

1 Concurrent photochemical whitening and darkening of ambient 2 brown carbon

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13 **Abstract.** The light-absorbing organic aerosol (OA), known as brown carbon (BrC), has important radiative impacts, however
14 its sources and evolution after emission remain to be elucidated. In this study, the light absorption at multiple wavelengths,
15 mass spectra of OA and microphysical properties of black carbon (BC) were characterized at a typical sub-urban environment
16 in Beijing. The absorption of BC is constrained by its size distribution and mixing state, being subtracted from total absorption
17 to obtain the absorption of BrC, then by applying the least-correlation of BC absorption with secondary BrC, the absorption
18 contributed by BC, primary BrC and secondary BrC was apportioned. The multi-linear regression analysis on the factorized
19 OA mass spectra indicated the OA from traffic and biomass burning emission contributed to primary BrC. Importantly, the
20 moderately oxygenated OA (O/C=0.62) was revealed to highly correlate with secondary BrC. These OA had higher nitrogen
21 content, in line with the nitrogen-containing functional groups detected by the Fourier transform infrared spectrometer. The
22 photooxidation was found to result in reduced contribution of primary BrC about 20% but enhanced contribution of secondary
23 BrC by 30%, implying the concurrent whitening and darkening of BrC. This provides field evidence that the photochemically
24 produced secondary nitrogen-containing OA can considerably compensate some bleaching effect on the primary BrC, hereby
25 causing radiative impacts.

26 1. Introduction

27 Atmospheric absorbing organic aerosol (OA), known as brown carbon (BrC), is important contributor to anthropogenic
28 absorption besides black carbon (BC) (Laskin et al., 2015; Liu et al., 2020), particularly at shorter visible wavelengths (Bahadur
29 et al., 2012). Due to complex compositions of OA, the primary sources and subsequent evolution of BrC in the atmosphere
30 remains to be explicitly understood and causes uncertainties in evaluating the radiative impacts of BrC (Liu et al., 2020).

31 The chromophores of BrC are mainly aromatic compounds associated with certain functional groups (Liu et al., 2015c).
32 Particularly, compounds containing nitro, nitrated or other forms of nitrogen-containing functional groups are more absorbing
33 (Nakayama et al., 2013; Jacobson, 1999). It is well established that primary OA, especially from biomass burning, contains a
34 large fraction of BrC (Andreae and Crutzen, 1997; Rizzo et al., 2013; Bond, 2001). These primary BrC had a range of
35 absorptivity, which was found to be controlled by burning phases, with OA co-emitting with BC (the flaming phase) exhibiting
36 a higher absorptivity than OA-dominated smoldering phase (Liu et al., 2021). BrC can experience reactions with atmospheric
37 oxidants after emission. Previous studies (Satish et al., 2017; Satish and Rastogi, 2019) found nitrogenous compounds from
38 biomass burning were responsible for brown carbon over South Asia and the chromophores were photobleached in the
39 afternoon. Numerous field and laboratory studies found the decrease of BrC absorptivity due to photobleaching on
40 chromophores, with lifetime ranging from a few hours (Zhao et al., 2015; Liu et al., 2021) to a few days (Forrister et al., 2015),
41 which may depend on the concentration of ambient hydroxyl radical (Wang et al., 2014), also influenced by relative humidity
42 and particle volatility (Schnitzler et al., 2020). The absorptivity of BrC could be also enhanced due to addition of functional
43 groups by forming conjugated structure with aromatics. This was supported by a number of laboratory studies that BrC
44 absorptivity could be enhanced when forming nitrogen-containing organic compounds, such as the formation of nitro-
45 aromatics when aromatics reacted with NO_x (Nakayama et al., 2013), or produced organic amine after reacting with ammonia
46 (Updyke et al., 2012). The enhancement of BrC absorptivity could occur either through nitration of existing chromophores, or
47 formation of new secondary organic aerosol (SOA) chromophores through gas-phase oxidation.

48 The above findings mean the enhancement or bleaching of BrC absorptivity via photooxidation will coexist. The time scale
49 between both competing processes will ultimately determine the lifetime of BrC in the atmosphere. However, both processes
50 have been rarely investigated in the field to explicitly determine the BrC components which principally determine the
51 respective enhancement or decrease of its absorptivity, particularly in regions influenced by combined anthropogenic sources.
52 In this study, by measurements using multiple-wavelength absorption and microphysical properties of BC in a sub-urban region,
53 the absorption of BC, primary and secondary BrC was discriminated. In conjunction with source attribution via OA mass
54 spectra, we are able to link the segregated absorption with certain sources and investigate their primary information and
55 subsequent evolution. The competition between photobleaching and secondary formation of BrC was investigated in real world.

56 2. Experimental and instrumentation

57 2.1 Site description and meteorology

58 The experiment was conducted during springtime at the Beijing Cloud Laboratory and Observational Utilities Deployment
59 Base (117.12°E, 40.14°N), which is located in the northeast suburban area in Beijing (Fig S1a). The site is surrounded by the
60 northwest mountain ridge, without significant local primary anthropogenic emissions (Hu et al., 2021). The 72-h backward
61 trajectories with every 3 hours initializing from the site are analyzed by the HYSPLIT model (Draxier and Hess, 1998) using
62 the 3-hourly 1°×1° meteorological field from the GDAS reanalysis product. The obtained backward trajectories were further
63 clustered to group the similar transport pathways (Makra et al., 2011). The meteorological parameters, including the
64 temperature (T), ambient relative humidity (RH), wind speed (WS) and wind direction (WD) were measured by a monitoring
65 station on the site.

