Observations of aerosol optical properties at a coastal site in Hong Kong, South China

Jiaping Wang$^{1,2,3,4}$, Aki Virkkula$^{1,2,3,5,6}$, Yuan Gao$^{3,7}$, Shuncheng Lee$^{4}$, Yicheng Shen$^{1,2,3}$, Xuguang Chi$^{1,2,3}$, Wei Nie$^{1,2,3}$, Qiang Liu$^{1,2,3}$, Zheng Xu$^{1,2,3}$, Xin Huang$^{1,2,3}$, Tao Wang$^{1}$, Long Cui$^{4}$, and Aijun Ding$^{1,2,3}$

$^1$Joint International Research Laboratory of Atmospheric and Earth System Sciences, Nanjing, China
$^2$Institute for Climate and Global Change Research & School of Atmospheric Sciences, Nanjing University, Nanjing, 210023, China
$^3$Collaborative Innovation Center of Climate Change, Jiangsu Province, China
$^4$Department of Civil and Environmental Engineering, The Hong Kong Polytechnic University, Hung Hom, Kowloon, Hong Kong SAR, China
$^5$Finnish Meteorological Institute, Helsinki, Finland
$^6$Department of Physics, University of Helsinki, Helsinki, Finland
$^7$Department of Civil Engineering, the Chu Hai College of Higher Education, Castle Peak Bay, Hong Kong SAR, China

Correspondence to: Jiaping Wang (jiaping16@126.com) and Shuncheng Lee (shun-cheng.lee@polyu.edu.hk)

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Abstract. Temporal variations in aerosol optical properties were investigated at a coastal station in Hong Kong based on the field observation from February 2012 to February 2015. At 550 nm, the average light-scattering ($151 \pm 100 \, \text{Mm}^{-1}$) and absorption coefficients ($8.3 \pm 6.1 \, \text{Mm}^{-1}$) were lower than most of other rural sites in eastern China, while the single-scattering albedo (SSA = 0.93 ± 0.05) was relatively higher compared with other rural sites in the Pearl River Delta (PRD) region. Correlation analysis confirmed that the darkest aerosols were smaller in particle size and showed strong scattering wavelength dependencies, indicating possible sources from fresh emissions close to the measurement site. Particles with $D_p$ of 200–800 nm were less in number, yet contributed the most to the light-scattering coefficients among submicron particles. In summer, both $\Delta \text{BC} / \Delta \text{CO}$ and $\text{SO}_2 / \text{BC}$ peaked, indicating the impact of nearby combustion sources on this site. Multi-year backward Lagrangian particle dispersion modeling (LPDM) and potential source contribution (PSC) analysis revealed that these particles were mainly from the air masses that moved southward over Shenzhen and urban Hong Kong and the polluted marine air containing ship exhausts. These fresh emission sources led to low SSA during summer months. For winter and autumn months, contrarily, $\Delta \text{BC} / \Delta \text{CO}$ and $\text{SO}_2 / \text{BC}$ were relatively low, showing that the site was more under influence of well-mixed air masses from long-range transport including from South China, East China coastal regions, and aged aerosol transported over the Pacific Ocean and Taiwan, causing stronger abilities of light extinction and larger variability of aerosol optical properties. Our results showed that ship emissions in the vicinity of Hong Kong could have visible impact on the light-scattering and absorption abilities as well as SSA at Hok Tsui.

1 Introduction

Atmospheric aerosol strongly affects the earth’s radiative balance by scattering and absorbing incoming solar radiation, which, however, is still a large source of uncertainties in global climate forcing assessment (Stocker et al., 2013). The aerosol optical properties are responsible for the direct aerosol radiative forcing, depending on aerosol chemical composition and microphysical properties. Relative to another major component of radiative forcing, greenhouse gases, the shorter atmospheric lifetime of aerosols leads to more localized effects and regional differences in aerosol optical properties. Due to the spatial and temporal differences
of aerosol optical properties caused by the complex distribution of tropospheric aerosols, field monitoring of aerosol optical properties in different regions around the world is critical for exploring the variations in aerosol radiative forcing. Among the major aerosol radiative forcing drivers, mineral dust, sulfate, nitrate, and organic carbon generally have negative radiative forcing. Contrarily, absorbing aerosols, like black carbon (BC), can strongly absorb visible light, enhancing the warming effect of the atmosphere (Jacobson, 2001; Babu and Moorthy, 2001; Ding et al., 2016).

Light absorption and scattering of different kinds of aerosols have distinct wavelength dependencies that are approximately proportional to $\lambda^{-\text{AAE}}$ or $\lambda^{-\text{SAE}}$, respectively, where $\lambda$ is the wavelength and AAE and SAE are the Ångström exponents of absorption and scattering, respectively. Hence, the wavelength dependency of aerosol light scattering and absorption has been recognized as an efficient index to distinguish aerosol types (e.g., Russell et al., 2010; Moosmüller and Chakrabartty, 2011; Devi et al., 2016). For instance, BC can strongly absorb light at all visible wavelengths, while other light-absorbing aerosols (some organic aerosol, soil, and dust) absorb more blue light than red light (Moosmüller et al., 2011; Bond et al., 2013; Ding et al., 2016). Therefore, the absorption Ångström exponent (AAE) is often related to the dominant absorbing aerosol type for a mixture of aerosols (Cazorla et al., 2013). The AAE in externally mixed BC-dominated regions has been reported to be around 1 (Hegg et al., 2002; Bond and Bergstrom, 2006; Bond et al., 2013), while it is greater than 1 for some organic aerosol from biomass smoke and mineral dust due to their diverse light absorbing abilities at different wavelength ranges (Kirchstetter et al., 2004; Russell et al., 2010; Valezuela et al., 2015; Devi et al., 2016). Moreover, studies have shown that AAE of BC has a large variability, depending on the size of BC cores and coating thickness (e.g., Lack and Cappa, 2010). For non-coated BC with small diameters (e.g., 10 nm), AAE is close to 1, but large BC cores can have AAE less than 1 (e.g., Gyawali et al., 2009; Lack and Cappa, 2010). For coated BC particles, laboratory measurements of Schnaiter et al. (2005) reported that thickly coated BC by $\alpha$-pinene plus ozone SOA could decrease the AAE of up to 0.8. Coating of BC by purely scattering material may also result in AAE up to about 1.8 (Gyawali et al., 2009; Lack and Cappa, 2010). The scattering Ångström exponent (SAE) is often regarded as a qualitative indicator of the dominating particle size, that is, large values (SAE $> 2$) indicate a large contribution of small particles and small values (SAE $< 1$) a large contribution of large particles. For instance, Delene and Ogren (2002) reported that the influence of large sea-salt particles led to lower SAE. However, this interpretation is not quite unambiguous, as was shown, for example, by Schuster et al. (2006) and Virkkula et al. (2011). The SSA is the ratio of scattering to extinction coefficient, i.e., the sum of scattering and absorption coefficients. It equals 1 for purely scattering aerosol and is clearly lower, approximately 0.3, for pure BC particles (e.g., Schnaiter et al., 2003; Mikhailov et al., 2006). SSA varies significantly for smoke of different origin and age and correlates with the presence of BC in the combustion products (e.g., Dubovik et al., 2002).

