

Strong Primary Contribution to Brown Carbon Light Absorption in Tibet and Urban Areas:

Insights based on in situ measurements

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Abstract

To investigate optical properties, sources, and radiative effects of brown carbon (BrC), we conducted synchronous field campaigns in the Qinghai–Tibet Plateau (Yangbajing) and urban Guangzhou in July 2022, using multi-wavelength Aethalometer (AE33) and aerosol mass spectrometer (AMS) measurements. Total aerosol and BrC light absorption coefficients at 370 nm (Abs_{total} : 1.6 ± 1.6 $M\ m^{-1}$; BrC: 0.2 ± 0.3 $M\ m^{-1}$) in Tibet were an order of magnitude lower than Guangzhou, attributed to extremely low aerosol/organic aerosol (OA) mass concentrations. However, BrC fractions in Abs_{total} (15 % vs. 21 % at 370 nm) correlated with primary OA (POA) ratios, highlighting anthropogenic emission impacts even in this clean background. Diurnal variations (morning/evening peaks) of source-specific BrC absorption were regulated by local emissions (e.g., biomass burning, traffic emission) and regional secondary formation. Source apportionment revealed primary sources (biomass burning OA (BBOA), hydrocarbon-like OA (HOA)) dominated BrC absorption (> 75 %). The mass absorption cross-section (MAC) of HOA ($2.08\ m^2\ g^{-1}$ in Tibet; $2.57\ m^2\ g^{-1}$ in Guangzhou) was comparable to that of BBOA (1.11 – 2.54 in Tibet; $1.91\ m^2\ g^{-1}$ in Guangzhou), indicating the high light absorption capacity of BrC from fossil fuel. Integrated "simple forcing efficiency" (370–660 nm) showed primary emissions contributed > 98 % of total radiative forcing at both sites. This study advances understanding of BrC dynamics and sources in diverse environments, underscores primary sources' critical role in BrC absorption, and emphasizes the need for source-specific OA optical parameterization.

Keywords:

Brown carbon; Source apportionment; Multiple linear regression; Optical properties; Radiative forcing

1. Introduction

Light absorbing components of atmospheric aerosols comprise black carbon (BC) and light absorbing organic aerosols, known as brown carbon (BrC). The BrC exhibits significant absorption in the near-ultraviolet (300–400 nm) and visible ranges (Andreae and Gelencsér, 2006; Kirchstetter and Thatcher, 2012; Laskin et al., 2015), but it remains largely overlooked in most climate models (Chung et al., 2012). Feng et al. (2013) found that accounting for the strong absorption of BrC in model could shift the global mean direct radiative forcing of OA at the top of the atmosphere from a cooling of $-0.08\ Wm^{-2}$ to a warming of $+0.025\ Wm^{-2}$, emphasizing the significant role of BrC in global and regional direct radiative forcing of carbonaceous aerosols. Wang et al. (2025) revealed that dark BrC, emitted by wildfires and agricultural burning, strongly absorbs solar radiation, thereby generating a radiative effect of $+0.02$ to $+0.68\ Wm^{-2}$, which contributed comparable warming as BC. Current global model simulation studies have demonstrated that the impact of BrC can contribute 12–50 % of the total positive radiative forcing from light-absorbing atmospheric aerosols and can be regionally different, which emphasizes the spatially dynamic variation of BrC and its important role in atmospheric warming (Li et al., 2025; Brown et al., 2018; Feng et al., 2013; Wang et al., 2018; Xu et al., 2024),.

In general, the source of atmospheric BrC can be categorized by direct emission (primary BrC) and secondary generation (secondary BrC) (Laskin et al., 2015). Over the past decade, primary BrC was found to be dominated by biomass burning, and the absorption capacity of BrC from fossil fuel is often neglected (Saleh, 2020). However, more and more recent studies have shown that fossil fuels (vehicle emissions and coal combustions) are also important contributors to the BrC, which have been previously underestimated (Du et al., 2014; Kasthuriarachchi et al., 2020a; Xie et al., 2017; Huang et al., 2022; Tang et al., 2020). Wang et al. (2022b) conducted a comprehensive review of the relationship between the source and light absorption characteristics of BrC based on the measurement results in China, positing that the emission and light absorption capacity of BrC from fossil fuel combustion might be comparable to or even exceed those induced by biomass burning. This finding raises the question of how much primary fossil and non-fossil sources contribute to ambient BrC. Regarding secondary BrC, its formation involves complex gaseous, particulate, and liquid-phase reactions from diverse precursors (Laskin et al., 2015). E.g., the nitro compounds formed through NO_3 oxidation or OH oxidation under high NO_x chemistry can lead to a significant enhancement in light absorption within the ultraviolet–visible (UV–Vis) range (Li et al., 2020; Jiang et al., 2019). Aqueous reactions of carbonyl groups with reduced nitrogenous organic compounds, such as organic amines and ammonium, are also found to be important BrC sources (Powelson et al., 2014; Tang et al., 2022).

The complex sources and multi-forming pathways of BrC make its global simulation a great challenge, e.g., mass absorption cross-section (MAC), which serves as a crucial optical parameter in simulating the BrC light absorption and further its radiative forcing, is still not clear due to the impact of multiple factors. E.g., various studies have demonstrated that the BrC light absorption properties are susceptible to the sources (Tang et al., 2020), photochemical aging (Yu et al., 2016; Wong et al., 2017; Lee et al., 2014; Zhao et al., 2015), humidity (Kasthuriarachchi et al., 2020b), acidity of aerosols (Mo et al., 2017; Phillips et al., 2017), and structure of chromophores (Laskin et al., 2014; Hems and Abbatt, 2018). In general, the MAC of secondary BrC was found to be generally lower than primary sources in ambient BrC (Qin et al., 2018; Zhang et al., 2022b). Smog chamber experiments also show that the MAC of aged coal combustion emission ($0.14 \pm 0.08 \text{ m}^2 \text{ g}^{-1}$) was much lower than that of primary emissions ($0.84 \pm 0.54 \text{ m}^2 \text{ g}^{-1}$) (Ni et al., 2021). If the impact of the atmospheric aging on MAC is not considered in the model, the simulated BrC light absorption may be overestimated by 45 % to 128 % (Li et al., 2025). In addition, Li et al. (2025) pointed out that the uncertainty in the effective MAC of primary BrN (i.e., absorptive nitrogenous component of BrC) from anthropogenic and biomass burning has the greatest impact (reaching ± 76 % uncertainty) on the simulated BrN light absorption, highlighting the key role of source-specific MAC in the global simulation of BrC light absorption. Further clarification of the MAC of BrC from different sources in the ambient air is imperative, which can greatly help to improve the understanding of BrC light absorption and the model simulation. However, the total contribution to ambient BrC from SOA and POA and their source-specific MAC values is still ambiguous due to varied regions and circumstances, which warrants further study.

Due to the complexity and diversity of BrC, in conjunction with black carbon (BC), which share similar combustion sources and complex mixture states (Bond and Bergstrom, 2006; Cappa et al., 2012; Cappa et al., 2019), it remains challenging to measure the source-specific BrC light absorption in the

76 ambient air directly. The field measurement on ambient BrC begins with filter-based offline methods
77 using solvent (water or organic solvent) extraction (Bond and Bergstrom, 2006). One of the limits of the
78 offline technique is the low time resolution (usually 12–24 hours), which cannot reflect the dynamic
79 variation of BrC during a day. As measurement techniques have developed, online methods for aerosol
80 light absorption at different wavelengths have become available (Lack et al., 2014). The most widely
81 used instruments include multi-wavelength Aethalometers (AE31/AE33) (Drinovec et al., 2015) and
82 three-wavelength multi-pass Photo Acoustic Spectroscopy (PAS) (Lack et al., 2012). The online
83 measurements can provide dynamic variations in BrC light absorption at high time resolution (1 minute).
84 Together with source apportionment techniques, source contribution to BrC at high time resolution can
85 be obtained, which can greatly aid in understanding the dynamic characteristics of ambient BrC and its
86 sources.

87 Currently, most of the field studies focusing on ambient BrC and its sources are conducted in urban
88 areas (Qin et al., 2018; Sun et al., 2021; Wang et al., 2019b; Zhong et al., 2023), while the BrC
89 contribution in regional background areas is still limited. The Qinghai–Tibet Plateau (QTP) region,
90 spanning approximately 2.5 million square kilometers, is acknowledged as the world's highest plateau
91 (Yao et al., 2012). The atmosphere in QTP can substantially influence the climate in the world. E.g., the
92 light-absorbing carbonaceous aerosols have been identified as one of the main factors causing the
93 accelerated glacier retreat across the QTP (Usha et al., 2022; Kang et al., 2019; Chelluboyina et al., 2024).
94 Compared to the southern and northern regions of the QTP, anthropogenic BC is the dominant driver of
95 glaciers melt in the central QTP, whose radiative forcing can be up to 17 times greater than that in the
96 southern QTP (Li et al., 2017; Ming et al., 2013). Moreover, a portion of the glaciers in central QTP are
97 situated in the headwaters of the Yangtze River, thus, the melting directly impacts the livelihoods of
98 millions of people downstream. The melt sensitivity of these glaciers to light-absorbing impurities (such
99 as OA, BC and mineral dust) is not only higher than that of their southern counterparts but also carries
100 greater regional ecological and social significance (Li et al., 2017). Although BrC is an important light-
101 absorbing OA, limited studies on its variation and sources have been conducted in the central QTP, and
102 most have focused on the edge of the southern region (Wang et al., 2019a; Zhang et al., 2021; Tian et al.,
103 2023). We found that cross-border transport of biomass burning from South Asia was responsible for a
104 significantly higher BrC light absorption contribution in the southern QTP than in the central and
105 northeastern regions. However, OA in the central QTP not only originated from long-range transport (Xu
106 et al., 2018), but anthropogenic activities also have a significant impact on the OA sources, such as local
107 biomass burning and fossil fuels (Xiang et al., 2024). Field measurements in the central QTP can improve
108 our understanding of how human activities in remote regions impact light-absorbing aerosols and,
109 consequently, glacial melting and radiative forcing. In central QTP, only one source apportionment of the
110 light absorption of offline water-soluble BrC was reported (Zhu et al., 2024), which showed a large
111 contribution of biomass burning (29 %), fossil fuel combustion (17 %) and secondary contribution (54 %)
112 to the total BrC light absorption. Further clarification is still needed. In addition, although the online BrC
113 data have been reported in the Tibet region, the dynamic variation of BrC and its source contribution is
114 seldom shown (Zhang et al., 2021; Wang et al., 2024; Chen et al., 2024; Wang et al., 2019a; Tian et al.,
115 2023; Zhu et al., 2021; Zhu et al., 2017). The investigation of the dynamic variation of BrC, e.g., diurnal

variation, can greatly promote the understanding of BrC fate in the ambient air, which shall be further studied.