66 2.2 Measurements of BC microphysics and absorption coefficient

67 In this study, the ambient aerosols were sampled by a large-flow (1.05 m³ min⁻¹) air particle sampler (TH-1000C II) with a
68 PM_{2.5} impactor (BGI SCC 1.829) and dried by a silica drier before measurement. The single particle soot photometer (SP2,
69 DMT., USA) used continuous laser at $\lambda=1064\text{nm}$ to incandesce light-absorbing aerosols (such as BC) for irradiating detectable
70 visible light. The incandescence signal was used to measure the refractory black carbon (rBC) mass. The SP2 incandescence
71 signal was calibrated using the Aquadag standard (Acheson Inc., USA), and a factor of 0.75 was applied to correct for ambient
72 BC (Laborde et al., 2012). The scattering signal was calibrated by monodispersed polystyrene latex spheres (PSL). The BC
73 core diameter (D_c) was calculated from the measured BC mass by assuming a BC density of 1.8 g cm⁻³ (Bond and Bergstrom,
74 2006). The leading edge only (LEO) method was applied to reconstruct the scattering signal of BC, which was used to
75 determine the coated particle diameter (D_p) by a Mie-lookup table with the inputs of scattering and incandescence signal of
76 each BC particle (Liu et al., 2014; Taylor et al., 2015). The mass median diameter (MMD) is derived from the D_c distribution,
77 which is determined as below and above MMD the rBC mass concentration is equal (Liu et al., 2019a). The bulk coating
78 thickness (D_p/D_c) is calculated as the cubic root of ratio of the total coated BC volume divided by the total volume of rBC.

79 The mass absorption cross section (MAC) (in m² g⁻¹) of each BC particle can be calculated using the measured coated and
80 uncoated BC sizes by applying the Mie core-shell calculation. The absorption coefficient of BC at certain wavelength, $\sigma_{\text{abs,BC}}$
81 (λ) is determined by multiplying the calculated MAC and rBC mass concentration at each size:

$$82 \sigma_{\text{abs,BC}}(\lambda) = \sum_i \text{MAC}(\lambda, D_{p,i}, D_{c,i}) m(\log D_{c,i}) \Delta \log D_{c,i} \quad (1)$$

83 where $m(\log D_{c,i})$ denotes the BC mass concentration at each logarithmic bin of D_c . The SP2 measurement at $\lambda=1064\text{nm}$ longer
84 than mostly populated BC size means the derived coatings and subsequent calculation of MAC is relatively independent of
85 particle shape within uncertainty of 21% (Liu et al., 2014; Hu et al., 2021).

86 The absorption coefficients at wavelengths $\lambda=375, 470, 528, 635$ and 880 nm were measured by a Micro-Aethalometer
87 (MA200, Aethlabs, San Francisco, CA, USA). Aerosol particles were collected on filter tapes, on which the light attenuation
88 was measured continuously with a time resolution of 30 s. The loading effect of filters was automatically corrected by

89 measuring attenuation at two different sampling flow rates on two spots in parallel (Drinovec et al., 2015). Moreover, a multi-
 90 scattering correction factor (C-value) of 3.5, 3.2 and 2.4 at the wavelengths 370 nm, 528 nm and 880 nm, respectively were
 91 utilized to correct attenuation for the multiple light scattering effect. It was obtained by comparing the absorption coefficient
 92 with a photoacoustic soot spectrometer (PASS-3, DMT) (Hu et al., 2021).

93 2.3 Attribution of primary and secondary BrC absorption coefficient

94 The absorption coefficient of BC at different λ is calculated using the measured uncoated core and coated size as mentioned
 95 above. The absorption coefficient of total BrC is obtained by subtracting the BC absorption coefficient from the total absorption
 96 at certain wavelength, expressed as:

$$97 \sigma_{\text{abs, BrC}}(\lambda) = \sigma_{\text{abs, total}}(\lambda) - \sigma_{\text{abs, BC}}(\lambda) \quad (2)$$

98 where the absorption coefficient of BC ($\sigma_{\text{abs, BC}}$) is obtained from the SP2 measurement, $\sigma_{\text{abs, total}}(\lambda)$ is the total light absorption
 99 of aerosols measured by the MA200. The absorption coefficient of secondary BrC, the absorption not contributed by primary
 100 sources, is obtained by subtracting the absorption of all primary sources from the total absorption (Crilley et al., 2015),
 101 expressed as:

$$102 \sigma_{\text{abs, secBrC}}(\lambda) = \sigma_{\text{abs, total}}(\lambda) - \sigma_{\text{abs, pri}}(\lambda) \quad (3)$$

103 where $\sigma_{\text{abs, pri}}(\lambda)$ is the light absorption from primary sources. Here an assumption is made that light absorption from primary
 104 aerosols is all from combustion sources, and these sources necessarily contain BC (Wang et al., 2018). Therefore, the total
 105 absorption from primary sources can be obtained by scaling a factor from the mass concentration of BC, expressed as:

$$106 \sigma_{\text{abs, pri}}(\lambda) = \left(\frac{\sigma_{\text{abs, total}}}{[\text{rBC}]} \right)_{\text{pri}} \cdot [\text{rBC}]$$

107 (4)