There are several ways to assess the sources of aerosols, for instance by comparing observed particle concentrations with other tracers. CO is a by-product of the incomplete oxidation. Due to its long lifetime (about 1–2 months) in the troposphere, CO can act as a tracer of anthropogenic emissions (Jennings et al., 1996). A strong positive correlation between BC and CO has been found in previous studies concerning source identifications (Pan et al., 2011; Jennings et al., 1996). The BC / CO ratio is considered a good indicator to determine BC emission and to recognize source characteristics. Also, the emission ratio of BC and CO varies significantly from different sources, making it an effective index for validating emission inventories (Girach et al., 2014). The SO$_2$ / BC ratio can be also used for assessing the sources since both BC and SO$_2$ are emitted in fossil fuel combustion (Bond et al., 2013).

The Pearl River Delta (PRD) region in southern China has undergone fast industrialization with increasing emissions of particulate and gaseous pollutants (Wang et al., 2003). In particular, the growing crisis of high particulate matter (PM) levels in the Pearl River Delta (PRD) region is of great concern due to its adverse effects on regional and continental atmospheric environment (Wang et al., 2009; Ding et al., 2013; Lam et al., 2005; Liu and Chan, 2002; Verma et al., 2010). Hong Kong is a typical coastal city in the PRD region. Under the influence of the East Asian monsoon, this region is controlled by the southerly winds bringing marine inflow from the South China Sea in summer, while in winter it is downwind from the North China Plains and East China and is dominated by the continental outflow (Ding et al., 2013; Lam et al., 2001; Zhou et al., 2013). Thus, it is an ideal place for exploring the characteristics of optical properties for continental and marine aerosols.

There have been studies concerning aerosol optical properties and light-absorbing aerosols in the PRD region. Man and Shih (2001) did field observations of light-scattering and absorption coefficients from September 1997 to April 1999 in Hong Kong. Cheng et al. (2006a) investigated the seasonal variation patterns of BC concentrations in Hong Kong as well as the potential sources of BC by continuous measurement from June 2004 to May 2005, using a model AE-42 Aethalometer (Magee Scientific Inc., Berkeley, CA). Cheng et al. (2008) presented the 1-month record of aerosol optical measurements with related chemical apportionment at Xinken in the PRD region and reported a relatively low SSA at this polluted rural site. Mixing states of light-absorbing aerosols were also investigated using optical closure experiments during the campaign (Cheng et al., 2006b; Tan et al., 2016). However, long-term observations of several key aerosol optical properties (including wavelength dependencies of light scattering and absorption SSA), and studies on
the relationships between optical properties and particle size as well as their quantitative linkage to multi-scale transport) have been limited in Hong Kong over the past decade.

In this study, we aim at demonstrating the temporal variations of aerosol optical properties at a coastal station in Hong Kong and investigating the relationships between aerosol optical properties and size distributions based on field observations. Source analyses are conducted by comparing observed BC–CO ratios as well as the SO2–BC ratios. Transport pattern and origins of aerosols were quantitatively studied based on Lagrangian particle dispersion modeling (LPDM). Characteristics of local aerosol optical properties dominated by different aerosol source regions were also compared and illustrated.

2 Methodology

2.1 Sampling site

The Hok Tsui (HT) monitoring station is situated on the southeast tip of Hong Kong Island facing the South China Sea (22.22° N, 114.25° E 60 m above sea level) with an almost vertical drop to the sea. This station has a view of the sea for over 180° from the northeast to the southwest and is 20 km away from the urban area of Hong Kong on the northwest. Owing to the characteristics of the location mentioned above, it is an ideal background monitoring site for identifying both the long-range transport of polluted continental/marine air mass caused by anthropogenic emissions and relatively clean marine air mass in different seasons. For more details about the HT site, please refer to Wang et al. (2009) and papers cited therein.

2.2 Absorption measurement

Light-absorption measurement was conducted using a model AE-31 Aethalometer (Magee Scientific Company, Berkeley, California, USA) from 1 February 2012 to 30 September 2013 and 1 March 2014 to 28 February 2015. Sample air was obtained through a stainless steel inlet with a PM2.5 cut-off, protected with a rain cap. Prior to entering the instrument, sample air was heated to ensure a moderate relative humidity. The sample inlet was approximately 1.5 m above the roof of the measurement station building, which was about 4 m above the ground. The sample flow provided by the internal pump was set to 4.0 L min⁻¹. The AE-31 Aethalometer performs continuous measurements of BC concentrations at seven wavelengths (370, 470, 520, 590, 660, 880 and 950 nm) with a time resolution of 5 min. In this work, without specific notes, BC concentrations refer to the aethalometer data measured at λ = 880 nm. Sample flow on the Aethalometer display was checked once a week to ensure the flow was within 0.2 L min⁻¹ of previous week and flow calibration was conducted once a month using an independent flowmeter. The inlet cyclone was cleaned every month.

In order to correct the systematic errors of filter-based absorption technique, the light absorption coefficients (σap) at all wavelengths were calculated by using the method presented by Collaud Coen et al. (2010), where the Cref factor was set to be 4.26 according to the value from Cabauw (CAB) station reported in the same paper. CAB station is located near populated and industrialized areas, which was to some extent similar to Hok Tsui station (near most of cities in the Pearl River Delta region). The reported average Cref value at CAB was 4.26 ± 0.11, and it varies from 2.60 to 4.75 (Collaud Coen et al., 2010). There was no MAAP (Multi-Angle Absorption Photometer) or any other reference absorption instrument available, so determining Cref at Hok Tsui was not possible and the published mean Cref at CAB station was used. However, to present an upper estimate for σap, the Cref calculated as 3.51 for the clean marine site of Mace Head (MHD); Collaud Coen et al., 2010), was also used and the respective average σap and SSA are presented in the discussions. Since the Cref is responsible for the largest uncertainty in the calculation of σap (Collaud Coen et al., 2010) we did not make further uncertain analyses by using the uncertainties related to the other factors within the algorithm. Absorption coefficients were presented under standard temperature and pressure (STP; 273.15 K, 1013 hPa). Measured BC concentrations were corrected following the algorithm presented by Virkkula et al. (2007).

2.3 Scattering measurement

Light-scattering coefficients (σsp) at wavelength of 450, 550 and 700 nm were measured using an integrating nephelometer (model 3563, TSI Inc, St. Paul, MN, USA). The averaging time was set to 5 min. Calibration was conducted once a month using CO2 and filtered air as described in the user manual. An internal heater was used to maintain a moderate relative humidity during measurement. Raw σsp data were corrected for truncation errors following the method from Anderson and Ogren, (1998), where the scattering coefficients were determined by calculating the Ångström exponents from uncorrected scattering coefficients and the correction factors with no-cut inlet. Scattering coefficients were then corrected to STP using pressure and temperature readings from the nephelometer.