In this study, the real-time measurement of OA and light absorption of aerosols was carried out in the background site of central QTP. For comparison, a concurrent campaign was also carried out during the same period in a typical megacity of China, Guangzhou. The comparison results from distinct two environments at the same time can help better understand the light absorption capacity of BrC in QTP. In both campaigns, an online multi-wavelength Aethalometer (AE33) was applied to characterize the dynamic variation of BrC. The positive matrix factorization (PMF) method together with aerosol mass spectrometer (AMS) data was used to apportion the sources of OA. The multiple linear regression (MLR) was used to explore the possible dynamic source contribution to BrC in two representative regions. Finally, source-specific BrC to light absorption and the radiative forcing in remote and urban areas was shown.

2. Methodology

2.1. Sampling sites

The field campaigns were simultaneously conducted from July 3 to August 3, 2022 in Tibet, and from July 16 to August 5, 2022 in urban Guangzhou, as depicted in Fig. 1. The Yangbajing site (YBJ site; 30.2°N, 90.45°E; 4,300 m above sea level [a.s.l.]) serves as a background location situated in the central QTP, approximately 90 km northwest of Lhasa City. With a permanent population of around 6,000, YBJ was supposed to be influenced by mixed plumes of regional transportation, while local anthropogenic emissions (e.g., light traffic flow and/or fuel combustion for domestic heating or cooking activities) were also found (Liu et al., 2021; Xiang et al., 2024). The urban site (GIG site; 23.1 °N, 113.4°E; 53 m a.s.l.) is set on the campus of the Guangzhou Institute of Geochemistry, Chinese Academy of Sciences (CAS), located in the downtown area of Guangzhou (permanent population: ~20 million). The GIG site is surrounded by transportation, commercial, and residential areas (Chen et al., 2021). During the whole campaigns, lower temperature and Relative Humidity (RH) were observed in the YBJ site (13.4 ± 4.6 °C, 52.1 ± 21.9 %) than those in the GIG site (29.5 ± 2.8 °C, 82.9 ± 12.7 %), which is expected due to the high altitude of Yangbajing (4,300 m) compared to Guangzhou (53 m). Southeast and north winds were dominant throughout the observation period at the YBJ site with an average wind speed of 2.3 ± 1.6 m s⁻¹, while the GIG site was dominated by south and southwest winds at a speed of 2.2 ± 0.9 m s⁻¹, as shown in Fig. 1. The date and time used in this study are reported using Beijing Time (BJT: UTC +8h).

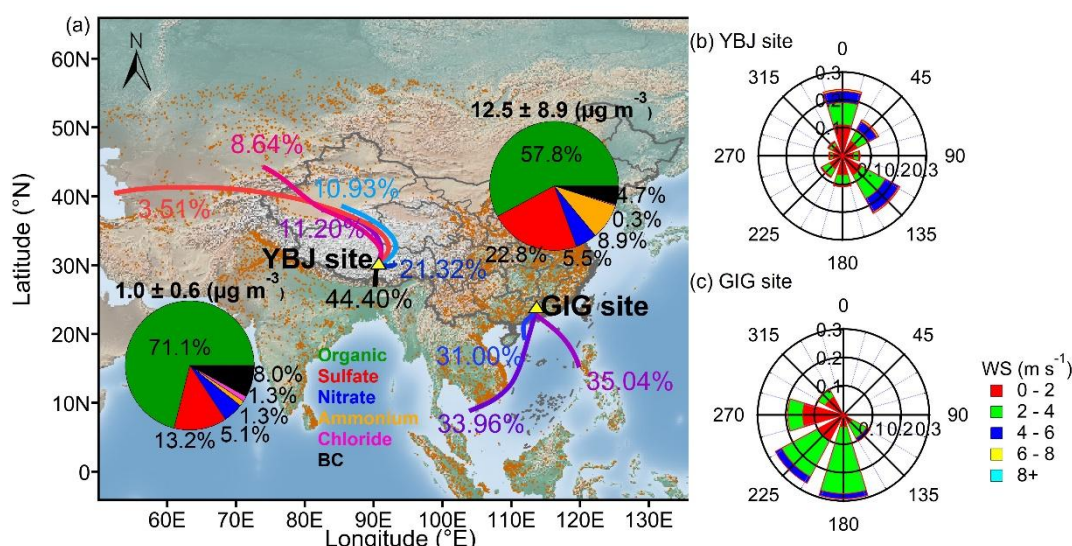


Figure 1. (a) Map of the locations of Yangbajing (YBJ) and Guangzhou (GIG) sites (yellow triangles). Solid colored lines represent the average back trajectory clusters during the whole campaign and the corresponding contributions plotted using the MeteoInfo version 2.2.6 developed by Wang (2019) (download from <http://www.meteothink.org>, last access time: 23 June 2025). The orange dots on the map indicate the location of the fire spot (download from *Active Fire Data Earthdata (nasa.gov)*, last access time: 14 December 2023). The pie charts represent the chemical compositions of submicron particulate matter (PM₁) along with their contributions at both sites during this campaign. The rose plots colored by wind speed (WS) at (b) the YBJ site and (c) the GIG site are also shown.

2.2. Light absorption coefficient measurement

The aerosol light absorption coefficients at both sites were measured by multi-wavelength Aethalometer (Model AE33, Magee Scientific Corp., Berkeley, CA, USA) at seven wavelengths (370, 470, 520, 590, 660, 880, 950 nm) with a high time resolution of 1 minute (Drinovec et al., 2015). Ambient aerosols were introduced into AE33 through a PM_{2.5} cyclone at a flow rate of 5 L min⁻¹. The AE33 collected aerosols via continuous pumping to a specific location on the filter belt. It then measured the transmitted light that passed through this sample-containing spot and a corresponding blank filter film spot. The instantaneous light-absorbing aerosols were determined by analyzing the variation in the attenuation rate of the transmitted light across the particulate-loaded filter membrane. To accurately reflect the real optical absorption coefficients of airborne aerosols, a real-time compensation parameter (k value) and a fixed filter multiple scattering parameter ($C_{\text{ref}} = 1.57$ for tetrafluoroethylene (TFE)-coated glass filter) are required to correct the optical attenuation coefficient measured on the filter membrane. For the k value, the "dual-point" measurement technique of AE33 avoids the "aerosol loading" effect during single filter tape membrane sampling, enabling real-time calculation corrections for load compensation parameters (Drinovec et al., 2015). Multiple studies have shown that the C value (2.8–7.8) varies with different wavelengths and observation locations (Qin et al., 2018; Collaud Coen et al., 2010; Tian et al., 2023; Zhang et al., 2021). In this study, the C_{final} values of 3.34 ($2.23 \times C_{\text{ref}}$) for the YBJ site and 3.6 (2.3×1.57) for the GIG site were applied based on the previous studies in Tibet (Zhang et al., 2021) and Guangzhou (Cai et al., 2024), also using the AE33 instrument.

2.3. OA measurement and source apportionment

The main chemical compositions of submicron aerosols including OA, nitrate, sulfate, ammonium,

and chloride were measured using a soot particle time-of-flight aerosol mass spectrometer (SP-AMS; Aerodyne Research Inc., Billerica, MA, USA) (Onasch et al., 2012) at YBJ site and a time-of-flight aerosol chemical speciation monitor (ToF-ACSM; Aerodyne Research Inc., Billerica, MA, USA) (Fröhlich et al., 2013) at GIG site. During this campaign, both the SP-AMS and ToF-ACSM shared the same sampling inlet with co-located AE33. The setup diagram can be found in Fig. S1. The time resolution of SP-AMS and ToF-ACSM was 1 min and 40 s, respectively. For SP-AMS, dual vaporization mode and tungsten-only V mode were alternatively applied every 4 minutes. In this study, tungsten-only mode, which resembled the traditional HR-ToF-AMS, was applied. The ionization efficiency (IE) calibration was done using monodispersed NH_4NO_3 aerosols before and after campaigns. The relative ionization efficiency (RIE) of sulfate and ammonium is 1.26 and 4.24 for SP-AMS and 1.22 and 3.39 for ACSM, while a default RIE of 1.4 was used for OA. A constant collection efficiency (CE) of 0.5 was used for YBJ measurement due to the quite low mass fraction of ammonium nitrate ($< 40\%$), while the composition-dependent collection efficiency (average CDCE = 0.52) (Middlebrook et al., 2012) was applied in GIG measurement. The SP-AMS data were processed using SQUIRREL (v1.65) and PIKA (v1.25A), embedded in Igor Pro (v6.37; WaveMetrics, Inc., Lake Oswego, OR, USA), while ToF-ACSM data were processed using Tofware 3.2.4 (Tofwerk AG, Thun, Switzerland).