108 where [rBC] is the mass concentration of rBC measured by the SP2, $\left(\frac{\sigma_{\text{abs, total}}}{[\text{rBC}]} \right)_{\text{pri}}$ is the scaling factor to derive the
 109 absorption of primary combustion sources from [rBC]. This factor is obtained using the minimum R-squared (MRS) approach
 110 (Wu and Yu, 2016), by adjusting the factor until a minimum correlation between $\sigma_{\text{abs, secBrC}}$ and [rBC] is reached because the
 111 absorption from secondary sources are least likely to covary with that from primary sources (Wang et al., 2019a). This method
 112 has been used in urban and sub-urban environment to obtain the primary BrC associated with combustion sources. Different
 113 sources may exhibit different ratios of $\left(\frac{\sigma_{\text{abs, total}}}{[\text{rBC}]} \right)_{\text{pri}}$, however there were no sporadic pollution events during the experimental
 114 period, uniform sources are therefore considered, and this ratio tends to represent a mean for the experiment. Being different
 115 from previous studies, an auxiliary characterization of rBC mass measured by the SP2 is used here to avoid the possible
 116 interference from absorption measured by the same instrument. The $\left(\frac{\sigma_{\text{abs, total}}}{[\text{rBC}]} \right)_{\text{pri}}$ ratio at $\lambda=375$ nm, 470 nm, 528 nm, 635
 117 nm and 880 nm is calculated to be 20.7, 17.0, 14.4, 11.7 and 5, respectively (Fig. S2), which falls within the reported values
 118 from previous studies 11-50 (Zhang et al., 2020; Wang et al., 2019a). This scenario assumes a relatively consistent absorption

119 relative to BC mass concentration from sources during experiment. This however may not include some sporadic events when
120 sources with distinct OA or BC mass fraction may be introduced and alter the single $\left(\frac{\sigma_{abs,total}}{[rBC]}\right)_{pri}$ ratio. The $\sigma_{abs,secBrC}$
121 therefore represents the overall mean value during the experimental period but this ratio will vary with seasons and locations.
122 The σ_{abs} of primary BrC can then be calculated as:
123
$$\sigma_{abs,priBrC}(\lambda) = \sigma_{abs,BrC}(\lambda) - \sigma_{abs,secBrC}(\lambda) \quad (5)$$

124 where $\sigma_{abs,BrC}$ and $\sigma_{abs,secBrC}$ is calculated from Equation (2) and (3), respectively. When evaluated the contribution of
125 photooxidation to the absorption coefficient, the absorption coefficient during photooxidation (12:00-17:00) was divided by
126 the respective average absorption coefficients of primary and secondary BrC.

127 **2.4 Composition measurement**

128 The mass concentration and chemical composition of non-refractory sub-micron PM (NR-PM₁) including organic aerosols
129 (OA), nitrate (NO₃⁻), sulfate (SO₄²⁻), chloride (Cl⁻) and ammonium (NH₄⁺) were determined with a High-Resolution Time-of-
130 Flight Aerosol Mass Spectrometer (HR-ToF-AMS, Aerodyne Research Inc., USA). The setup, operation, and calibration
131 procedures of the AMS have been described elsewhere (Canagaratna et al., 2007). During this field observation, the AMS was
132 operated in V-mode for the quantification of mass concentrations. The composition-dependent collection efficiencies were
133 applied (Middlebrook et al., 2012), and the ionization efficiency was calibrated using 300 nm pure ammonium nitrate (Jayne
134 et al., 2000). Elemental ratios of OA including oxygen-to-carbon (O/C), hydrogen-to-carbon (H/C) and nitrogen-to-carbon
135 (N/C) were determined to the improved-ambient method (Canagaratna et al., 2015).
136 Positive Matrix Factorization (PMF) (Paatero and Tapper, 1994) was performed on the inorganic and organic high-resolution
137 mass spectra to distinguish OA components from different sources (Zhang et al., 2011; Ulbrich et al., 2009; Decarlo et al.,
138 2010). The mass spectra of the combined matrix for m/z < 120 were excluded in PMF analysis. Five OA factors were identified.
139 The diagnostics of PMF is summarized in Text S1 and Fig. S6.

140 **2.5 Offline Fourier transform infrared spectrometer (FTIR) analysis**

141 Particulate Matter (PM) samples were collected once a day onto prebaked (600°C, 4h) quartz fiber filters (Whatman, QMA,
142 USA) using a large-flow (1.05 m³ min⁻¹) air particle sampler (TH-1000C II). The collected filter samples were stored in the
143 refrigerator at -20°C before analysis. The infrared spectra of collected samples were measured by a Fourier transform infrared
144 spectrometer (FTIR, Thermo Scientific, USA) equipped with an iD5 attenuated total reflectance accessory (diamond crystal)
145 to quantify the chemical functional groups over the wavenumbers range of 550-4000 cm⁻¹ with a resolution of 0.5 cm⁻¹. The
146 NO and NO₂ symmetric stretch in the FTIR spectra can characterize the functional groups associated with nitrogen-containing
147 organics (Coury and Dillner, 2008). Fig. S3 shows typical examples of FTIR spectra and the assigned functional groups for
148 the three pollution levels during experiment. The peak at 1110 cm⁻¹ corresponds to the background of the quartz fiber filter
149 overlapped with some X-H bending vibrations, which is subtracted for the following analysis. The characteristic organic nitrate

150 spectra appear at wavenumbers 860 cm^{-1} (NO symmetric stretch), 1280 cm^{-1} (NO_2 symmetric stretch) and $1630\text{-}1640\text{ cm}^{-1}$
151 (NO_2 asymmetric stretch) (Bruns et al., 2010). After baseline calibration, The FTIR peaks of 1630 cm^{-1} and 860 cm^{-1} are
152 integrated the absorption areas above the baseline. The summed integrated area of -NO and - NO_2 are hereby used to indicate
153 the nitrogen-containing organics. There was no discernable peak of carbonyl group for our infrared spectrum, and the peak of
154 OH at 2500 cm^{-1} - 3400 cm^{-1} for the carboxylic acid is not discernable neither, thus the influence of ketone and carboxylic acid
155 may be of less importance for our dataset.