The σsp and σap data were used for calculating SSA as SSA = σsp/(σsp + σap). The nephelometer took its sample from a total suspended particle inlet (TSP) but the Aethalometer through a PM2.5 inlet, so it may seem somewhat uncertain which size range the SSA represents. However, BC is the most important light-absorbing constituent in aerosol particles and it is well known that it is in the submicron size range. In larger particles there might be some light-absorbing dust particles, but their contribution at this site can be considered to be negligible. Therefore it is reasonable to claim that the absorption coefficients derived from the aethalometer data represent absorption in the full TSP size range.
range even if there was a PM$_{2.5}$ inlet for the Aethalometer. Since the scattering coefficients were measured after a TSP inlet, it is also reasonable to say that the SSA represents that of TSP.

2.4 Particle size measurement and the use of the size distributions

An Ultrafine Particle Monitor (UFPM, model 3031, TSI Inc.) was used to measure the number size distribution of particles in the size range of 20 to 800 nm with six size bins of mobility diameter: 20–30, 30–50, 50–70, 70–100, 100–200, and 200–800 nm. The operating principle of a UFPM is based on the diffusion charging of particles, followed by size segregation within a differential mobility analyzer (DMA) and the detection of the aerosol via a sensitive electrometer. The UFPM was equipped with a model 3031200 environmental sampling system. The sample inlet was placed 2.0 m above the ground. Ambient air was continuously drawn through a size selective PM$_{10}$ inlet at a standard flow rate of 16.7 L min$^{-1}$. The sample then passed through a PM$_1$ cyclone to remove larger particles. The main sample stream was subsampled into the UFPM at a flow rate of 5.1 L min$^{-1}$. A Nafion dryer was installed upstream of the UFPM to ensure proper conditioning of the aerosol and to minimize effects due to water vapor. The remaining 11.7 L min$^{-1}$ of make-up air, drawn through a vacuum pump and exhausted, was routed through the Nafion dryer as purge air. The averaging time was set to 15 min.

The total mass concentrations of particles with a mobility diameter less than 800 nm were calculated using the following equation:

$$PM_1 = \sum_{i=1}^{n} N_i \rho_i \frac{\pi}{6} D_{p,i}^3,$$

where $N_i$ is the number concentration in each size bin, $\rho_i$ is the density of particles assumed to be 1.7 g cm$^{-3}$, and $D_{p,i}$ is the geometric mean of the upper and lower limit diameter in each size bin.

For spherical particles the aerodynamic diameter ($D_a$) is calculated from the mobility diameter ($D_m$) as $D_a = D_m \sqrt{\frac{\rho_p}{\rho_0}}$, where $\rho_p$ is the density of the particle and $\rho_0$ its the density of water. For $D_m = 0.8 \mu$m and $\rho_p = 1.7 \text{ g cm}^{-3}$, this yields $D_a = 1.0 \mu$m. In the results, therefore, the mass concentration calculated from the number size distributions was denoted as $PM_1$.

The size distributions were used for calculating scattering coefficients from the following equation:

$$\sigma_{sp}(\lambda) = \int Q_{sp}(\lambda, D_p, m) \frac{\pi D_p^2}{4} n(D_p) dD_p,$$

where the scattering efficiencies ($Q_{sp}$) were calculated by using the BHMIE code (Bohren and Huffman, 1983). We assumed that the $D_p$ of each particle is equal to the geometric mean of the upper and lower limit diameter in its size bin for modeling, and the aerosol is ammonium sulfate with the refractive index $m = m_t = 1.52$ (Chamaillard et al., 2006). The refractive index used in the modeling could, in principle, be varied and iterated until the measured and modeled scattering coefficients match, as was done, for example, by Virkkula et al. (2011). However, due to the different size ranges and low number of size bins of the size distributions, this kind of iteration is not reasonable for the data in this work.

Both the PM$_1$ and the $\sigma_{sp}$ calculated from the number size distributions have uncertainties due to the uncertainties of the UFPM. The first is the wide range of particle diameters within the size bins and the use of the geometric mean of the bin limits for the whole bin. This yields the highest uncertainty for the bin that measures particles in the size range 200–800 nm that can easily be calculated assuming all particles in that size range were 800 nm instead of the geometric mean 400 nm. This calculation is theoretical in the real atmosphere, however, and yields unrealistically high uncertainties and will not be analyzed further. Another source of uncertainty is related to the instrument itself. Hillemann et al. (2014) found that the number of concentrations measured by UFPM are typically within a range of ±20% from the reference values measured with a scanning mobility particle sizer (SMPS). Also, Gómez-Moreno et al. (2015) compared the UFPM with an SMPS and found that the size distributions measured by UFPM and SMPS were similar in the sense that the peak concentrations were observed at the same size. In the same study it was also observed that in the size channels corresponding to particle diameters <100 nm the UFPM overestimated the number concentrations and in the two largest channels it underestimated the number concentrations. These are the channels that measure the particles that have the highest mass and that scatter light most efficiently. It may therefore be argued that both the PM$_1$ and the modeled $\sigma_{sp}$ are underestimated.

It was mentioned above that the PM$_1$ concentrations were calculated using the density of 1.7 g cm$^{-3}$, which deserves reasoning. The densities of major inorganic aerosol compounds such as ammonium sulfate and sodium chloride are 1.76 and 2.165 g cm$^{-3}$ (e.g., Tang, 1996). Zhang et al. (2008) estimated that the density of sulfuric acid-coated soot is 1.7 g cm$^{-3}$. Ambient aerosols contain many unknown compounds (such as organics) and also some water (even after drying to RH < 50%). Therefore, densities of ambient aerosols can vary within a certain range based on their compositions. Densities of real atmospheric aerosols have been measured in several campaigns. Quinn et al. (2001) determined aerosol densities on a cruise across the Atlantic Ocean. The density of submicron aerosols, averaged from observations at very different regions, was 1.73 ± 0.24 g cm$^{-3}$. Pitz et al. (2003) determined the mean apparent particle density of 1.6 ± 0.5 g cm$^{-3}$ for urban aerosol. Saarikoski et al. (2005) found that at a boreal forest site the average density was 1.66 ± 0.13 g cm$^{-3}$. Based on these publications it is reason-
able to use the density value of 1.7 g cm\(^{-3}\) for the estimation of aerosol mass concentration from the number size distributions of particles smaller than 800 nm of the mobility diameter. It has to be noted. However, there is uncertainty, in the aerosol mass concentration since the density was not measured at this site.

2.5 Supporting measurements

CO data were used to help analyze aerosol sources since they typically originate from incomplete combustion like BC. Hourly mixing ratios of carbon monoxide were measured with a nondispersive infrared absorption instrument (Teledyne API model 300) at Hok Tsui station.