The positive matrix factorization (PMF) (Ulbrich et al., 2009; Zhang et al., 2011) was applied to the OA spectral matrix to resolve the sources of OA at both sites. More detailed information can be found in Text S1 and Table S2. The final OA source apportionment at the YBJ site and GIG site are shown in Fig. S4. At the YBJ site, five OA factors were finally resolved using free PMF with PMF3.05 (Ulbrich et al., 2009), including hydrocarbon-like OA (HOA, 11 %) mainly from traffic emissions, biomass burning OA (BBOA, 9 %), biofuel-OA (13 %), less-oxidized oxygenated OA (LO-OOA, 42 %), and more-oxidized oxygenated OA (MO-OOA, 25 %). At the GIG site, OA was resolved based on Multilinear Engine 2 (ME-2; SoFi 6.8) (Canonaco et al., 2013), and the standard BBOA mass spectra (Hu et al., 2016; Hu et al., 2013) were introduced as an external constraint to fully constrain (a value = 0) BBOA factor at the GIG site. Finally, five factors were chosen with MO-OOA (49 %) and LO-OOA (22 %) dominating the total OA, followed by cooking-related OA (COA, 13 %), BBOA (8 %), and HOA (8 %).

2.4. Calculation of BC and BrC light absorption coefficients

The Absorption Ångström exponent (AAE) method was widely used to distinguish BC and BrC light absorption coefficients measured by Aethalometer (Lack and Langridge, 2013). Previous studies demonstrated that the aerosol light absorption at 880 nm was entirely dominated by BC (Kirchstetter et al., 2004). The absorption coefficients of BrC at shorter wavelengths (370, 470, 520, 590, and 660 nm) can be calculated by combining the AAE (Lack and Langridge, 2013) with BC light absorption at 880 nm, using the following Equations:

$$\text{Abs}_{\text{BC}}(\lambda) = \text{Abs}(880) \times \left(\frac{880}{\lambda}\right)^{\text{AAE}_{\text{BC}}} \quad (1)$$

$$\text{Abs}_{\text{BrC}}(\lambda) = \text{Abs}(\lambda) - \text{Abs}_{\text{BC}}(\lambda) \quad (2)$$

Here, $\text{Abs}_{\text{BC}}(\lambda)$ and $\text{Abs}_{\text{BrC}}(\lambda)$ (M m^{-1}) represent light absorption coefficients of BC and BrC at wavelength λ (nm), respectively. $\text{Abs}(\lambda)$ represents the total aerosol light-absorbing coefficients at the

wavelength λ , which can be calculated by the mass concentration of BC ($\mu\text{g m}^{-3}$) multiplied by mass absorption cross-section (MAC; $\text{m}^2 \text{g}^{-1}$) at wavelength λ , denoted as $\text{Abs}(\lambda) = \text{BC}(\lambda) \times \text{MAC}(\lambda)$. In this study, default MAC values for BC (18.47, 14.54, 13.14, 11.58, 10.35, and $7.77 \text{ m}^2 \text{g}^{-1}$ for 370, 470, 520, 590, 660, and 880 nm, respectively) were used (Drinovec et al., 2015). AAE_{BC} denotes the wavelength dependence of pure BC particles, which was usually assumed as 1 to calculate the light absorption coefficient of BC (Tian et al., 2023; Zhang et al., 2022b). However, due to the multiple effects (e.g., lensing effect) caused by the mixing of non-BC/coating materials and BC during atmospheric evolution, the light absorption of BC can be enhanced (Jacobson, 2001; Liu et al., 2017; Peng et al., 2016). Moreover, AAE_{BC} was also found to be varied as a function of absolute wavelength values (Luo et al., 2022), which was suggested as 0.8–1.4 for BC particles (Lack and Langridge, 2013; Kasthuriarachchi et al., 2020a; Zhai et al., 2022; Corr et al., 2012). We summarized the AAE_{BC} values from the literatures in Table S3, and found all the actually used values are between 1 – 1.18. E.g., within one campaign (Kasthuriarachchi et al., 2020), 93% of directly measured AAE from ambient core BC in Singapore was found to be less than 1.1. An AAE_{BC} of 1.4 represents a typical high value, considering the mixing of non-absorbing material with BC in the extreme case (Lack and Langridge, 2013). For our study, we found ~82% of data was negative value when the AAE_{BC} of 1.4 was applied, which is unrealistic to apply here. Thus, to investigate the uncertainty of BrC estimated here, AAE_{BC} of 0.8 to 1.2 was applied, which results in 2–13 % and 4–14 % of BrC contribution to total light absorption from 370 to 660 nm for the YBJ site and GIG site, respectively.

2.5. Sources contribution to BrC light absorption based on multiple linear regression (MLR)

To further understand the sources of BrC light absorption, a multiple linear regression (MLR) was used to apportion the BrC light absorption to different components resolved by the source apportionment of OA (Kasthuriarachchi et al., 2020a; Qin et al., 2018; Tian et al., 2023; Zhang et al., 2021), using the following Equations:

$$\text{Abs}_{\text{BrC}}(\lambda) = \sum(a \times \text{Factor}) + \text{intercept} \quad (3)$$

Here, $\text{Abs}_{\text{BrC}}(\lambda)$ (M m^{-1}) represents the BrC light absorption coefficient at a wavelength of λ (370, 470, 520, 590 and 660 nm); a represent the regression coefficients of different OA components, which can be regarded as MAC ($\text{m}^2 \text{g}^{-1}$) values (Kasthuriarachchi et al., 2020a); Factor ($\mu\text{g m}^{-3}$) represents the mass concentration of OA from different sources (see Sect. 2.3).

2.5.1. Uncertainty analyses for the MLR method

Multicollinearity is an important factor leading to inaccurate estimation of regression coefficients in the MLR model. To estimate the uncertainty of the MLR method, we tested different scenarios with varied PMF factor combinations based on their correlation. To compare with other published results and simplify the calculation process, the MLR uncertainty at 370 nm was estimated. In general, four cases were chosen for the YBJ campaign, and three cases were chosen for the GIG campaign by combining or excluding collinear factors. The detailed information can be found in Text S2 and Table S5. All the cases yield similar low coefficients for the SOA factor and high coefficient values for the POA factors, suggesting the validity of MLR analysis shown here.

With each solution, the uncertainty of the regression coefficient (i.e., MAC) for the individual OA components input into the MLR model was evaluated using Monte Carlo simulations. For the Monte Carlo calculation input, the uncertainty of the PMF factor mass concentration needs to be evaluated. A bootstrap analysis (100 iterations; (Ulbrich et al., 2009) was applied, which shows a 9–36 % uncertainty for the PMF factors at the YBJ site and 3–9 % at the GIG site, as shown in Table S6. The uncertainty for the coefficient of BrC at 370 nm was estimated to be 43 % for the YBJ site and 36 % for the GIG site based on the lower (0.8) and upper limit of (1.2) previously reported AAE_{BC} range. The total uncertainties of each coefficient for each PMF factor were then calculated by Monte Carlo with 10,000 simulations.

Considering that biomass burning is widely reported as an important source of BrC light absorption and regarded as a warming agent affecting global climate (Wang et al., 2025), we consider all biomass burning related contributing sources when run the MLR model in the case of eliminating the collinearity problem in this study. As previously reported in the literature (Kasthuriarachchi et al., 2020a; Qin et al., 2018), the MAC of COA is nearly zero, thus, the light absorption by COA at GIG site was not considered in this study. We finally combined the BBOA factor and Biofuel–OA factor as BBOA (case 3) for the YBJ site and included BBOA, HOA as input (case 3) for the GIG site (Table 1). The SOA (LO–OOA and MO–OOA) at GIG site were not inputted here due to the collinearity with BBOA, the light absorption contribution and MAC values (case 3) at GIG site will be lower limits for SOA. For its upper limits, Case 2 in the supporting information (Text S2; Table S5) was also shown (Fig. S9). At the YBJ site, the MAC uncertainty of BBOA (26.4 %), HOA (20.8 %), LO–OOA (56.3 %), and MO–OOA (57.9%) was found, as shown in Table S7. For the GIG site, the MAC uncertainty of HOA (5.8 %), BBOA (6.8 %), and intercept (21.6 %) was estimated. With the final solution, the total light absorption calculated from the predicted regression coefficients (i.e., MAC) showed strong agreement with the measured values, as indicated by the slopes (YBJ: 0.9; GIG: 0.9) and Pearson correlation coefficients (YBJ: $R = 0.8$; GIG: $R = 0.7$). These results suggest the robustness of the regression analyses. However, a notable intercept (0.37 M m^{-1} , representing 13 % of the total light absorption) was observed in the MLR model at the GIG site. This intercept indicates a portion of BrC light absorption that could not be explained by the OA factors, potentially due to uncertainties associated with the MLR method. The detailed discussion of MAC and BrC light absorption contributions is shown in Sect.3.2.

Table 1. Regression coefficients (MAC) of the final case of multiple linear regression (MLR) at 370, 470, 520, 590, and 660 nm at the YBJ site and GIG site. Note that the SD represents the Standard deviation purely calculated from the MLR model.