156 3. Results and Discussion

157 3.1 Source attributed OA

158 The overview results are shown in Fig. S1. The organics dominated the aerosol compositions for most time, but occasionally
159 nitrate was the most abundant component (Fig. S1g). Note that the nitrate here may also include components containing in
160 organics besides ammonium nitrate. Backward trajectories (Fig. S1a-d) showed that the most abundant PM_{10} concentration was
161 associated with air masses transported in shorter distance from southern regions (C1), but the longer and faster northerly
162 transported air mass from cleaner north (C2) could dilute the concentrations.

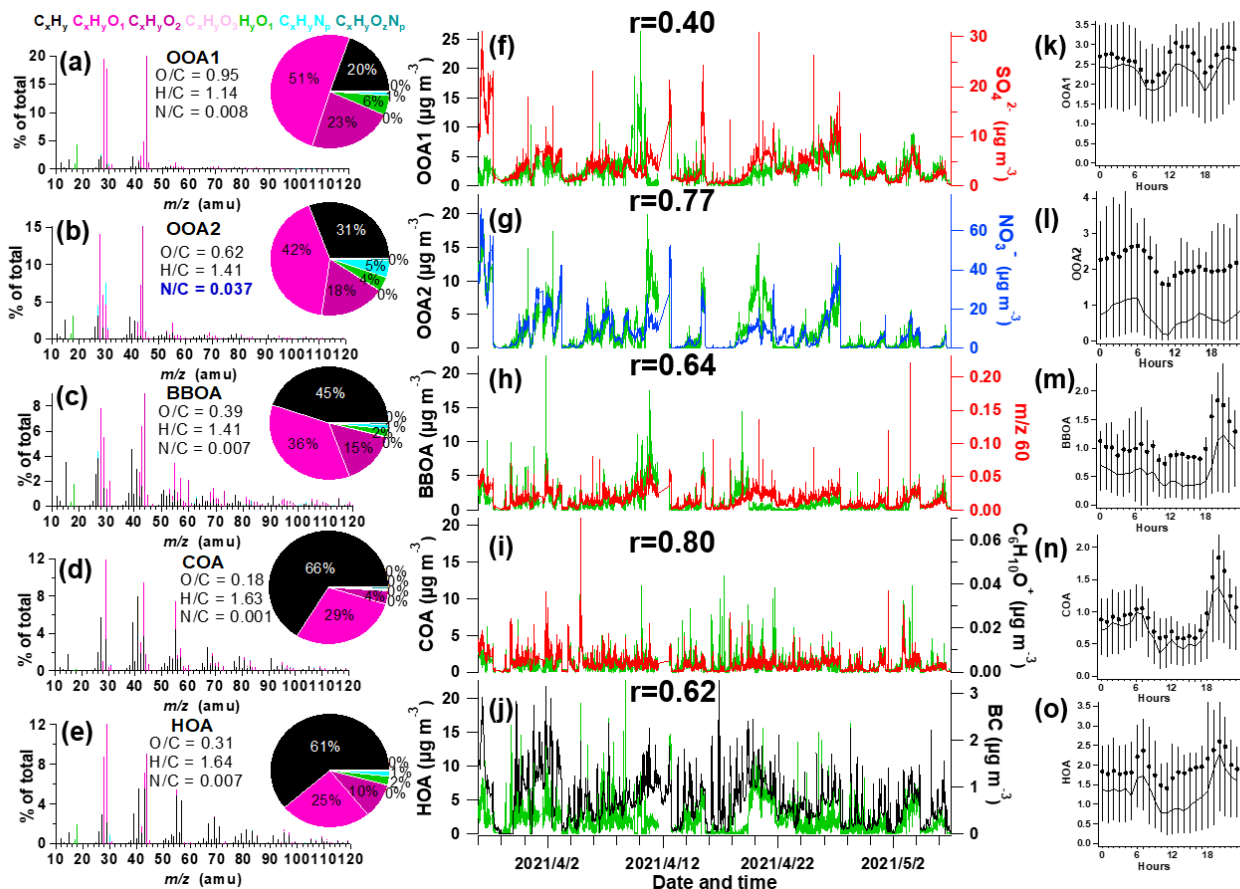
163 The resolved OA factors by the PMF analysis are shown in Fig. 1, including the mass spectra, time series and diurnal profiles
164 of each PMF factor with corresponded external and internal tracers. Three primary OA (POA) were identified as hydrocarbon-
165 like OA (HOA), cooking-related OA (COA), biomass burning OA (BBOA), with O/C of 0.31, 0.18 and 0.39 respectively.
166 These POA had considerable fraction of hydrocarbon fragments (C_xH_y), indicating their less aged status. The HOA profile was
167 characterized by higher contributions of aliphatic hydrocarbons and has dominated ion tracers such as m/z 41 (C_3H_5^+), 43
168 (C_3H_7^+), 55 (C_4H_7^+) and 57 (C_4H_9^+). The HOA concentration correlated with BC ($r=0.62$), which emits from traffic emissions.
169 The diurnal variation exhibited strong morning and afternoon rush-hour peaks of mass concentration. This factor was consistent
170 with the mass spectra of previously measured HOA from on-road vehicle emissions in urban cities (Zhang et al., 2005; Aiken
171 et al., 2009; Sun et al., 2016; Hu et al., 2017), which has m/z peaks characteristic of hydrocarbon fragments in series of $\text{C}_n\text{H}_{2n+1}^+$
172 and $\text{C}_n\text{H}_{2n-1}^+$. The mass spectrum of HOA shows overall similarity to those of primary OA emitted from gasoline and diesel
173 combustion sources ($r=0.68$) (Elser et al., 2016).