In addition to the measurements at the HT station, the following supporting data measured at two nearby sites were used in the analyses. \( \text{SO}_2 \) is the precursor of sulfate, the most important light-scattering constituent and it is also one of the major pollutants of ship emission. \( \text{PM}_{2.5} \) concentrations can be used for a semi-quantitative quality check of the aerosol mass concentrations calculated from the size distributions. Hourly \( \text{SO}_2 \) and \( \text{PM}_{2.5} \) concentrations at Eastern Station (about 7 km away from HT station, the location shown in Fig. 1b), were downloaded from the open-access dataset from the website of the Hong Kong Environmental Protection Department (HKEPD).

The hourly averaged meteorological parameters, including air temperature, relative humidity (RH), wind direction, wind speed, and precipitation were obtained from the dataset in the HKEPD in which meteorological data from the near-site meteorological station (Waglan Island, WGL) were used for analyzing this paper. The location of the WGL station is shown in Fig. 1b.

2.6 Backward Lagrangian particle dispersion modeling (LPDM)

Transport and dispersion simulations were conducted using a Lagrangian particle dispersion modeling (LPDM) following the method developed by Ding et al., (2013). LPDM was conducted by using the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model, developed in the Air Resource Laboratory (ARL) of the US National Oceanic and Atmospheric Administration (Draxler and Hess, 1998; Stein et al., 2015). In each simulation, particles were released at a height of 100 m above the ground level at the site and went backward in time for a 7-day period. LPDM calculations were driven with GDAS (Global Data Assimilation System) data (http://ready.arl.noaa.gov/HYSPLIT.php). Particle positions were calculated in each hour and gridded concentrations were in a spatial resolution of 0.01° latitude by 0.01° longitude.

Knowing the transport characteristics of air masses, the next step was to explore the source profile of light absorbing particles affecting the regional aerosol optical properties in Hong Kong. Since BC is the most significant light-absorbing constituent of aerosols, the PSC of BC to observed air masses was calculated using the MIX Asian emission inventory (Li et al., 2017) together with LPDM results. The MIX emission inventory has a horizontal grid resolution of 0.25° × 0.25° in longitude and latitude and it is considered the anthropogenic emissions from transportation, residential, industry, and power generation in continental area. In each grid cell, the emission rate was multiplied by the footprint retrorplume, and the sum of this potential source contribution of all grid cells can provide the total BC concentration resulting from emissions during a certain period (Ding et al., 2013). The maps of averaged source contribution profile of BC in different seasons were calculated covering 70–140° in longitude and 0–50° in latitude. This method to calculate the PSC of target pollutants has been adopted in a previous study by Ding et al. (2013). The major advantage of this method is that it captures the potential contribution of target pollutants to the receptor due to the transport of air mass containing the information of anthropogenic emissions.

In this study, the MIX emission inventory provided relatively high spatial resolution of BC emission rates, considering its major anthropogenic sources in China and nearby Asian countries. However, marine emission is not included in the MIX database. To investigate the possible influence of marine sources, like ship emissions, on the observed aerosol concentrations at this coastal site, we used the observed aerosol concentrations together with the LPDM footprint. We used the following concentration-weighted equation to calculate the potential source contribution from each grid cell:

\[
A_{x(i,j)} = \frac{\sum_{t=1}^{n} (x_t \cdot R_t(i,j))}{\sum_{t=1}^{n} R_t(i,j)},
\]

Table 1. Statistical summary of data measured at Hok Tsui station. Scattering coefficients \( (\sigma_{sp}) \) and absorption coefficients \( (\sigma_{ap}) \) at \( \lambda = 550 \text{ nm} \) are corrected to STP (1013 mbar, 273.15 K), Ångström exponents of scattering and absorption (SAE, AAE), single-scattering albedo (SSA), total particle number concentration, \( (N_{\text{total}}) \) geometric mean diameter (GMD), and \( \text{PM}_1 \).
Figure 1. (a) Map showing the location of Hok Tsui (HT) monitoring station with emission inventory in Asia. (b) Locations of monitoring stations mentioned in this paper and (c) wind rose plot at WGL in Hong Kong.

where $x$ is the selected optical property or other parameters, and we chose $\sigma_{ap}$, $\sigma_{sp}$ and PM$_1$ in this study. $R$ represents the retroplume with 3-day backward time, while $i$ is the time step and $n$ is the total number of the time steps. The interpretation of Eq. (3) is that it shows the average value of the property $x$ observed at the receiving site when air masses have come from over grid cell $i, j$. The method is analogous to that presented by Stohl et al. (1996) and the concentration-weighted trajectory (CWT) methods reviewed by Cheng et al. (2015). The major difference is that in the present approach, the footprints were used instead of single trajectories for each time step.

3 Results and discussions

3.1 Aerosol optical properties and their relationships with particle size

3.1.1 Overall results of aerosol optical properties and related parameters

Table 1 shows a basic statistical summary of all measured parameters. The light absorption coefficients at $\lambda = 550$ nm were interpolated between the $\sigma_{ap}$ at 520 and 590 nm. The mean absorption and scattering coefficients at $\lambda = 550$ nm during the whole measurement period were $8.3 \pm 6.1$ and $151 \pm 100 \, \text{Mm}^{-1}$, respectively. As mentioned in the methods, the above-mentioned $\sigma_{ap}$ was calculated by using the $C_{\text{ref}}$ of CAB. If, instead, we use the $C_{\text{ref}}$ of MHD, $\sigma_{ap} = 10.1 \pm 6.1$, which may be considered as an upper estimate. Table 2 summarizes the light-scattering and absorption coefficients and single-scattering albedos observed in this study and in selected other studies using comparable instruments (Man and Shih, 2001; Xu et al., 2002; Yan et al., 2008). On average, the $\sigma_{ap}$ was lower than that measured at Lin’an regional background station in the rural area of the Yangtze River Delta region (Xu et al., 2002). Compared to the value measured at Hok Tsui 15 years ago, $\sigma_{ap}$ was lower than that observed in Hok Tsui, from November 1997 to February 1999 (Man and Shih, 2001). Being the most significant light-absorbing constituent of aerosols, a similar decrease of BC concentration was also found. Table 3 presents the mean BC mass concentrations reported in other comparable studies. The overall average of BC mass concentrations in this study was $1.4 \pm 1.1 \, \mu g \, m^{-3}$ (Table 1), which was lower than the values observed at the same site in 2004–2005 (with a mean of $2.4 \, \mu g \, m^{-3}$ using a AE-42 Aethalometer; Cheng et al., 2006a). A decreasing trend of BC concentration was found at the Panyu station in the PRD region with a decreasing rate of approximately $1 \, \mu g \, m^{-3}$ per year from 2004 to 2007 (Wu et al., 2009). Compared to the other rural sites in South China, BC levels in Hok Tsui station were lower than the concentra-
Table 2. Summarization of aerosol light-scattering coefficients, absorption coefficients and single-scattering albedo observed in this study and reported in other studies.