YBJ site					
MAC (Average \pm SD)	Wavelength (nm)				
	370nm	470nm	520nm	590nm	660nm
biofuel–OA	1.11 ± 0.11	0.42 ± 0.05	0.11 ± 0.03	0.11 ± 0.02	0 ± 0.03
HOA	2.08 ± 0.3	1.11 ± 0.14	0.18 ± 0.10	0.16 ± 0.06	0 ± 0.07
LO–OOA	0.15 ± 0.08	0.14 ± 0.03	0.07 ± 0.02	0.04 ± 0.02	0.01 ± 0.02
MO–OOA	0.18 ± 0.18	0.19 ± 0.08	0 ± 0.06	0.02 ± 0.04	0 ± 0.04
Intercept	0 ± 0.02	0.02 ± 0.01	0.04 ± 0.01	0.02 ± 0	0.03 ± 0
GIG site					
MAC (Average \pm SD)	Wavelength (nm)				
	370nm	470nm	520nm	590nm	660nm
BBOA	1.91 ± 0.21	0.90 ± 0.10	0.35 ± 0.06	0.40 ± 0.04	0.20 ± 0.02
HOA	2.57 ± 0.28	1.40 ± 0.13	0.72 ± 0.07	0.38 ± 0.06	0.13 ± 0.03

2.6. Calculation of radiative forcing

The estimation of the direct radiative forcing caused by BrC was conducted using a model known as the “simple forcing efficiency (SFE)”, which can provide a radiative forcing (W g^{-1}) based on a given mass of aerosols. Although the resulting value of SFE is lower than that projected by climate models, it still serves as a valuable tool for gauging the sensitivity of various input parameters (Bond et al., 2006; Chylek and Wong, 1995). In this study, we used a modified version of the wavelength-dependent SFE without considering mass scattering and expressed it as follows (Tian et al., 2023; Zhang et al., 2020a; Zhang et al., 2022b):

$$SFE(\lambda) = \frac{S(\lambda)}{4} \times \tau_{atm}^2 \times (1 - F_c) \times 4\alpha_s \times MAC(\lambda) \quad (4)$$

Here, $S(\lambda)$ ($\text{W g}^{-1} \text{nm}^{-1}$) is the wavelength-dependent solar irradiance based on the ASTM G173–03 Reference Spectra (<https://www.nrel.gov/grid/solarresource/spectra-am1.5.html>); τ_{atm} represents atmospheric transmission (0.79); F_c is the cloud fraction (0.6); α_s is the surface albedo (global average 0.19) (Chen and Bond, 2010). $MAC(\lambda)$ ($\text{m}^2 \text{g}^{-1}$) is the mass absorption cross-section of different OA components at wavelength λ at a 1 nm resolution. The $MAC(\lambda)$ can be calculated according to the power-law fitting results between the MAC from each OA component at different wavelengths (370 to 660 nm) in this study. Note that the SFE represents a straightforward calculation designed to ascertain the relative significance of diverse optical properties of radiative forcing. However, to accurately determine forcing efficiency, a comprehensive radiative transfer model is still necessary (Efremenko and Kokhanovsky, 2021).

3. Results and discussion

3.1. Overview of Aerosol light absorption

The summarized total aerosol light absorption coefficients (Abs_{total}) as a function of wavelength at two sites are shown in Fig. 2 and Table S8. The more detailed temporal evolutions of Abs_{total} at two sites are displayed in Fig. S7. In general, the average Abs_{total} ranged from 0.6 to 1.6 M m^{-1} (370 to 950 nm) at the YBJ site, while the 7–8 times higher Abs_{total} (4.2–13.2 M m^{-1}) was observed at the GIG site (Fig. 2a and Table S8). This discrepancy was primarily attributed to the much higher aerosol mass concentration in urban Guangzhou (PM_{10} : $12.5 \pm 8.8 \mu\text{g m}^{-3}$) compared to the background Tibetan region (PM_{10} : $1.0 \pm 0.6 \mu\text{g m}^{-3}$) (Figs. 2a and 2b). The campaign-averaged BrC light absorption coefficients (Abs_{BrC}) and its contribution to total aerosol light absorption increased with decreasing wavelength at both sites (AAE of BrC is 2.6 for YBJ and 3.2 for GIG; Figs. 2c, 3b, and 3d), indicating strong BrC light absorption at shorter wavelengths. At 370 nm, where BrC contributes the most to the total light absorption, Abs_{BrC} in Tibet ($0.2 \pm 0.3 \text{ M m}^{-1}$) was found to be a factor of 13 lower than that in Guangzhou ($2.9 \pm 2 \text{ M m}^{-1}$). This finding is consistent with the much lower OA mass concentrations (by a factor of ~10) at the YBJ site (OA: $0.7 \pm 0.4 \mu\text{g m}^{-3}$ vs $6.9 \pm 5.8 \mu\text{g m}^{-3}$ at GIG; Figs. 2b and 2c). A positive correlation between BrC light absorption and OA mass concentration was also observed across different field studies (Fig. 2d; Table S4). This relationship aligns with the findings of Wang et al. (2022a), who reported that

BrC light absorption at 365 nm positively correlates with OA mass concentration across various sources.

To facilitate a visual comparison with previous studies, the literature-reported BrC light absorption results at 365 or 370 nm are summarized in Fig. 3e and Table S4. This wavelength was chosen due to its higher BrC abundance across different wavelengths, which leads to less uncertainty. At the YBJ site, the campaign-averaged Abs_{BrC} at 370 nm (0.2 M m^{-1} ; $0.03\text{--}0.6 \text{ M m}^{-1}$ for 5–95 % range; Table S4) was lower than the values reported across the Qinghai–Tibet Plateau (QTP) ($0.6\text{--}14.9 \text{ M m}^{-1}$), e.g., the BrC light absorption at YBJ site (central QTP) was much lower than those observed at the edges of the QTP, such as Ngari ($5.9\text{--}10.7 \text{ M m}^{-1}$), Qomolangma Station (QOMS, 4.4 M m^{-1}), and Gaomeigu (12.3 M m^{-1}), where elevated values are primarily influenced by the cross-border transport of biomass burning plumes during pre-monsoon or post-monsoon period (Zhang et al., 2021; Tian et al., 2023; Zhu et al., 2017). The low BrC light absorption at YBJ can be attributed to the extremely low mass loading of OA ($0.7 \pm 0.5 \mu\text{g m}^{-3}$), which is influenced by wet deposition from precipitation and specific atmospheric circulation patterns during July, a monsoon period (Xu et al., 2018; Zhao et al., 2013). In addition, the absolute BrC light absorption coefficient at YBJ was comparable to levels measured in certain remote regions, such as the Arctic ($0.04\text{--}0.2 \text{ M m}^{-1}$ at 365 nm) (Barrett and Sheesley, 2017; Yue et al., 2022; Yue et al., 2019), highlighting the relatively clean atmospheric conditions at YBJ as a background site in Tibet. In general, BrC light absorption in urban areas tends to be higher or comparable to that in QTP regions due to typically higher OA mass concentrations (Fig. 2d).

Although a relatively large discrepancy in the absolute light absorption coefficient of BrC was observed between the YBJ and GIG sites, the contribution of BrC to total aerosols ($\text{Fraction}_{\text{BrC}}$) was comparable at both sites (15 % vs. 21 %), highlighting the significant role of BrC in light absorption at both locations. We summarized $\text{Fraction}_{\text{BrC}}$ across different studies and regions, as shown in Fig. 3e and Table S4, which reveals that $\text{Fraction}_{\text{BrC}}$ values generally fall within a range of 8–58 %. Notably, the $\text{Fraction}_{\text{BrC}}$ in the Qinghai–Tibet Plateau (QTP) is comparable to that in urban areas. Furthermore, a clear trend shows that $\text{Fraction}_{\text{BrC}}$ at the edges of the northeastern (such as Qinghaihu Lake) and south QTP (such as Ngari) tends to be higher than in the central QTP (such as Beiluhe, Namco, and YBJ site), and the $\text{Fraction}_{\text{BrC}}$ in Northern China is generally higher than in Southern China. To better understand the factors driving the $\text{Fraction}_{\text{BrC}}$, we investigated the relationship between the contribution of primary organic aerosol (POA) and secondary OA (SOA) to total OA and the $\text{Fraction}_{\text{BrC}}$. As illustrated in the inset plot of Fig. 3e, a positive correlation was observed, suggesting that the enhanced POA contribution will increase the BrC light absorption contribution in carbonaceous aerosol. For sites at the edges of the QTP and in northern Chinese cities, where the $\text{Fraction}_{\text{BrC}}$ is higher, OA is significantly influenced by the combustion of solid fuels, including coal and biomass, particularly during winter. This finding signifies the importance of POA in contributing to ambient BrC light absorption.

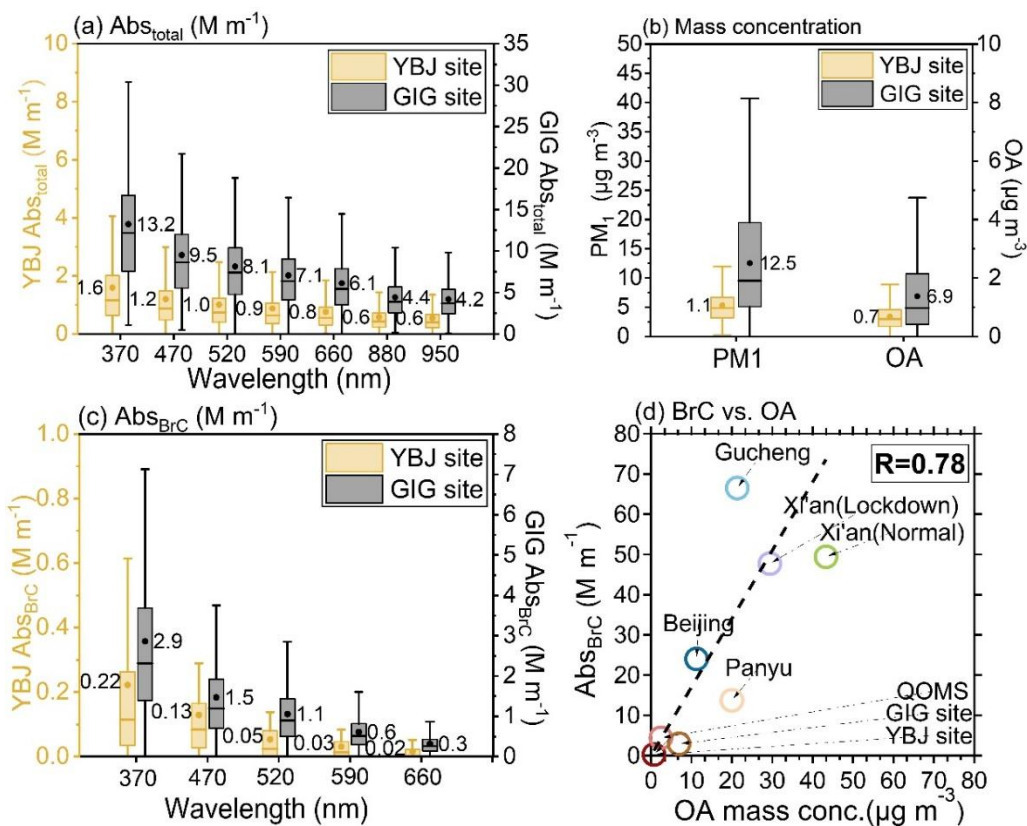


Figure 2. Box plot of the light absorption coefficient of total aerosols (Abs_{total}) (a) and BrC (Abs_{BrC}) (c) from 370nm to 950nm at YBJ (left Y axis) and GIG (right Y axis) sites. The Abs_{BrC} was separated from the Abs_{total} based on Eq. (1) and (2) using $AAE_{BrC}=1$, (b) Box plot of PM_{10} and OA mass concentrations at the YBJ site and GIG site. The whiskers indicate the 90th and 10th percentiles, the upper and lower boundaries of boxes indicate the 75th and 25th percentiles, the lines in the boxes indicate the median values, and the markers for the mean values. (d) Scatter plot of the BrC absorption coefficient as a function of OA concentration (data from the literature (Table S3)).