174 The OA from cooking sources (COA) is also characterized by prominent hydrocarbon ion series, however, with higher signal
175 at $\text{C}_n\text{H}_{2n-1}^+$ than $\text{C}_n\text{H}_{2n+1}^+$. COA had apparent fragments of both C_4H_9^+ and $\text{C}_3\text{H}_3\text{O}^+$, and has a higher ratio of $\text{C}_3\text{H}_3\text{O}^+/\text{C}_3\text{H}_5\text{O}^+$
176 (3.1), $\text{C}_4\text{H}_7^+/\text{C}_4\text{H}_9^+$ (2.2) than HOA (0.9–1.1), with cooking-related fragments of $\text{C}_5\text{H}_8\text{O}^+$ (m/z 84), $\text{C}_6\text{H}_{10}\text{O}^+$ (m/z 98) and
177 $\text{C}_7\text{H}_{12}\text{O}^+$ (m/z 112) (Sun et al., 2011b; Mohr et al., 2012). The COA shows overall similar spectral pattern to the reference
178 spectra of COA ($r=0.92$) (Elser et al., 2016). Its minor peak at noon and larger peak in the evening (Fig. 11) also corresponded
179 with the lunch and dinner time respectively. There was only a minor peak at noon for COA, which may be due to the sub-urban
180 nature of the site where the major aerosols from cooking sources may have been processed and lost the signature near source.

181 The feature of this factor was also observed in sub-urban environment (Huang et al., 2021).
182 The BBOA factor was identified based on the prominent signals of m/z 60 ($C_2H_4O_2^+$) and 73($C_3H_5O_2^+$), which are known
183 fragments of levoglucosan (Cubison et al., 2011). And BBOA also correlated with potassium (K^+ , $r = 0.80$), which are indicator
184 of biomass burning (Pachon et al., 2013; Brown et al., 2016). The m/z 60 and 73 together with a unique diurnal variation have
185 been shown to be a robust marker for the presence of aerosols from biomass burning emissions in many urban locations (Sun
186 et al., 2016). The BBOA shows very similar mass spectral patterns to previously reported reference spectra of biomass burning
187 ($r=0.94$) (Elser et al., 2016). The BBOA factor that was identified in spring accounted for 12.8% of the total OA in Beijing,
188 similar to previous reports (Hu et al., 2017). Biomass (Cheng et al., 2013) and solid fuel burning emissions (Sun et al., 2014)
189 have been widely observed to importantly contribute to the primary OA in this region. This off-road combustion source was
190 particularly abundant during wintertime for residential heating activities (Shen et al., 2019; Yang et al., 2018; Liu et al., 2016),
191 while boiler for industry use (mostly using coal as fuel) was in operation throughout the year (Liu et al., 2015b). During the
192 springtime of the experiment, the residential heating activities dropped due to increased ambient temperature thus the BBOA
193 may be mainly contributed by the industry sector.

194 Two types of oxygenated organic aerosols (OOA) were identified, in moderate (OOA2, $O/C=0.62$) and high oxidation state
195 (OOA1, $O/C=0.95$), respectively, which is very similar to the spectra of OOA factors resolved in other cities (Hayes et al.,
196 2013; Ulbrich et al., 2009). The average mass spectrum of OOA2 in this study is characterized by m/z 29 (mainly CHO^+), 43
197 (mainly $C_2H_3O^+$) and m/z 44 (CO_2^+), similar to the semi-volatile OOA spectrum identified in other locations (Sun et al., 2011a;
198 Zhou et al., 2016). On average, OOA2 accounts for 42% and 18% of $C_xH_yO^+$ and $C_xH_yO_2^+$ ions, respectively (Fig. 1b). These
199 results clearly indicate that OOA2 was primarily composed of less oxygenated, possibly freshly oxidized organics. Notably,
200 OOA2 had a substantially higher N/C than other factors ($N/C=0.037$), and had highest correlation with nitrate ($r=0.77$) and
201 with $C_xH_yN_z$ and $C_xH_yN_zO_p$ fragments ($r=0.83$). This factor therefore tends to largely result from nitrogen-containing OA and
202 its elevation at night may be also associated with dark oxidation by nitrate radical.