<table>
<thead>
<tr>
<th>Site</th>
<th>Period</th>
<th>$\sigma_{ap}$</th>
<th>$\sigma_{sp}$</th>
<th>SSA</th>
<th>Instrumentation</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hok Tsui, Hong Kong (rural, coastal)</td>
<td>Feb 2012–Feb 2015</td>
<td>8.3 ± 6.1</td>
<td>151 ± 100</td>
<td>0.93 ± 0.05</td>
<td>AE31, Magee Scientific Nephelometer, TSI, Inc.</td>
<td>This work</td>
</tr>
<tr>
<td>Cape D’Agular (Hok Tsui), Hong Kong (rural, coastal)</td>
<td>Nov 1997–Feb 1998</td>
<td>25.72</td>
<td>64.77</td>
<td></td>
<td>PSAP, Radiance Research Nephelometer, Radiance Research</td>
<td>Man and Shih (2001)</td>
</tr>
<tr>
<td>Cape D’Agular (Hok Tsui), Hong Kong (rural, coastal)</td>
<td>Mar–Apr 1998</td>
<td>15.79</td>
<td>38.65</td>
<td></td>
<td>PSAP, Radiance Research Nephelometer, Radiance Research</td>
<td>Man and Shih (2001)</td>
</tr>
<tr>
<td>Xinken, PRD, China (non-urban, regionally polluted)</td>
<td>Oct–Nov 2004</td>
<td>70 ± 42</td>
<td>333 ± 137</td>
<td>0.83 ± 0.05</td>
<td>MAAP, Thermo, Inc. Nephelometer, TSI, Inc.</td>
<td>Cheng et al. (2008)</td>
</tr>
<tr>
<td>Shangdianzi, China (rural)</td>
<td>Sep 2003–Jan 2005</td>
<td>17.54 ± 13.44</td>
<td>174.6 ± 189.1</td>
<td>0.88 ± 0.05</td>
<td>AE31, Magee Scientific Nephelometer, EcoTech</td>
<td>Yan et al., (2008)</td>
</tr>
<tr>
<td>Lin’an, China (rural)</td>
<td>Nov 1999</td>
<td>23 ± 14</td>
<td>353 ± 202</td>
<td>0.93 ± 0.04</td>
<td>PSAP, Radiance Research Nephelometer, Radiance Research</td>
<td>Xu et al. (2002)</td>
</tr>
<tr>
<td>Granada, Spain (urban)</td>
<td>Dec 2005–Nov 2007</td>
<td>21 ± 10</td>
<td>60 ± 30</td>
<td>0.68 ± 0.07</td>
<td>MAAP, Thermo, Inc. Nephelometer, TSI, Inc.</td>
<td>Lyamani et al., (2010)</td>
</tr>
<tr>
<td>Alomar station, Norway (background, coastal)</td>
<td>Jun–Aug 2008</td>
<td>0.40 ± 0.27</td>
<td>5.41 ± 3.55</td>
<td>0.91 ± 0.05</td>
<td>PSAP, Radiance Research Nephelometer, TSI, Inc.</td>
<td>Mogo et al. (2012)</td>
</tr>
</tbody>
</table>

Table 3. Comparison of mean concentration of BC with other studies.

<table>
<thead>
<tr>
<th>Site</th>
<th>Environment</th>
<th>Period</th>
<th>Inlet</th>
<th>BC ($\mu$g m$^{-3}$)</th>
<th>Instrumentation</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hok Tsui, Hong Kong</td>
<td>Rural, coastal</td>
<td>Feb 2012–Feb 2015</td>
<td>PM$_{2.5}$</td>
<td>1.4 ± 1.1</td>
<td>AE31, Magee Scientific</td>
<td>This work</td>
</tr>
<tr>
<td>Cape D’Agular (Hok Tsui), Hong Kong</td>
<td>Rural, coastal</td>
<td>Jun 2004–May 2005</td>
<td>PM$_{2.5}$</td>
<td>2.4 ± 1.8</td>
<td>AE42, Magee Scientific</td>
<td>Cheng et al. (2006a)</td>
</tr>
<tr>
<td>Yongxing Island, China</td>
<td>Oceanic rural, (South China Sea)</td>
<td>May–Jun 2008</td>
<td>PM$_{2.5}$</td>
<td>0.54 (rainy season)</td>
<td>AE42, Magee Scientific</td>
<td>Yu et al. (2013)</td>
</tr>
<tr>
<td>Maofengshan, China</td>
<td>Rural, PRD</td>
<td>May–Jun 2008</td>
<td>PM$_{10}$</td>
<td>2.62 (rainy season)</td>
<td>AE42, Magee Scientific</td>
<td>Yu et al. (2013)</td>
</tr>
<tr>
<td>Toulon, France</td>
<td>Semi-urban, coastal</td>
<td>Jun 2005–Oct 2006</td>
<td>PM$_{2.5}$</td>
<td>0.95 (winter)</td>
<td>AE31, Magee Scientific</td>
<td>Saha and Despiau (2009)</td>
</tr>
<tr>
<td>Hyytiälä, Finland</td>
<td>Boreal forest</td>
<td>Dec 2004–Dec 2008</td>
<td>PM$_{2.5}$</td>
<td>0.32 ± 0.34</td>
<td>AE31, Magee Scientific</td>
<td>Hyvarinen et al. (2011)</td>
</tr>
<tr>
<td>Voerde-Spellen, Germany</td>
<td>Rural</td>
<td>Sep–Oct 1997</td>
<td>PM$_{2.5}$</td>
<td>0.8 ± 0.3</td>
<td>AE-10 IM, G1V</td>
<td>Kuhlbusch et al. (2001)</td>
</tr>
<tr>
<td>Preila, Lithuania</td>
<td>Rural, coastal</td>
<td>Mar–Apr 2002</td>
<td>PM$_{2.5}$</td>
<td>0.84</td>
<td>AE40, Magee Scientific</td>
<td>Andriejauskiené (2008)</td>
</tr>
</tbody>
</table>

The seasonal cycles of target parameters were analyzed based on hourly-averaged data classified as four seasons: winter (December–February), spring (March–May), summer (June–August), and autumn (September–November). Seasonal averaged values of selected parameters are listed in
Table 4. Summary of seasonal average value of target pollutants.