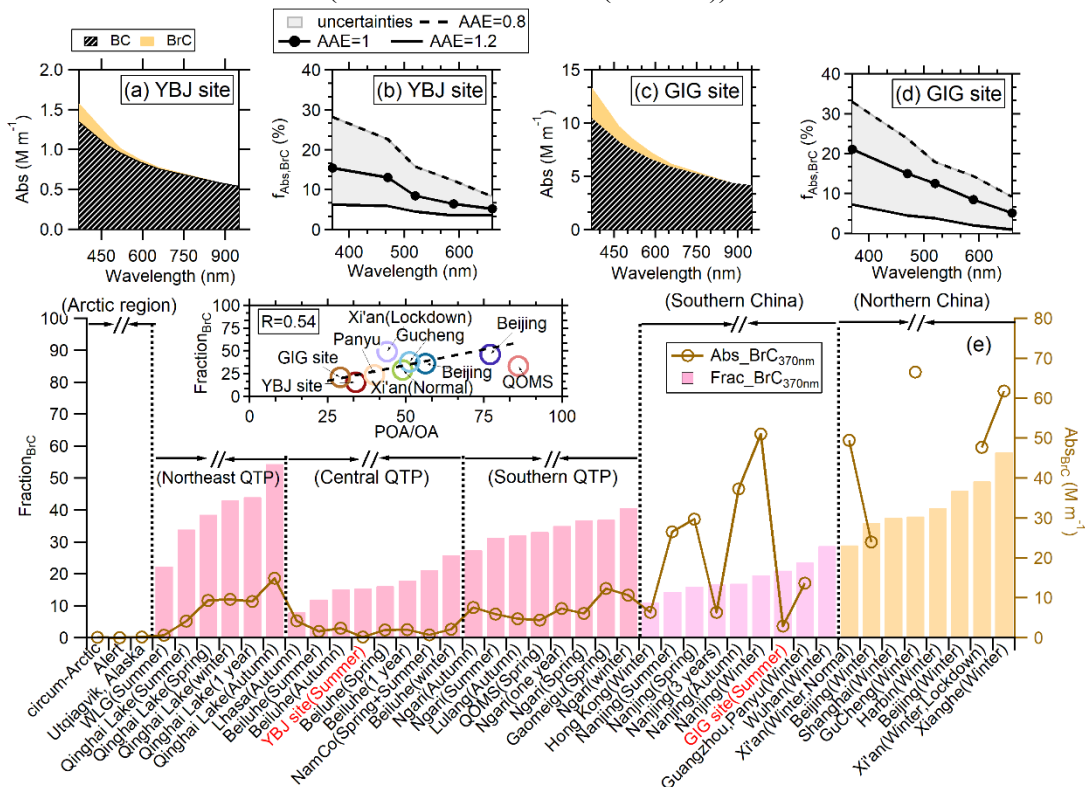


Figure 3. The absolute light absorption coefficients of BC (Abs_{BC}) and BrC (Abs_{BrC}) at wavelengths from 370 to 880 nm at the (a) YBJ site and (c) GIG site, respectively. The BrC was calculated based on the AAE of BC (AAE_{BC})=1. The contribution of BrC light absorption at wavelengths from 370 to 660 nm at the (b) YBJ site and (d) GIG site, respectively. The grey-filled area represents variations in the BrC light absorption fraction caused by the AAE_{BC} from 0.8 (dashed line) to 1.2 (solid line). The circle makers are the average value estimated based on AAE_{BC} =1. (e) The summary of BrC light absorption coefficients (the brown circles) and their contributions to total light absorption coefficients at 370 nm from the literature results. The results were categorized according to the locations of their observation sites (Arctic region, Qinghai–Tibet Plateau (QTP) region, Southern China, and Northern China). The inset plot represents the relationship between the BrC light absorption fraction and the contribution of primary OA to total OA. The detailed information is provided in Table S4.

To further elucidate the dynamic evolution of BrC light absorption, the diurnal variations of $Abs_{BC,370}$ and $Abs_{BrC,370}$ were displayed in Fig. 4 (Abs_{total} at different wavelengths was shown in Fig. S8). $Abs_{BC,370}$ and $Abs_{BrC,370}$ at both sites peaked simultaneously, indicating combustion source for BC are also important contributor to the BrC in both studies. At the YBJ site, the peaks at 08:00 and 22:00 align with the NO_x ($NO + NO_2$) and CO, which were due to the local or regional anthropogenic emissions (such as vehicle emissions and other combustion activities) in Tibet. In other periods, a background $Abs_{BrC,370}$ value of 0.19 M m^{-1} was found at YBJ site. The diurnal patterns of $Abs_{BC,370}$ and $Abs_{BrC,370}$ at the GIG site, which showed an enhancement at 10:00 and a stronger peak at 21:00, were slightly different from those at the YBJ site. This difference suggests distinct source emissions or secondary formation patterns between the two locations (Fig. 4b and Fig. S8b). Higher increase of NO_2/NO and CO at night in Guangzhou was also found, implying the potentially important contribution of vehicles and other combustion sources, e.g., biomass burning, at this urban site.

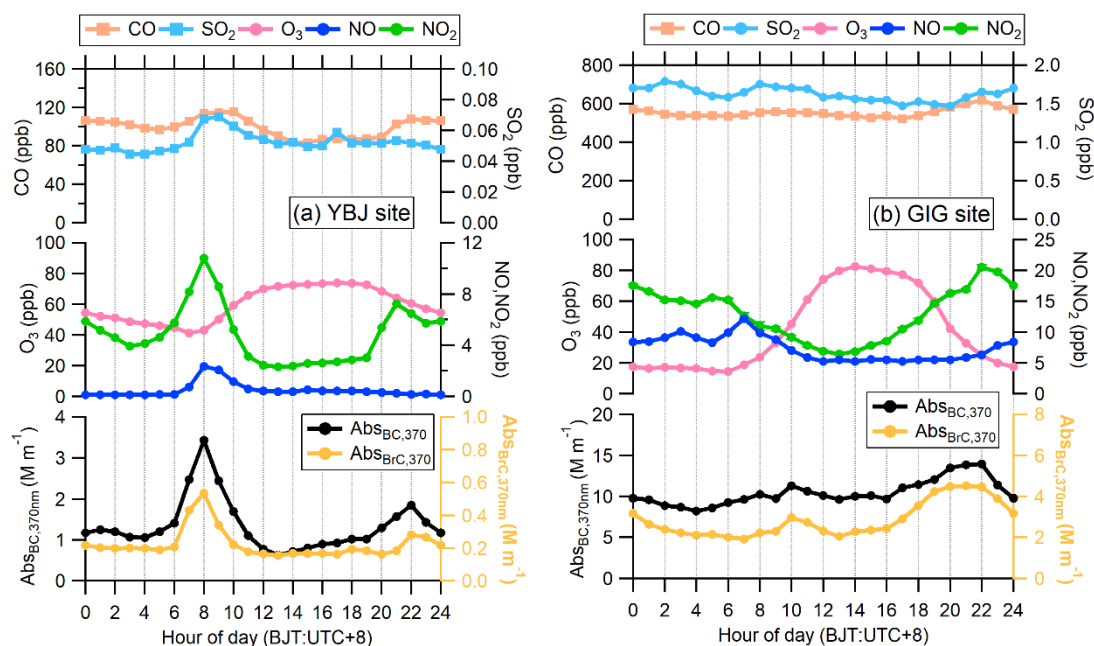


Figure 4. Diurnal variations of gas pollutants and BC/BrC light absorptions at 370 nm at (a) YBJ site and (b) GIG site.

3.2. Optical properties of source-specific BrC

To further identify the source of absorptive BrC in this study, we applied the multiple linear regression (MLR) method to attribute BrC light absorption at different wavelengths to OA components from various sources, based on the PMF analysis conducted at the YBJ and GIG sites. The detailed

methodology is described in Sect.2.5.

At the YBJ site in Tibet, the MLR model identified BBOA (the sum of BBOA and Biofuel-OA, as defined in Sect.2.5.1 and Text S1), HOA, LO-OOA, and MO-OOA as the main contributors to BrC light absorption. As shown in Fig. 5a, secondary organic aerosol (SOA, comprising LO-OOA and MO-OOA) dominated the total OA mass concentration (67 %), followed by BBOA (22 %) and HOA (11 %). However, for BrC light absorption, POA (including BBOA and HOA) emerged as the most significant contributor, with BBOA contributing the most (40 %), followed by HOA (38 %) and SOA (22 %), signifying the key role played by biomass burning and vehicle emissions for BrC light absorption at the YBJ site. Indeed, both the BBOA ($R = 0.77$) and HOA ($R = 0.64$) show better correlations with BrC light absorption at 370 nm than other factors (Fig. S10). A similar trend was observed at the GIG site in urban Guangzhou (Fig. 5d), where POA (including BBOA, HOA) represented only 29 % of the total OA mass concentration but contributed disproportionately to BrC light absorption (89 %). These findings highlight the critical role of primary BrC in light absorption during our measurement campaign. In the following, we will discuss each source contribution to BrC in detail.