203 The mass spectrum of OOA1, which was characterized by a dominant peak at m/z 44 (mainly CO_2^+), a highest O/C (0.95). On
204 average, OOA1 contributes 51% of the $C_xH_yO^+$ signal and 23% of the $C_xH_yO_2^+$ signal (Fig. 1a). OOA1 showed particularly
205 high correlation with sulfate ($r=0.40$) because of their similar volatilities (Huffman et al., 2009; Jimenez et al., 2009). The
206 slight enhancement at noon for OOA1 (also for OOA2) soon after morning rush-hour indicated the likely rapid formation of
207 SOA through photooxidation. This significantly higher mean OOA2 than median value in the diurnal pattern indicated that
208 this OA type was largely associated with pollution events. Both OOA1 and OOA2 showed nighttime peak maybe due to
209 reduced boundary layer.

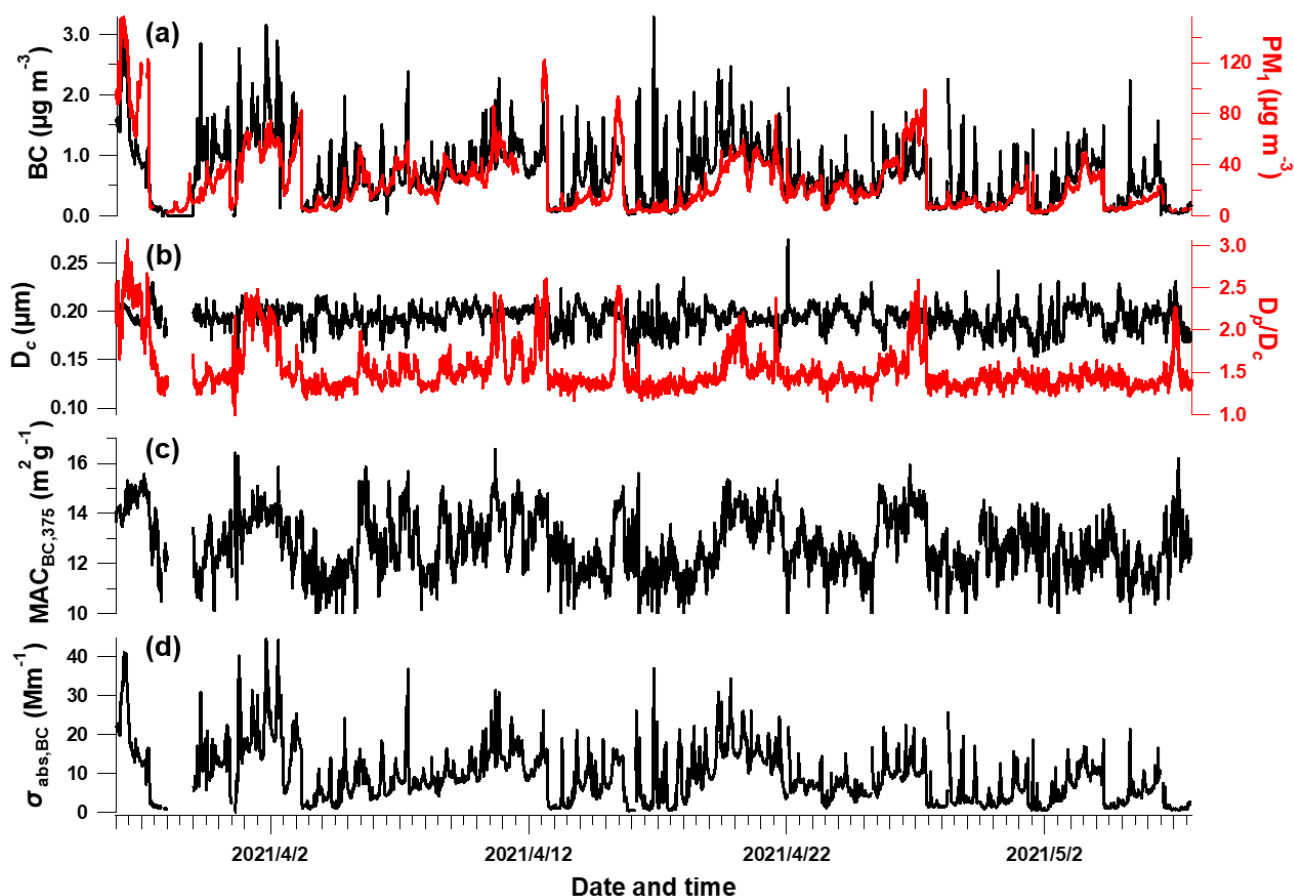


210
 211 **Figure 1. Information of source-apportioned organic aerosols by the PMF analysis. Mass spectra of (a) oxygenated OA1 (OOA1), (b)**
 212 **oxygenated OA2 (OOA2), (c) biomass burning OA (BBOA), (d) cooking-related OA (COA), (e). hydrocarbon-like OA (HOA), (f-j)**
 213 **Temporal variations of each PMF factor and the corresponding marker species. (k-o) Diurnal profiles of each factor. The lines, dots**
 214 **and whiskers denote the median, mean and the 25th/75th percentiles at each hour respectively.**