<table>
<thead>
<tr>
<th></th>
<th>Winter</th>
<th>Spring</th>
<th>Summer</th>
<th>Autumn</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AVG ± SD MED</td>
<td>AVG ± SD MED</td>
<td>AVG ± SD MED</td>
<td>AVG ± SD MED</td>
</tr>
<tr>
<td>σ_{ap, 550 nm} (Mm^{-1})</td>
<td>10.9 ± 7.1 9.6</td>
<td>7.5 ± 4.8 6.4</td>
<td>5.5 ± 5.8 3.8</td>
<td>7.4 ± 4.5 6.3</td>
</tr>
<tr>
<td>σ_{sp, 550 nm} (Mm^{-1})</td>
<td>193 ± 102 176</td>
<td>148 ± 89 133</td>
<td>64 ± 62 49</td>
<td>140 ± 82 130</td>
</tr>
<tr>
<td>SSA (550 nm)</td>
<td>0.94 ± 0.03 0.95</td>
<td>0.93 ± 0.05 0.95</td>
<td>0.90 ± 0.06 0.92</td>
<td>0.94 ± 0.03 0.95</td>
</tr>
<tr>
<td>BC (µg m^{-3})</td>
<td>2.0 ± 1.2 1.8</td>
<td>1.3 ± 1.0 1.2</td>
<td>0.9 ± 1.1 0.6</td>
<td>1.5 ± 0.9 1.4</td>
</tr>
<tr>
<td>CO (ppbv)</td>
<td>459 ± 186 423</td>
<td>275 ± 154 262</td>
<td>117 ± 84 91</td>
<td>270 ± 134 252</td>
</tr>
<tr>
<td>AAE</td>
<td>1.1 ± 0.2 1.1</td>
<td>1.0 ± 0.3 1.0</td>
<td>0.7 ± 0.4 0.7</td>
<td>0.9 ± 0.3 0.8</td>
</tr>
<tr>
<td>SAE</td>
<td>1.3 ± 0.3 1.3</td>
<td>1.2 ± 0.5 1.2</td>
<td>1.4 ± 0.6 1.5</td>
<td>1.7 ± 0.4 1.8</td>
</tr>
<tr>
<td>N_{total} (cm^{-3})</td>
<td>7690 ± 3821 6768</td>
<td>8620 ± 4868 7540</td>
<td>7003 ± 4460 5943</td>
<td>7808 ± 3970 7077</td>
</tr>
<tr>
<td>GMD (nm)</td>
<td>72 ± 17 70</td>
<td>68 ± 15 66</td>
<td>58 ± 17 54</td>
<td>69 ± 17 68</td>
</tr>
<tr>
<td>PM_{1} (µg m^{-3})</td>
<td>27 ± 19 22</td>
<td>25 ± 19 21</td>
<td>13 ± 20 7</td>
<td>22 ± 14 19</td>
</tr>
</tbody>
</table>

Figure 2. (a) Seasonal cycle of scattering coefficient, σ_{ap}; absorption coefficient, σ_{sp}; single-scattering albedo; SSA; temperature; and precipitation; where bold solid lines represent median values, diamonds show the monthly averages, and thin solid lines are percentiles of 75 and 25%. (b) Seasonal cycles of BC, CO, SO_{2}, and PM_{2.5} concentrations, where blue solid lines represent median values, diamonds show the monthly averages, the boxes are 25th and 75th percentiles, and the thin bars represent 10th and 90th percentiles.

The highest σ_{ap} and σ_{sp} values were observed in winter (10.9 ± 7.1 and 193.5 ± 102 Mm^{-1}, respectively), which were more than twice that of summer. A similar pattern was observed in a previous study in Hong Kong in 1997–1999 (Man and Shih, 2001). Compared with other rural/background sites, the average SSA_{550 nm} at Hok Tsui was 0.94 ± 0.03 during autumn, which was higher than that measured at Xinken, PRD, China, in the same season (0.83 ± 0.05), while this value was 0.90 ± 0.06 in summer, which was slightly lower than that observed at a coastal station in Norway; also in summer (0.91 ± 0.05, Mogo et al., 2012).

Figure 2 presents the monthly variation of measured optical properties and meteorological parameters. A clear seasonal cycle of aerosol optical properties is shown with σ_{ap} and σ_{sp}, having peaked in January and reaching the lowest level in July. While it was lighter in winter, the aerosol was the darkest in summer, especially in August, with a seasonal mean SSA of 0.87. Averaged seasonal values of 1-SSA in 36 wind sectors are presented in Fig. 3a. These figures show the disparity of SSA from different wind directions. Overall, air plume coming from the southwest to the north (225–360°) had higher 1-SSA, i.e., lower SSA, than that from the east (45–135°). Ding et al. (2013) reported that the contribu-
Figure 3. (a) Seasonal mean value of 1-SSA in 36 wind sectors during the whole period. (b) Map of averaged 7-day retroplume when SSA is below 0.9 compared with (c) averaged 7-day retroplume during the whole period.

Figure 4. Averaged diurnal variations of (a) $\sigma_{sp}$, (b) $\sigma_{ap}$, (c) BC, (d) CO, (e) SAE, (f) AAE, (g) SSA, and (h) PM$_1$ in four seasons.

tion of anthropogenic emissions from Guangdong and Hong Kong was the highest in August, which means more freshly emitted urban aerosols were brought to the monitoring station with lower SSA in this month (Cheng et al., 2008). The main synoptic process contributing to this kind of sub-regional transport is tropical cyclones. Ding et al. (2004) explained the mechanism of how these tropical cyclones influence the development of sea–land breeze and also explained further about sub-regional and urban air mass accumulation in the South China. Zhang et al. (2013) found an important
influence of tropical cyclones in ozone and haze pollution in this region in summer, based on an analysis of 13-year data.

Another possible reason for the relatively low SSA in August is that the air mass came mainly from the southwest of the site (Fig. 1), a main waterway for ocean-going vessels in Hong Kong (Yau et al., 2012). These vessels emitted considerable amount of light-absorbing carbon from diesel engines during combustion. Similar pattern was also observed in the seasonal diagrams of BC, SO$_2$, PM$_{2.5}$, and CO, which are typical components of ship exhaust (Fig. 2, Hong Kong Air Pollutant Emission Inventory for 2013 from Hong Kong Environmental Protection Department: http://www.epd.gov.hk/epd/english/environment intimk/air/data/emission_inve.html).

Figure 3b demonstrates the averaged 7-day retroplume of the times when SSA was lower than 0.9. Compared with the overall averaged 7-day retroplume during the whole measurement period (Fig. 3c), darker aerosols were mostly from two main types of regions in the vicinity: one was the nearby continental area with fresh polluted air masses from urban Hong Kong and neighboring PRD cities; another branch was from the ocean side. Fresh emission of passing ships or fast transport from southern Asia could lead to higher proportion of BC in the air plumes and thus caused lower SSA.

Figure 4 shows the diurnal cycles of $\sigma_{sp}$, $\sigma_{sp}$, BC, CO, and PM$_{1}$ for four seasons. There was an increase in $\sigma_{ap}$ after sunrise with a peak occurring before noontime. It might be associated with a combined effect of increased human activities and turbulence mixing in the boundary layer in the morning. This pattern was more significant in summer, although the pollution level was relatively low. This phenomenon supports the explanation of turbulent mixing from a middle or upper planetary boundary layer (PBL) because of a stronger vertical mixing in summer. The PM$_{1}$ also showed a daytime maximum concentration but with the peak in the afternoon (Fig. 4c). For $\sigma_{sp}$, morning peaks were not as significant as $\sigma_{ap}$. The decrease in $\sigma_{sp}$ in the early afternoon might be caused by a further development of PBL or mixing layer, in which the air pollutants experienced a substantial dilution, resulting in lower concentrations of pollutants at the ground surface. Diurnal variations and fluctuations of CO mixing ratios show a similar pattern with $\sigma_{sp}$ but a relatively smaller variability.