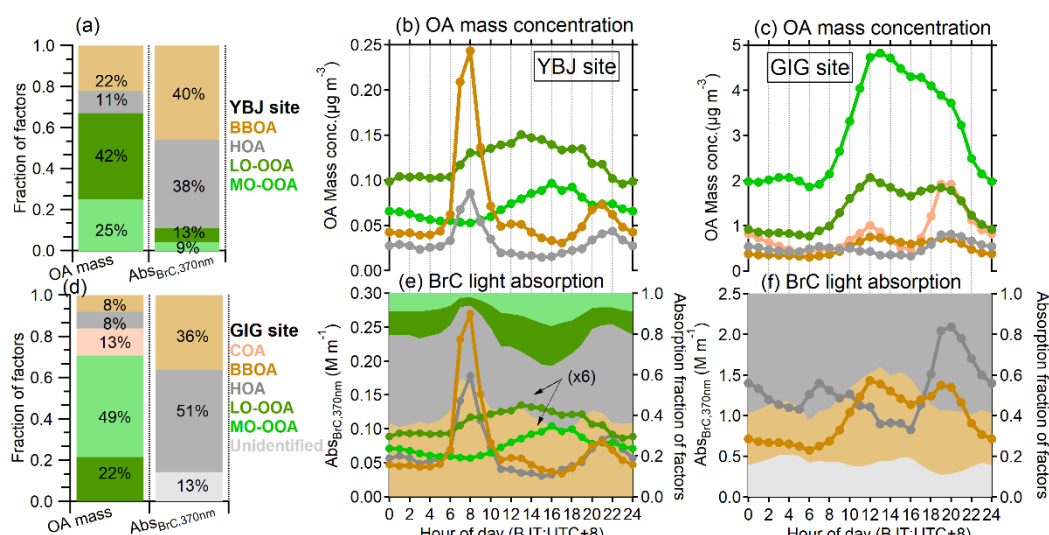


Figure 5. (a) The contributions of different OA factors to total OA (left bars) and contributions of different OA factors to BrC light absorption at 370 nm (right bars) at the YBJ site and GIG site. The diurnal variations of OA factors mass loading and the light absorption of different OA components at 370 nm at the YBJ site (b, e) and GIG site (c, d). The “x6” indicates the light absorption of LO-OOA and MO-OOA at 370 nm expanded by 6 times to better guide the eyes for the trend of diurnal patterns.

3.2.1. Optical properties of BBOA

The diurnal variations of the biomass burning OA (BBOA) light-absorbing coefficients at 370 nm for the YBJ site exhibited strong correlation with BBOA mass concentration (Fig. 5b, 5e), characterized by distinct peaks at 08:00 and 21:00 local time. This temporal pattern aligns with the observation reported in urban Lhasa, which exhibits a peak at 9:00 and 22:00, respectively (Zhao et al., 2022). The BBOA peaks at the YBJ site (central QTP) in the morning and night are primarily influenced by local anthropogenic activities, i.e., the traditional Weisang ritual. The Weisang activity, prevalent throughout Tibet, including the sparsely populated YBJ region, typically occurs twice daily in the morning and evening. Wherever Tibetans are living, almost all have a Weisang furnace. This cultural practice involves the combustion of specific organic materials, including wormwood, cypress branches, highland barley,

ghee lamp, and zanba in dedicated stoves, generating characteristic mulberry smoke for religious purposes (Zhang et al., 2020b; Zhang et al., 2022a). Additionally, traditional residential solid biofuel combustion (primarily yak dung and wood) for cooking and space heating contributes significantly to BBOA loading in the plumes (Shen et al., 2021). The AMS spectral similarity between ambient BBOA and emissions from both Weisang activities and solid biofuel combustion (Zhang et al., 2022a), showing the highest signals of m/z 41, 43 and 55, and high abundance of hydrocarbon ions above m/z 60, provides further evidence for their dominant contributions to regional BBOA in Tibet. The hourly wind roses show that the wind direction at the YBJ site exhibits a regular diurnal pattern, with a gradual change from the north wind in the morning to the stronger southeast wind in the afternoon (Fig. S11). Combining the Bivariate polar plots, the higher morning peak (6:00 to 11:00) of BBOA was from the nearby residential area under low wind speed condition (Figs. S12, 13), while the lower evening peak was due to the dilution of stronger wind speed in the evening (17:00 to 24:00) (Figs. S12, 13).

Compared to Tibet, the urban Guangzhou exhibited fundamentally different diurnal patterns in both BBOA mass concentration and BrC light absorption coefficients (Fig. 5c, 5f), where the BBOA peak occurs around noon and nighttime, reflecting the different biomass burning activity between Tibet and Guangzhou. Previous literature (Cai et al., 2023; Wang et al., 2017) reported that the agriculture burning at the suburban areas likely dominated the BBOA mass concentration at Guangzhou areas.

The mass absorption coefficient (MAC, $\text{m}^2 \text{g}^{-1}$), a crucial optical parameter for BrC characterization, quantifies the light absorption capacity per unit mass of OA. Our measured MAC values for BBOA ($1.11\text{--}2.54 \text{ m}^2 \text{g}^{-1}$ at YBJ vs. $1.91 \pm 0.21 \text{ m}^2 \text{g}^{-1}$ at GIG; Table 1, S4) fall within the lower range of previously reported values ($0.6\text{--}8 \text{ m}^2 \text{g}^{-1}$; Fig. 6, Table S10). This variability in MAC values from similar combustion sources, as documented in numerous studies (Budisulistiorini et al., 2017; Chen and Bond, 2010; Martinsson et al., 2015; Saleh et al., 2014), arises from multiple factors, including fuel composition, combustion conditions, and efficiency. The relatively low MAC values observed at both sites in this study can be attributed to several reasons: 1) Both studies were conducted in the summer time, which coincided with intense solar radiation and elevated oxidant concentrations, promoting photobleaching of chromophores in fresh BBOA (Sumlin et al., 2017). Recent experimental evidence (Hems et al., 2021) demonstrates rapid (minutes to hours), nonlinear photobleaching kinetics of fresh BrC, highlighting the complex nature of these atmospheric processes. Especially for the Guangzhou samples, BBOA from regional transport was likely subjected to a longer oxidation process, which led to a lower MAC at the GIG site. 2) Combustion conditions and fuel type significantly influence MAC values (Martinsson et al., 2015). Zhang et al. (2022a) systematically characterized Tibetan biofuel emissions, revealing that Weisang materials and yak dung produce abundant OA with relatively low light absorption efficiency due to incomplete combustion. Similarly, Moschos et al. (2024) showed higher MAC values for hardwoods ($0.8\text{--}1.6 \text{ m}^2 \text{g}^{-1}$) versus animal dung ($0.2\text{--}0.7 \text{ m}^2 \text{g}^{-1}$); 3) At YBJ site, the multiple linear regression (MLR) model incorporates biofuel-OA, which contains cooking OA, typically characterized by weak/no absorption (Kasthuriarachchi et al., 2020a; Qin et al., 2018). The spectral similarity between COA and biofuel-OA prevents complete separation in PMF analyses. Sensitivity analyses (Sect.2.5.1 and Text S1; Table S5) demonstrate MAC_{BBOA} variability ($1.11\text{--}2.54 \text{ m}^2 \text{g}^{-1}$) under different model assumptions, with maximum values ($2.54 \text{ m}^2 \text{g}^{-1}$) obtained when excluding biofuel-OA contributions.

3.2.2. Optical properties of HOA

The light-absorbing coefficient of HOA at the YBJ site exhibited a pronounced peak at 08:00 and a minor peak at 22:00 (Fig. 5b, 5e), aligning with the diurnal variation of HOA mass concentration. In the Tibetan Plateau region, heavy-duty diesel trucks, which are critical for transporting essential goods across this remote area, constitute a significant emission source alongside gasoline vehicles (Liu et al., 2021; Xiang et al., 2024). Our observation site is located less than 1 km (straight-line distance) from the G6 Beijing–Tibet Expressway. The Bivariate polar plots also show that the HOA was affected by the wind direction, with the morning peak affected by the north and northeast plumes, while the evening peak was affected by the southeast and northeast plumes with stronger wind speed. The traffic emission during the night is more regional than that in the morning, which was supported by the fact that the NO mass concentration was only enhanced during the morning (2.3 ppb) but not during the night, while an obvious NO₂ peak at both periods (10 ppb and 7.2 ppb) was observed (Fig. 4a). Combining the Rose plots and Bivariate polar plots (Fig. S12), the evening peak of HOA was also affected by the southeast and northeast traffic emissions plumes with stronger wind speed, which also supported the regional nature of evening HOA peak. At the urban GIG site in Guangzhou, HOA light absorption coefficients displayed a bimodal distribution with a moderate morning peak at 07:00 and a stronger evening peak at 20:00 (Figs. 5c, 5f), consistent with typical urban traffic emissions for rush hours (Chen et al., 2021). The much higher evening peak than the morning peak coincides with rush-hour traffic congestion, suggesting intensified vehicular emissions during these periods, as well as the effects of meteorological conditions (e.g., reduced boundary layer height). The different diurnal variation of light absorption on both HOA and BBOA between Tibetan and Guangzhou observations highlights the regionally specific emission drivers from different areas.