215 3.2 Segregated aerosol absorption

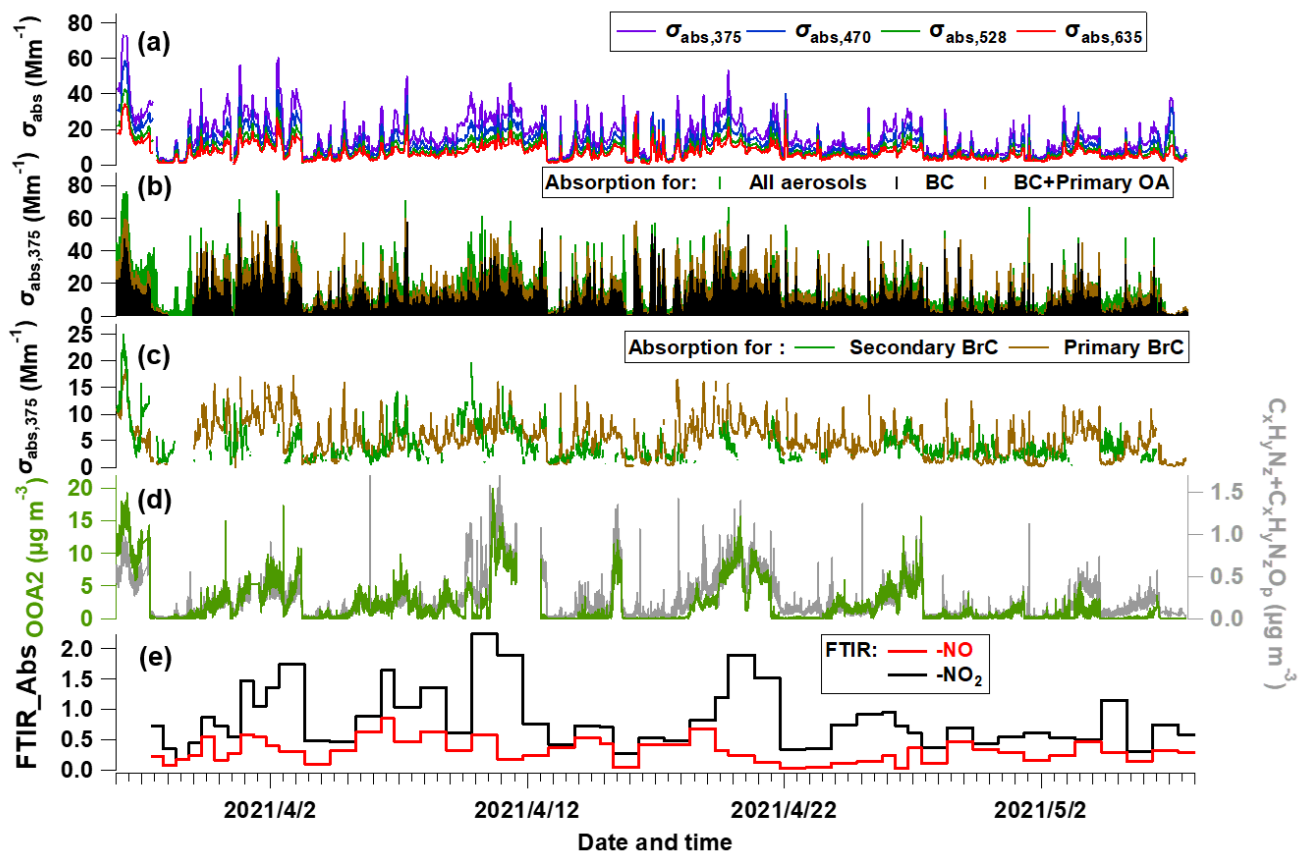
216 Fig. 2 shows the time series of BC properties, including the BC mass concentration, D_p/D_c , D_c , MAC and light absorption
 217 coefficient of BC (section 2.2). The MMD of BC core varied between 93 – 274 nm which may correspond to the source-
 218 specific information (Liu et al., 2019b) or coagulation process during ageing. The coating of BC (indicated by D_p/D_c) showed
 219 sporadic enhancement which was closely associated with enhanced PM concentration (Fig. 2a). This was consistent with
 220 previous studies that high coatings of BC occurred during heavier pollution due to the enhanced secondary formation of
 221 condensable materials to particle phase (Ding et al., 2019; Zhang et al., 2018). This clearly indicates the variation of mixing
 222 state of BC and this will potentially influence its MAC and absorption Ångström exponent (AAE) (Liu et al., 2015a). It will
 223 introduce considerable uncertainties to use constant MAC or AAE to derive the absorption coefficient of BC at multiple
 224 wavelengths. The MAC estimated using the measured BC core size and coatings (Fig. 2c) is thus used to derive the $\sigma_{\text{abs,BC}}$
 225 (section 2.2, shown in Fig. 2d). The $\sigma_{\text{abs,BC}}$ was $9.1 \pm 7.3 \text{ Mm}^{-1}$ during experimental period. MAC of BC at $\lambda=375\text{nm}$ showed to

226 be at $8.4 - 16.6 \text{ m}^2 \text{ g}^{-1}$ with enhanced absorption when high coatings.