3.1.3 Optical properties and their relationships with particle size

Wavelength dependencies of aerosol light scattering and absorption are closely related to aerosol size and dominating aerosol types. To find out the difference of light absorbing materials, Fig. 5a displays the relationship of SSA with AAE, color-coded with a BC mass fraction of submicron particles (PM$_{1}$ was calculated from the particle number size distributions measured with the UFPM.) It shows that aerosols with high SSA had a lower BC fraction and that AAE varied greatly in the lower value region, indicating the dominance of scattering particles. Such air masses were likely of longer transport time and the BC aerosols had mixed well with light-scattering aerosols during transport. Contrarily, the low SSA values mostly occurred when AAE were closely distributed around 1.0 and in these cases BC took up a higher proportion (red dots in Fig. 5a), showing freshly emitted BC plumes.

Figure 5b and c demonstrate the relationships between particle size and their scattering Ångström exponents as well as their darkness. It can be observed that SAE generally increased with decreasing SSA. Dark aerosols with low SSA were mostly small in size with low GMD but a high BC fraction. These small dark aerosols had higher SAE (1.5 to 2.0). The wide range of SAE was possibly due to the mixed control by continental aerosols and large sea-salt aerosols.

Figure 6 shows the scatter plot of $\sigma_{sp}$ calculated using Eq. (2) versus the measured $\sigma_{sp}$. The slope of $\sigma_{sp, submicron}/\sigma_{sp, obs}$ was 0.86, indicating that submicron particles were the major light-scattering components in the air masses arriving at the Hok Tsui station. For most of the time in the study period, the simulated $\sigma_{sp}$ was lower than the observed $\sigma_{sp}$. This is probably because the particle size distribution data from the UFPM only used the scanned submicron particles with mobility diameter from 20 to 800 nm (see Fig. 6b and c) in the calculation, but the nephelometer, equipped with a TSP inlet, measured light-scattering coefficients from all particles with a wider-sized range. The relatively limited number of particle size bins in the UFPM probably also leads to uncertainties for the calculation of $\sigma_{sp}$.

Hence, this result can only provide a rough image of the relationships between particle light scattering and their size distribution at the Hok Tsui station. It can be observed that particles with $D_{sp}$ less than 200 nm contributed the largest fraction of the total number of submicron particles but very little to the total scattering, whereas the small amount of larger particles ($D_{sp}$: 200–800 nm) contributed the most to the total light scattering.

The scatter plot (Fig. 6a) also shows that there were clusters of data where the modeled and measured $\sigma_{sp}$ fit close to the 1 : 1 line and clusters where the measured $\sigma_{sp}$ was clearly larger than those modeled on. After computing the averaged retroplume of these clusters, it was found that the former data cluster is mostly associated with polluted continental air and the latter with stronger winds and sea-salt particles (figures were not shown).

3.2 Source identification

Figure 7 shows the spatial distribution of BC / CO emission ratios in East China and the nearby regions calculated using the MIX emission inventory. It can be seen that the BC / CO emission ratio was higher in Shanxi Province (higher than 25 ng m$^{-3}$ ppbv$^{-1}$), Taiwan (approximately 20 ng m$^{-3}$ ppbv$^{-1}$), and the regions along the coastline of East China. As reported by previous studies, the BC / CO emission ratio from industrial coal burning ranges
from 1.9 to 20 and it was 5.6–13.3 ng m$^{-3}$ ppbv$^{-1}$ from open biomass burning (Wang et al., 2011; Zhang et al., 2009). For diesel vehicles, the BC / CO emission ratio was 14–39 and it was 15.6 ng m$^{-3}$ ppbv$^{-1}$ for ship emission calculated from a previous study in southern Asia (Dickerson et al., 2002).

A strong correlation between BC and CO and a high slope of $27 \times 10^{-3}$ g BC g CO$^{-1}$ were found from a previous study using C-130 aircraft flew over the Arabian Sea and northern
Indian Ocean (Dickerson et al., 2002; Mayol-Bracero et al., 2002).

In this study, ΔBC / ΔCO and SO₂ / BC ratios were investigated to study the source characteristics and the freshness of the fuel combustion sources. ΔBC / ΔCO (net growth of BC and CO; total concentration minus regional baseline. (Spackman et al., 2008) and SO₂ / BC were calculated with 1 h time resolution. The baseline of BC and CO was determined as 1.25th percentiles of data in each month (Pan et al., 2011). Monthly variation of ΔBC / ΔCO is displayed in Fig. 8 together with SO₂ / BC to demonstrate the fuel burning emission profile since SO₂ is a co-emitted species of fossil fuel combustion (Bond et al., 2013). Reference emission ratios of BC / CO and SO₂ / BC from previous studies (Bond et al., 2013; Li et al., 2015) are also plotted in Fig. 8.

In Hong Kong, major SO₂ emission was from navigation and public electricity generation, contributing 50 and 47% to total SO₂ emission (Emission Inventory 2013, HKEPD, http://www.epd.gov.hk/epd/english/environmentinhk/air/data/emission_inve.html). However, these two sources only took up 19 and 6% of CO emission and the largest contributor of CO reported in the emission inventory was road transport (59%). As shown in Fig. 8, ΔBC / ΔCO and SO₂ / BC ratios presented similar monthly variation patterns. The monthly mean ΔBC / ΔCO ranged from 1.5 to 20 µg m⁻³ ppbv⁻¹ during the whole study period. The highest values occurred in summer months for both ΔBC / ΔCO and SO₂ / BC and the ratios were relatively lower in winter. Since SO₂ has a short lifetime, which can easily deposit and transform into secondary aerosols, the synchronous elevation of ΔBC / ΔCO and SO₂ / BC in summer indicates that freshly emitted anthropogenic pollutants might be more easily influenced by the air masses in

![Figure 9. Scatter plot of hourly BC and CO in four wind sectors.](image)

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this coastal area. The decrease of ΔBC / ΔCO and SO₂ / BC in winter provided the evidence that this area was under the influence of contaminated air masses from a longer distance. Figure 9 displays the scatter plot of BC vs. CO in four wind sectors, giving an image of the freshness of polluted air masses and the intensities of combustion emissions from different directions. For wind directions from 180 to 360°, the data points show a good positive correlation, suggesting that most of the BC and CO emission sources in these areas were closer to the measurement site. Data points in the 0–180° wind direction sector were much more scattered with a lower slope of It should be ‘delta BC / delta CO’ of air pollutants coming from the northeast to the southeast. Through the transport of air plume, diffusion and deposition of air pollutants would decrease their concentrations that arrive to the receptor, and therefore lower the slope of ΔBC / ΔCO and their correlation coefficients.