In our study, the MAC of HOA for Tibet ($2.08 \pm 0.3 \text{ m}^2 \text{ g}^{-1}$) and urban Guangzhou ($2.57 \pm 0.28 \text{ m}^2 \text{ g}^{-1}$) (Table 1) were in the higher ranges of HOA MAC values reported by other studies ($0.4\text{--}2.04 \text{ m}^2 \text{ g}^{-1}$) (Table S10). Note that since the OOA factors in Guangzhou were not considered in the BrC source apportionment due to collinearity with BBOA, the HOA reported here shall be an upper limit. The results of both observation sites show that the HOA MAC ($2.08 \text{ m}^2 \text{ g}^{-1}$ at YBJ site vs. $2.57 \text{ m}^2 \text{ g}^{-1}$ at GIG site) is larger than the BBOA MAC ($1.11\text{--}2.54 \text{ m}^2 \text{ g}^{-1}$ at YBJ vs. $1.91 \pm 0.21 \text{ m}^2 \text{ g}^{-1}$ at GIG). To investigate whether the MAC from ambient BBOA or HOA is higher, we summarized the MAC results from different field studies in Fig. 6, yet no clear conclusion can be drawn. E.g., Higher HOA MAC than BBOA was observed for central Amazon study (De Sá et al., 2019) and Mexico study (Retama et al., 2022), while much higher BBOA MAC than HOA was found for Xianghe, Athens, Paris studies (Kaskaoutis et al., 2021; Wang et al., 2019b; Zhang et al., 2020c). For emission experiments, higher MAC for diesel exhaust than crop and wood have also been observed (Cheng et al., 2011; Du et al., 2014). In addition, very strong light absorption capacity (MAC; $5\text{--}6 \text{ m}^2 \text{ g}^{-1}$) (Table S10) induced by coal combustion (Wang et al., 2019b; Zhang et al., 2022b) was found, which is comparable to these extreme MAC values reported from biomass burning ($5\text{--}7.5 \text{ m}^2 \text{ g}^{-1}$) (Table S10) (Kaskaoutis et al., 2021; Zhang et al., 2020c). The diversity of MAC from different combustion sources implies the complex influences on light absorption from multiple factors, e.g., ambient oxidation, combustion efficiency and fuel types. The comparable and

diverse MAC between HOA and BBOA (Cappa et al., 2019; Zhong and Jang, 2014) warrants reevaluation of traffic aerosols' climate forcing and the necessity for clarifying MAC parameterizations for different sources.

3.2.3. Optical properties of SOA

Compared with the diurnal variation characteristics of BBOA and HOA from local emissions at YBJ site, light absorption coefficients of the less-oxidized oxygenated OA (LO-OOA) and more-oxidized oxygenated factor (MO-OOA) were characterized by high values during the day and low values at night (Fig. 5b, 5e). This day-enhanced temporal variation aligns with that of MO-OOA reported at NamCo (an observation station in central QTP) (Xu et al., 2018), which is influenced by the enhanced secondary BrC from photochemical oxidation during the day. In this study, the contribution of LO-OOA factor shows a broad increase to $Ab_{S_{BrC}, 370nm}$ during the day and peaks around 14:00 (20 %), while MO-OOA contribution (16 %) peaks approximately two hours later (16:00). The different diurnal variation of MO-OOA and LO-OOA suggests different formation pathways. In general, LO-OOA is characterized by freshly formed SOA showing a higher light absorption contribution (13 %) than that of MO-OOA (9 %), which more represents the regionally aging SOA (Figs. 5b, 5e). For the GIG site, the maximum direct light absorption contribution from SOA incorporated into case 2 (see Sect.2.5.1 and Text S1; Tables S4 and S6) is 33 % (Fig. S9).

The MAC of SOA (LO-OOA and MO-OOA) showed much lower values (0.15 and $0.18 \text{ m}^2 \text{ g}^{-1}$; Table 1, Fig. 6) than primary sources (e.g., BBOA and HOA), consistent with their higher degree of oxidation leading to the weaker light absorptivity of OA (Lee et al., 2014; Lambe et al., 2013). Lower MAC of SOA ($0\text{--}4 \text{ m}^2 \text{ g}^{-1}$) (Table S10) than POA ($0.4\text{--}8 \text{ m}^2 \text{ g}^{-1}$) has also been observed in multiple other ambient (De Sá et al., 2019; Retama et al., 2022; Tian et al., 2023; Washenfelder et al., 2015; Zhang et al., 2021; Zhang et al., 2022b) and laboratory studies (Ni et al., 2021; Hems et al., 2021; Zhong and Jang, 2014).

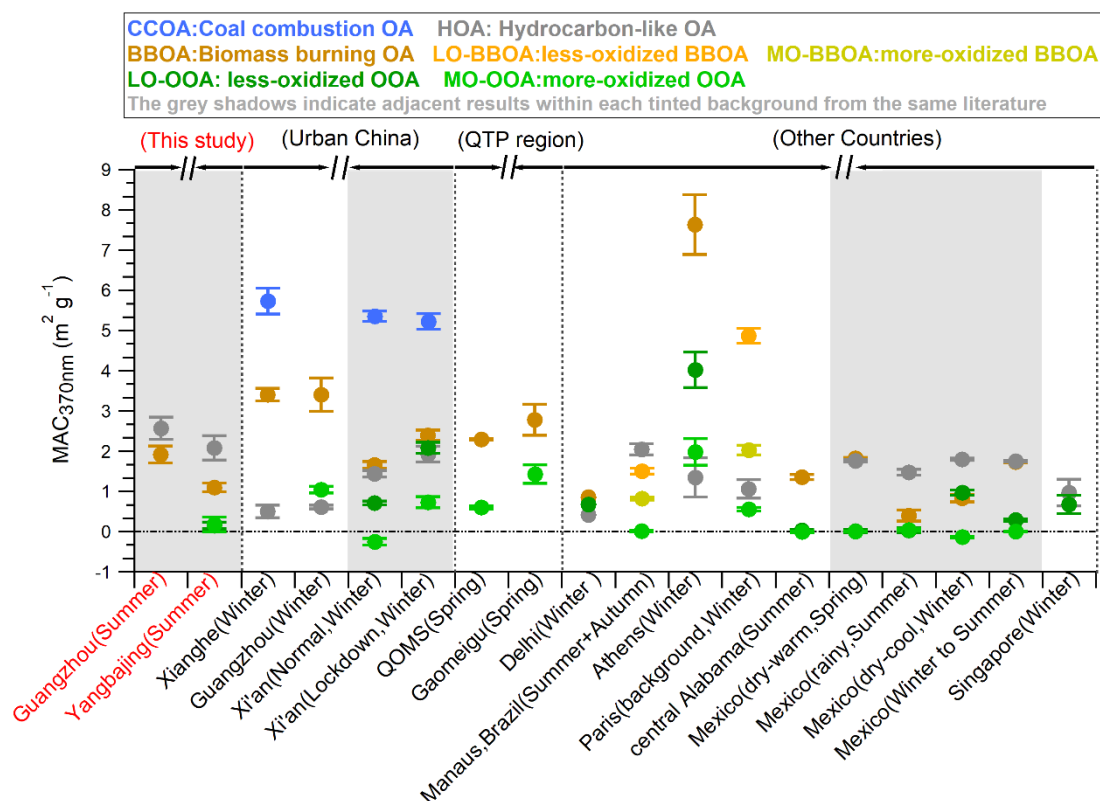


Figure 6. The literature summary of MAC from different BrC sources, which was obtained by the PMF–MLR method in different environments. All the results were categorized based on the locations of their observation sites (urban China, Qinghai–Tibet Plateau (QTP region), and other countries). Note the MO-OOA in Paris study represents OOA. Shadows indicate adjacent results within each tinted background from the same literature. The detailed information is provided in Table S10.

3.2.4. Summarized source contribution to BrC in the field studies

Figs. 7 (detailed data in Table S9) summarizes the light absorption contributions of OA from various sources across different regions, based on PMF and MLR analyses. Offline filter results are denoted by an asterisk in the pie chart. Based on this dataset, several features were found: 1) POA (primary organic aerosol) significantly influence ambient BrC light absorption on a global scale, accounting for 30–95 % of the BrC light absorption except the extremely low fraction (8.3 %) in Xi'an during summer (Lei et al., 2019). In particular, 3 quarters of the summarized study (based on number, as shown in Fig. S14) show that the POA contributes more than 50 % of BrC light absorption, signifying the important contribution of primary emission to total BrC. In the recent global model, Li et al. (2025) also found the primary emission dominated (77 %) the total light absorption from Nitrogen compounds. Note that the offline studies may underestimate the light absorption contribution of POA due to the application of soluble BrC (Bao et al., 2022; Chen et al., 2020; Lei et al., 2019; Zhong et al., 2023); 2) Key POA sources of BrC light absorption include coal combustion, biomass burning, and traffic emissions, with their light absorption contributions relative to total OA of 1–56 % (average: 33 %), 6–85 % (average: 38 %) and 4–83 % (average: 27 %), respectively (Fig. 7 and Table S9); 3) Coal combustion is especially evident in northern China during winter, with light absorption contribution of 30–89 % relative to POA (Lei et al., 2019; Sun et al., 2021; Wang et al., 2019b; Zhang et al., 2022b). In areas not subject to coal combustion, the biomass burning and traffic emissions dominate primary BrC light absorption, which can contribute

11–100 % (53 % for average) and 5–98 % (43 % for average) of primary BrC, respectively (Retama et al., 2022; De Sá et al., 2019; Kaskaoutis et al., 2021; Kasthuriarachchi et al., 2020a; Singh et al., 2021; Washenfelder et al., 2015; Zhang et al., 2020c). 4) The light absorption contribution from different sources showed significant spatial and temporal differences, e.g., HOA shows extremely high light absorption contribution in Singapore during summer (83 %), and in Mexico in both winter (54 %) and summer (49 %), while biomass burning dominated in central Alabama (85 %) and Paris (74 %). Given the pronounced spatial and temporal variations in source-specific light absorption, it is essential to conduct region-specific and season-specific observational research for a better understanding of the BrC sources and to better understand BrC sources and to better validate model simulations.

