30.36)	29
6	0.89(0.06)	24	19.09(14.74)	24	233.36(200.68)	27
7	0.80(0.10)	17	12.17(10.26)	17	58.60(69.85)	24

Table 6. The mean optical properties for each cluster in summer (June, July and August of 2004).

Table 7. The mean optical properties for each cluster in winter (December 2003, January and February 2004).

Cluster No.	SSA (at 525 nm)	Number of trajectories	$\sigma_{\rm abs} {\rm Mm}^{-1}$	Number of trajectories	$\sigma_{\rm sca} {\rm Mm}^{-1}$	Number of trajectories
1	0.92(0.03)	22	11.70(14.31)	22	154.72(212.33)	23
2	0.91(0.04)	49	9.02(7.98)	49	102.02(108.69)	49
3	0.93(0.03)	14	27.85(20.87)	15	414.57(396.15)	16
4	0.89(0.06)	37	3.76(2.59)	37	35.23(34.84)	37
5	0.91(0.03)	10	16.04(10.82)	16	213.83(198.59)	12
6	0.94(0.01)	19	21.83(13.79)	22	387.93(239.88)	22
7	0.92(0.02)	19	1.96(1.53)	19	23.00(17.15)	21
8	0.91(0.03)	34	2.38(1.56)	34	22.72(13.59)	35
9	0.90(0.03)	21	3.89(3.81)	22	35.53(29.97)	29
10	0.92(0.03)	23	1.50(1.38)	23	18.28(13.80)	24

The above analysis revealed significant differences in aerosol properties among the different clusters in summer. The clusters with the trajectories passing through polluted regions had not only higher absorption and scattering coefficients but also higher SSA values, while the clusters with mean trajectories passing through cleaner regions had lower absorption and scattering coefficients and lower SSA. This behavior means that the aerosols coming from the polluted regions were more efficient at scattering light than those coming from clean regions during the summer period.

The average trajectories for each cluster in winter showed different flow patterns from those in summer. The total trajectories were clustered into 10 groups (Table 7). The pathway of air masses for clusters #1, #2, #4, and #7–#10 were from the northwest, while the air masses for cluster #3 was from the west, and those for clusters #5 and #6 took curved pathways to the north and south, respectively. According to the map of the site shown in Fig. 1, most cities with dense population and high industrial activities are in the sectors from the west to the southeast direction of the station, thus, clusters #4 and #7–#10 mainly reflect the relatively cleaner air masses at the site, while clusters #3 and #6 indicate the directions affected by air masses coming from polluted regions. The cluster-mean aerosol absorption and scattering coefficients are given in Table 7, which demonstrated again

that the aerosol optical properties were tightly related to the pathways of air masses for each cluster. Low cluster-mean aerosol absorption and scattering coefficients were found for clusters #4 and #7-#10, while high aerosol absorption and scattering coefficients were observed for clusters #3 and #6. The pathways for clusters #1 and #2 were very similar. Both passed along the way slightly north to Zhangjiakou city. The cluster-mean levels of aerosol absorption and scattering coefficients for trajectory clusters #1 and #2 were higher than those of clusters #4 and #7-#10, a probable explanation for this feature was that the air masses of these two clusters were partly influenced by the emissions from the highly polluted city of Zhangjiakou and its surroundings. The pathway for cluster #5 was complicated, as revealed by the 36-h backward trajectory (Fig. 5b). The cluster mean trajectory shows the air masses that originated from the relatively cleaner regions in the northwest, and then passed through more polluted regions in the northeast and east, and the mixed influence of the cleaner and polluted air masses resulted in a relatively high level of aerosols at the site as given in Table 7.

Similar to the summer cases, the SSA values in winter were also slightly higher when the air masses were associated with polluted pathways and relatively lower when the air masses were coming from cleaner regions.

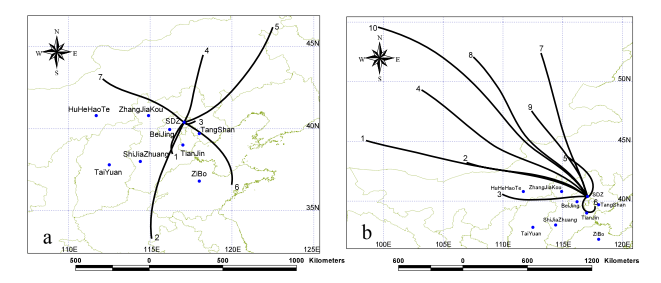


Fig. 5. The cluster mean 36h backward trajectories for summer and winter periods at SDZ (a) Summer period (June–August 2004), (b) Winter period (December 2003–February 2004).

Table 8. The mass concentration and percentage of selected chemical species in fine particles sampled during the summer period (17 July–2 August 2004) at SDZ site and some reported results obtained at Beijing urban sites.

Sample No.	Mass Conc. $(\mu g/m^3)$	$NO_3^ (\mu \text{ g/m}^3)$	SO_4^{2-} $(\mu g/m^3)$	$NH_4^+ (\mu g/m^3)$	OC $(\mu g/m^3)$	EC $(\mu g/m^3)$	Ions ^a /Mass (%)	EC/ Mass (%)	OC/Mass (%)	SSA (at 525 nm)
Summer										
#1 (polluted)	153.89	15.76	57.65	33.99	11.12	1.97	69.8	1.3	7.2	0.94
#2 (polluted)	132.66	16.21	58.72	35.67	6.64	2.59	83.4	2.0	5.0	0.93
#6 (clean)	32.79	4.85	7.15	4.24	4.61	0.89	49.5	2.71	14.1	0.87
#7 (clean)	33.95	2.51	9.69	4.02	3.43	0.76	47.8	2.2	10.1	0.87
#8 (clean)	12.84	0.14	3.20	1.02	1.58	0.29	34.0	2.2	12.3	0.83
#9 (clean)	20.54	2.32	3.50	1.70	3.36	0.86	36.6	4.2	16.4	0.81
Chegongzhuang ^b , Beijing	74.0	4.6	17.1	5.7	13.4	6.3	37.1	8.5	18.0	
BNU ^c , Beijing	73.3	12.2	16.0	10.4	11.5	5.2	52.7	7.1	15.7	
CS ^c , Beijing	82.2	13.3	19.2	11.0	9.3	6.6	52.9	8.0	11.31	
YH ^c , Beijing BMEMC ^d , Beijing	75.4 80.3	13.2	19.7	9.8	11.2 9.4	5.9 4.3	56.6	7.8 5.4	14.85 11.7	