To investigate the transport pattern of air masses that arrived at the site during the study period, Fig. 10 shows averaged 7-day retroplume for four seasons. As presented in Ding et al. (2013), it shows a distinguished different transport pattern under the influence of Asian monsoons. During summer, the majority of air masses came from the south and nearby PRD cities. Due to the dominance of relatively clean marine air, emission from passing ships or local activities in adjacent regions could make visible effects on the temporal variations of air pollutants. The source distribution was more complicated during winter. Driven by the winter monsoon, cold and dry air masses that were transported along the coastline of East China and from central China took up a higher proportion in winter months (Ding et al., 2013). There were also air masses passing through Taiwan and the East China Sea during the cold season (Fig. 10a).

Since BC is the most significant light-absorbing constituent of aerosols, to evaluate the potential source contribution of light absorbing particles on regional optical properties, averaged PSC maps of BC for different transport times and seasons were calculated using the method described in Sect. 2.6 and illustrated in Fig. 11. However, here we only calculated the PSC from emission over land because the available emission inventory from the MIX database is mainly focused on land areas. As shown in Figs. 11a and 10b, BC concentrations were influenced by the transport from nearby cities within a short time, especially Shenzhen and urban Hong Kong. Long-range transport of BC from the South and East China also played an important contribution. It was also shown that BC coming from the continental area through longer distances took up a higher proportion of the pollutant level in winter (Fig. 11d) than that of the summer (Fig. 11c). During summer, local emission was the biggest BC contributor.

Figures 12a–11c illustrate the average levels of σ_{sp}, σ_{sp} and PM₁ and the corresponding frequency of occurrence for air masses passing through different regions calculated using Eq. (3). Together with the shipping routes density map.
(Fig. 12d), it can be observed that the high levels of \( \sigma_{ap} \) and \( \sigma_{sp} \) were closely associated with the congested shipping lanes in the maritime space nearby Hong Kong. The high \( \sigma_{ap} \) and \( \sigma_{sp} \) were especially visible in the northeast due to the prevailing northeasterly wind from autumn to spring, transporting ship exhausts mainly through the Taiwan Strait. The belt-like zone with higher \( \sigma_{ap} \) and \( \sigma_{sp} \) was likely to reveal ship emission. As shown in Fig. 12d, there were dense shipping routes between Hong Kong and Singapore traveling through the Xisha Islands in the South China Sea where the routes were similar to the high \( \sigma_{sp} \) area in the south. During summer, Hong Kong was influenced by the southerly and southwesterly wind, bringing clean marine air to this region for most of time and leading to lower pollutant levels (Wang et al., 2009; Ding et al., 2013). Figure 12 indicates that Hong Kong could be affected by the passing vessels in the South China Sea due to controlling wind direction, driven by the summer monsoon.

3.3 Analyses of selected episodes

Figure 13 demonstrates the aerosol optical properties and BC-CO correlations associated with air masses from different source regions during selected episodes. The major source regions were Guangdong and urban Hong Kong (GH), ship emission (SP), North China (NC), and the aged continental area (AGC). The selection of the episodes was done by combining the footprints using LPDM and the variation trend of aerosol optical properties and \( \text{PM}_{2.5} \). The air pollution plumes coming from Guangdong and urban Hong Kong had the highest BC and CO concentrations (Fig. 13a), indicating a higher level of emission intensity and stronger light-extinction ability of aerosols from these regions. The slope of BC vs. CO was highest from ship emission \((0.012 \, \mu g \, m^{-3} \, ppbv^{-1})\), with high correlation \((r^2 = 0.84)\), showing that the ship emission source was close to the measurement station and its exhausts could largely affect the pollution level.

Figure 13b displays that Ångström exponents of scattering, from Guangdong and Hong Kong, were relatively high, compared to that from the aged continental area and North China, proving the dominance of smaller particles of emissions from PRD cities and passing ships. BC-containing particles transported from North and East China went through longer coating and deposition processes, which enlarged their size but decreased their concentrations when arriving to the measurement site. This can further explain the lower SSA in summer months.

Overall, the analyses suggest that aerosols from different source regions could cause great variances on regional aerosol optical properties. Thus, more ground observations of aerosol optical properties are needed to fully understand
the characteristics of different types of atmospheric aerosols and provide reference datasets for further investigation of aerosol radiative forcing and climatic effects.

4 Conclusions

Based on aerosol optical properties, relevant species and aerosol size measured at Hok Tsui station in Hong Kong, we studied the temporal variations and investigated the potential sources by using correlation analysis and Lagrangian dispersion modeling. Overall, the absorption coefficients at the site in the South China coastal region were lower than most of other rural sites in eastern China. Scattering coefficients observed in this study were almost twice the values monitored at the same station in 1998, yet BC concentrations decreased over 50% compared with the measurements in 2004. The darkest aerosols were smaller in particle size but showed strong scattering wavelength dependencies, indicating possible sources from fresh emissions close to the measurement site. Particles with $D_p$ of 200–800 nm were less in number, yet contributed the most to the light-scattering coefficients among submicron particles.

A remarkable correlation was found for BC and CO concentrations during episodes. $\Delta$BC/ΔCO range from 1.5 to 20 µg m$^{-3}$ ppb$^{-1}$ during whole period. Both $\Delta$BC/ΔCO and SO$_2$/BC peaked in summer months and were relatively low in spring and autumn. In summer, the site was affected by nearby combustion sources, while in spring and autumn the observed air masses were more under the influence of well-mixed air masses from long-range transport. Multi-year backward LPDM and PSC analysis together with case studies provided detailed information of the transport of air masses and their impacts on aerosol optical properties. For summer months, air masses moved southward over Shenzhen, and urban Hong Kong brought air pollutants mainly from residential and transportation to the measurement site, showing strong light absorbing ability; ship exhausts were introduced into the southerly marine air with higher speed, showing a strong positive correlation between BC and CO. These fresh emission sources led to low SSA during the summer. For winter and autumn months, the air plume arriving at Hok Tsui station was a mixture of multi-source aerosol including air masses from South China, East China coastal regions, and aged aerosol transported over the Pacific Ocean and Taiwan, causing stronger abilities of light extinction and larger
Figure 12. Map of average property retroplume for (a) $\sigma_{ap}$, (b) $\sigma_{sp}$, and (c) PM$_1$ (the non-colored areas were where the total retroplume was smaller than $10^{-12}$ mass m$^{-3}$ h$^{-1}$ (i.e., air plumes barely passed through these regions). Due to the different time period of valid data from UFP, the non-colored areas were slightly different in (c). (d) Density map showing the ship routes near Hong Kong during 2013 and 2014.

Figure 13. (a) Scatter plots of BC and CO. (b) $\sigma_{ap}$ and SAE from different source regions during episodes GH: Guangdong and Hong Kong; SP: Ship; NC: North China; and the AGC: aged continental area.
variability of aerosol optical properties as well as pollutant concentrations.

5 Data availability

The data can be accessed upon contact with the corresponding authors.

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

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References