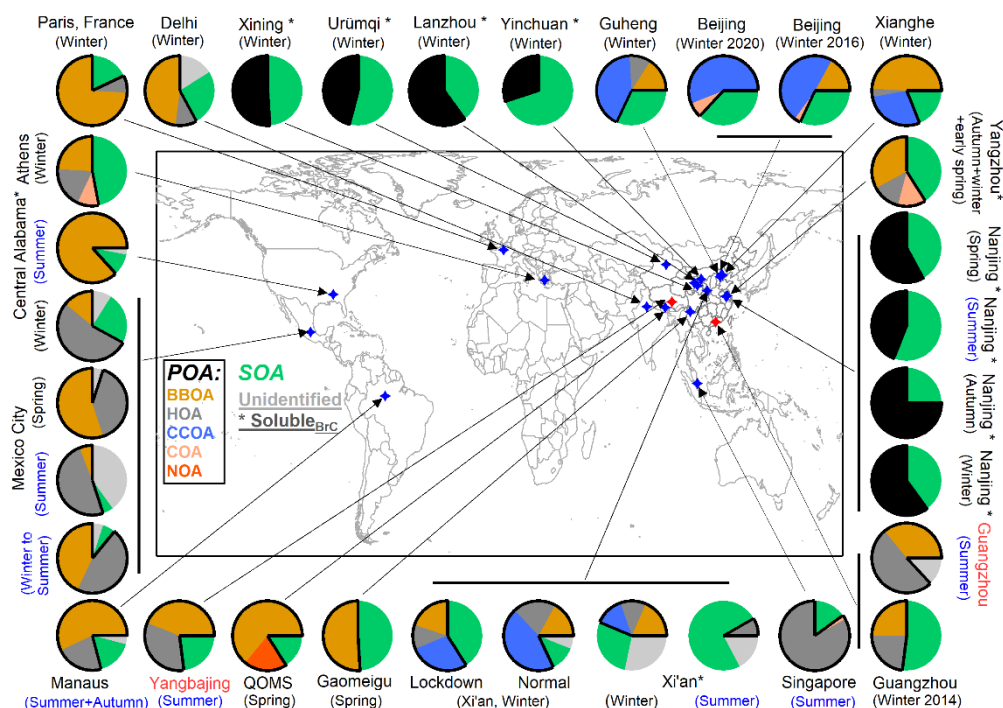


Figure 7. The summary of the contribution to BrC light absorption at 370 nm from different sources using the PMF-MLR method. The asterisks (*) represent the light absorption contributions of soluble BrC from different sources at 365nm. The sources include POA: biomass burning OA (BBOA), hydrocarbon-like OA (HOA), coal combustion OA (CCOA), cooking-related OA (COA), nitrogen-containing OA (NOA); The total SOA from oxygenated OA was used here. In addition, the unidentified fraction from the intercept of the MLR method was also shown. The detailed information on each pie is provided in Table S9.

3.3. Radiative effect of BrC from different sources

The simple forcing efficiency (SFE) of different organic aerosol (OA) components at the YBJ and GIG sites was estimated using Eq. (4) as described in Sect.2.6. This methodology has been widely applied to evaluate the climate impact of brown carbon (BrC) (Tian et al., 2023; Wang et al., 2019b; Zhang et al., 2022b; Zhong et al., 2023). As illustrated in Fig. 8, black carbon (BC) emerged as the dominant light-absorbing component, exhibiting the highest integrated SFE values of 294.7 W g⁻¹ at the YBJ site and 286.3 W g⁻¹ at the GIG site. In contrast, the integrated total SFE (from total OA absorption) across the 370–660 nm wavelength range was 21.4 W g⁻¹ at the YBJ site and 33.8 W g⁻¹ at the GIG site, representing approximately 9–13 % of the BC values. This is consistent with the well-documented strong light-absorbing properties of BC (Gustafsson and Ramanathan, 2016), while also highlighting the significant

role of BrC in short-term climate effects.

The SFE values for POA at the YBJ (19.2 W g^{-1}) and GIG (33.8 W g^{-1}) sites were comparable to those observed in other urban areas, such as Xi'an (approximately 33 W g^{-1} for HOA + BBOA + CCOA) during winter (Zhang et al., 2022b). POA, including BBOA and HOA, contributed significantly to the total BrC light absorption, accounting for over 80 % of the BrC-specific SFE in this study. These findings underscore the substantial influence of anthropogenic emissions on aerosol radiative forcing and their implications for regional and global climate systems.

From an environmental and climatic perspective, these results emphasize the critical role of both BC and BrC in modulating atmospheric radiative balance. The higher SFE of BC underscores its potent warming effect, while the non-negligible contribution of BrC, particularly from POA, highlights the importance of addressing anthropogenic sources. Further elucidating the fossil and non-fossil contribution, i.e., vehicle emission, coal vs. biomass burning combustion to the BrC, is vital for refining climate models, informing mitigation strategies, and developing policies aimed at reducing short-lived climate pollutants to mitigate near-term climate change and its associated environmental impacts.

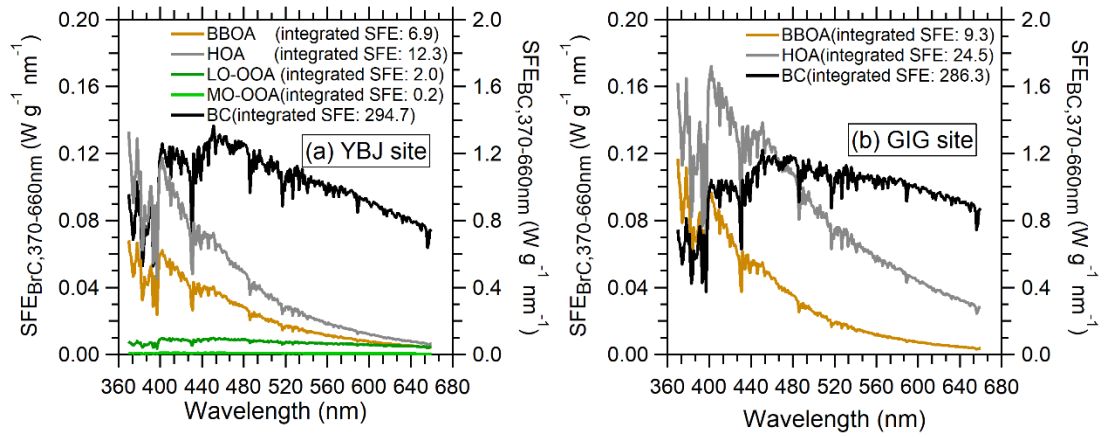


Figure 8. The Simple forcing efficiency (SFE) of BrC from different sources and BC from 370 to 660 nm at (a) YBJ and (b) GIG site.

4. Conclusions

To explore the optical properties, source contributions, and radiative effects of BrC in Qinghai-Tibet Plateau, the observations equipped with the AE33 and the SP-AMS were carried out in July 2022 in the Tibet background site (Yangbajing). For comparison, a simultaneous field observation was also conducted at the same periods in Guangzhou, a megacity with significant anthropogenic emissions. Our results reveal that the light absorption coefficient at 370nm from total aerosols (Ab_{total} , $1.6 \pm 1.6 \text{ M m}^{-1}$) and BrC (Ab_{BrC} , $0.2 \pm 0.3 \text{ M m}^{-1}$) in Tibet were approximately an order of magnitude lower than those in Guangzhou (Ab_{total} , $13.2 \pm 7 \text{ M m}^{-1}$; Ab_{BrC} , $2.9 \pm 2 \text{ M m}^{-1}$), consistent with the extremely low mass concentrations of total aerosols and organic aerosols (OA) observed at the Tibetan site. The minimal aerosol loading and weak light absorption in Tibet underscore its pristine atmospheric background.

Despite a large discrepancy in absolute light absorption coefficient, the BrC light absorption contribution to total aerosols (15 % and 21 %) is comparable between the two sites at 370 nm, highlighting BrC's significant contribution to total aerosol light absorption in Tibet. The summarized field studies demonstrate a positive correlation between BrC fraction in Ab_{total} as a function of POA/OA

($R = 0.54$), suggesting primary emissions contribute more effectively to BrC light absorption than BC, and POA is a more important source of BrC light absorption than SOA at these two sites. Source apportionment via the PMF–MLR method identified biomass–burning OA (BBOA) and hydrocarbon–like OA (HOA) from vehicles as the major contributors ($> 80\%$) to the BrC light absorption in both sites. The summarized field studies show that POA can account for 30–95 % of the BrC light absorption on a global scale. All these finding signifies the critical role of primary emission in OA light absorption. The main primary source for BrC includes biomass burning/biofuels, coal combustion, and vehicle emissions.

Diurnal variations of BrC and its sources exhibited distinct patterns with morning and nighttime peaks. Weisang activity and traffic rush hours in Tibet significantly influenced BrC levels, underscoring even remote regions' vulnerability to anthropogenic activities. Notably, traffic–related BrC contributions remain substantial in Guangzhou and even in the Tibet background, which necessitates that explicit light absorption parameters for fossil-derived BrC (e.g., from vehicle emissions) be considered in model simulations. Current literature remains inconclusive regarding MAC differences between HOA and BBOA, though fuel type, combustion efficiency, and aging effects critically influence BrC MAC, emphasizing the need for enhanced field measurements and parameterization of source-specific MACs.

Based on this field study, the integrated total SFE of BrC across the 370–660 nm wavelength range can account for approximately 7 % and 12 % of the BC (294.7 W g^{-1} in Tibet and 286.3 W g^{-1} in Guangzhou). In total, primary emission contributes over 98 % of the total SFE at both sites. These findings reinforce the urgency of controlling primary emissions and call for regional–specific parameterizations in climate models to improve assessments of BrC's radiative effects. In general, our study promotes the understanding of BrC dynamic variation and its sources at clean background Tibet and typical urban areas, emphasizing the strong influences of anthropogenic to radiative forcing.

Data availability

The data shown in the paper are available on request from the corresponding authors (weiwei.hu@gig.ac.cn and shanhuang_eci@jnu.edu.cn).

Author contributions

WH, SH, PY, NM, BY, MS designed the research. SH, PY, NM, BY, WZ, ZL, LL, TP, TF, JW conducted the field measurements. GZ and XB supported the AE33 instrument. WZ, LL, ZL, and YC analyzed the data. LL and SH supported the SP-AMS data analysis and OA source analysis for the Tibet campaign. WZ wrote the paper. WH, SH, ZL, TP, TF, JW, YC, GZ, XB, and XW reviewed and commented on the paper.

Competing interests.

The authors declare that they have no conflict of interest.

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1038