a Ions= $(NO_3^- + SO_4^2^- + NH_4^+)$

Filter sampling was conducted during subsets of the measurement period, and limited analyses of aerosol chemical composition and the corresponding aerosol optical properties are presented in Table 8. For comparison, we also list some results of aerosol chemical composition in PM_{2.5} (particles with aerodynamic diameter less than 2.5 μ m) obtained during the summer time at Beijing urban sites in Table 8.

The filter samples were collected at SDZ during the period from 17 July to 2 August 2004. Among the samples,

the samples #6-#9 were sampled on the "clean" days and the samples #1 and #2 on the polluted days. They represented cases when the site was influenced by polluted or cleaner air masses based on the calculated backward trajectories and the mass concentrations of each sample. Table 8 shows that the sum of major light scattering aerosol components, such as sulfate, nitrate, ammonium and OM (using a factor of 1.4 to estimate the mass of organic matter OM (He et al., 2001; Malm et al., 1994) accounted for 80%–90%

b He et al. (2001)

^c Sun et al. (2004)

^d Yu et al. (2006)

of total mass of $PM_{2.1}$ (particles with aerodynamic diameter less than $2.1~\mu m$) for samples collected in polluted days and 51%-69% in cleaner days. Among the major light scattering aerosol components, the sum of sulfate, nitrate and ammonium accounted for about 70% or more of total $PM_{2.1}$ mass when the site was under polluted conditions, and less than 50% for the cleaner days. However, OC fraction was relatively small, representing only about 5%-7% of the $PM_{2.1}$ mass under polluted conditions and 10%-16% of the mass during the cleaner days. Moreover, the ratio of elemental carbon (light absorbing aerosol) to $PM_{2.1}$ mass was relatively lower when the site was under polluted conditions.

The fractions of OC under polluted conditions were lower than those obtained during the summer time in the Beijing urban areas, although the concentration levels of OC were still comparable to the reported values for the regions (He et al., 2001; Sun et al., 2004; Yu et al., 2006). Different from OC, however, the fraction of the sum of sulfate, nitrate and ammonium in PM2.5 under polluted conditions was significantly higher than the reported values for the summer time measurements (He et al., 2001; Sun et al., 2004; see Table 8). Because OC is an important contributor to aerosol light scattering (Novakov et al., 2005), the lower fraction of OC under polluted conditions suggested that the corresponding higher SSAs were mostly due to the high fraction of secondary aerosols with sulfate, nitrate and ammonium in fine particles. On the other hand, when the site was under "clean" conditions, the dominant residential emissions (with more EC being released due to the combustion of coal at lower temperature) in the relatively cleaner regions, such as Inner Mongolia (Cao et al., 2006), result in a relatively higher fraction of elemental carbon and lower SSA of aerosols. This analysis is consistent with the results of the trajectory clustering analysis that indicated aerosols with air masses from polluted regions had more light scattering ability during the summer time.

It should be noted that the relationship between aerosol SSA and chemical compositions depends on many factors, such as size distributions, shape and mixing state of aerosols (Fuller et al., 1999). To fully understand the variability of SSA with the different air masses, further investigations are still needed.

4 Conclusions

From April 2003 to January 2005, in situ measurements of aerosol optical properties were conducted at a rural site in Northern China, the Shangdianzi Global Atmosphere Watch (GAW) regional station, about 150 km from Beijing. Mean values (standard deviation) of the scattering and absorption coefficients at this rural site in northern China for the entire measurement period were 174.6 Mm⁻¹ (189.1 Mm⁻¹) and 17.5 Mm⁻¹ (13.4 Mm⁻¹), respectively. These values were about one-third of reported value for scattering coefficient

and about one-fourth of that for absorption coefficient obtained in the Beijing urban area. The mean SSA for the entire period was 0.88 (0.05), which was about 0.07 higher than the value reported by Bergin et al. (2001) for Beijing urban area, and also higher than the value (0.85) in climate modeling simulations for China and India performed by Menon et al. (2002). Seasonal variations are observed for different aerosol properties. Both the aerosol absorption and scattering coefficients were lowest in winter, while the highest values appeared in summer for the absorption coefficient and in fall for the scattering coefficient. The mean SSA showed the lowest value in spring (0.85) and highest in winter (0.91).

The trajectory clustering analysis revealed that the scattering and absorption coefficients and SSA were all high when the air masses came from the densely populated and highly industrial areas (especially during the summer time). The SSA's associated with the polluted air masses were not only higher than those with the air masses coming from the "clean" directions, but also higher than the reported values measured at the pollution source regions. This fact suggests that there may be substantial secondary aerosol production during the transport of air masses from polluted regions during summer time. To fully understand the exact reasons for these results, further investigations on size distributions, chemical compositions, and mixing state of aerosols are needed. Aerosol optical properties measured at this rural area in northern China could serve as a basis for radiative forcing estimation and climate modeling for the region.

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