227
228 **Figure 2. Temporal evolution of BC-related properties. (a) rBC and PM₁ mass concentration, (b) BC core diameter and bulk coating**
229 **thickness (D_p/D_c), (c) calculated mass absorption cross section (MAC) at $\lambda=375\text{nm}$, (d) absorption coefficient of BC.**

230 Using the method above, the total ($\sigma_{\text{abs,total}}$) and attributed absorption of BC ($\sigma_{\text{abs,BC}}$), primary ($\sigma_{\text{abs,priBrC}}$) and secondary BrC
231 ($\sigma_{\text{abs,secBrC}}$) at $\lambda=375\text{nm}$ are shown in Fig. 3a-c. In Fig. 3b, the brown and green shades above the adjacent tracer indicate the
232 absorption coefficient of primary and secondary BrC, respectively. Fig. 3c shows that the absorption coefficient of primary
233 BrC was higher than secondary BrC for most time, but for certain periods they were equivalent or secondary BrC occasionally
234 exceeds primary BrC. The mean contribution of absorption coefficient for BC, primary BrC and secondary BrC is 51%, 27%
235 and 22% in this study. The tracers associated with nitrogen-containing organics, such as OOA2 (with highest N/C), $\text{C}_x\text{H}_y\text{N}_z$
236 and $\text{C}_x\text{H}_y\text{N}_z\text{O}_p$ fragments, and the FTIR measured $-\text{NO} + -\text{NO}_2$, are also shown in Fig. 3d-e.



237
 238 **Figure 3. Temporal evolution of segregated absorbing properties.** (a) Absorbing coefficients (σ_{abs}) at multiple wavelengths measured
 239 by the aethalometer, (b) σ_{abs} at $\lambda=375\text{nm}$ ($\sigma_{\text{abs},375}$) for all aerosols, primary OA and BC, (c) $\sigma_{\text{abs},375}$ for primary BrC and secondary
 240 BrC. (d) mass concentration of OOA2 and the $\text{C}_x\text{H}_y\text{N}_z$ and $\text{C}_x\text{H}_y\text{N}_z\text{O}_p$ fragments measured by the AMS. (e) FTIR-measured
 241 absorption of -NO and -NO₂ bonds.

242 3.3 Source attribution of BrC absorption

243 A multiple linear regression (MLR) analysis is performed to apportion the absorption coefficient of BrC with the PMF
 244 attributed OA factors, expressed as:

$$245 \sigma_{\text{abs,BrC}} = a_0 + a_1 \cdot [\text{OOA1}] + a_2 \cdot [\text{OOA2}] + a_3 \cdot [\text{BBOA}] + a_4 \cdot [\text{COA}] + a_5 \cdot [\text{HOA}] \quad (6)$$

246 where a_1 to a_5 represents the regression coefficients for each factor. The contribution of each source-specific OA factor to
 247 $\sigma_{\text{abs,BrC}}$ can be obtained. This analysis is performed for the total BrC, primary and secondary BrC respectively. The results are
 248 shown in Table 1. MLR on the total BrC shows high correlation ($r > 0.4$) with the factors of HOA, BBOA and OOA2, suggesting
 249 the potential importance of the primary biomass burning and traffic source along with OOA2 in governing absorption of BrC.
 250 MLR analysis on the primary BrC distinguishes its substantial correlation with BBOA ($r = 0.40$) and HOA ($r = 0.46$), while MRL
 251 on the secondary BrC has a high correlation with OOA2 only ($r = 0.44$). The MRL analysis links the apportioned absorption of
 252 physical properties with source-attributed chemical compositions, therefore validating and identifying the sources of primary

253 and secondary BrC.

254 **Table 1. Results of the multilinear regression analysis (MLR) between $\sigma_{\text{abs},375}$ and the five PMF-resolved OA factors, with $\sigma_{\text{abs},375}$ of**
255 **total BrC, primary and secondary BrC as dependent, respectively. All regression coefficients have passed the significance test with**
256 **$p < 0.01$. Partial correlations above 0.4 are marked in red. Since negative values appear when the COA participates, which is thus not**
257 **included in the final regression but the values using COA factor are shown in brackets.**

Dependent	$\sigma_{\text{abs},\text{BrC}}$		$\sigma_{\text{abs},\text{pri BrC}}$		$\sigma_{\text{abs},\text{sec BrC}}$	
	Regression coefficient	Partial correlation	Regression coefficient	Partial correlation	Regression coefficient	Partial correlation
Constant	2.26		1.67		1.47 (1.52)	
OOA1	0.57	0.23	0.04	0.02	0.46(0.46)	0.24 (0.24)
OOA2	1.22	0.53	0.37	0.25	0.74 (0.74)	0.44 (0.44)
BBOA	2.59	0.46	1.22	0.40	1.14 (1.18)	0.29 (0.29)
COA	1.30	0.22	1.45	0.36	/ (-0.25)	/ (-0.05)
HOA	1.70	0.47	1.17	0.46	0.49 (0.52)	0.20 (0.21)
R ²		0.77		0.63		0.55 (0.55)

258 Importantly, an oxygenated secondary OA factor (OOA2) is identified to significantly contribute to the secondary BrC. This
259 OOA has a moderate O/C (0.62) and a highest N/C of 0.037 among all factors. The high N/C means this factor contains the
260 most abundant nitrogen-containing fragments, implied as its high correlation with the $\text{C}_x\text{H}_y\text{N}_z$ and $\text{C}_x\text{H}_y\text{N}_z\text{O}_p$ fragments ($r=0.83$,
261 Fig. 3d) and with the FTIR absorption for $-\text{NO}_2$ and $-\text{NO}$ bonds ($r=0.69$, Fig. S4). The $-\text{NO}$ bond is mostly related to the
262 organic nitrates (RONO_2), and $-\text{NO}_2$ peak could result from both organic nitrates and nitro-organics (Bruns et al., 2010). There
263 is no discernable peak for organic amines. These all consistently imply that the OOA2 factor contained substantial fraction of
264 nitrogen-containing organics, and these compounds have contributed to the absorption of secondary BrC.

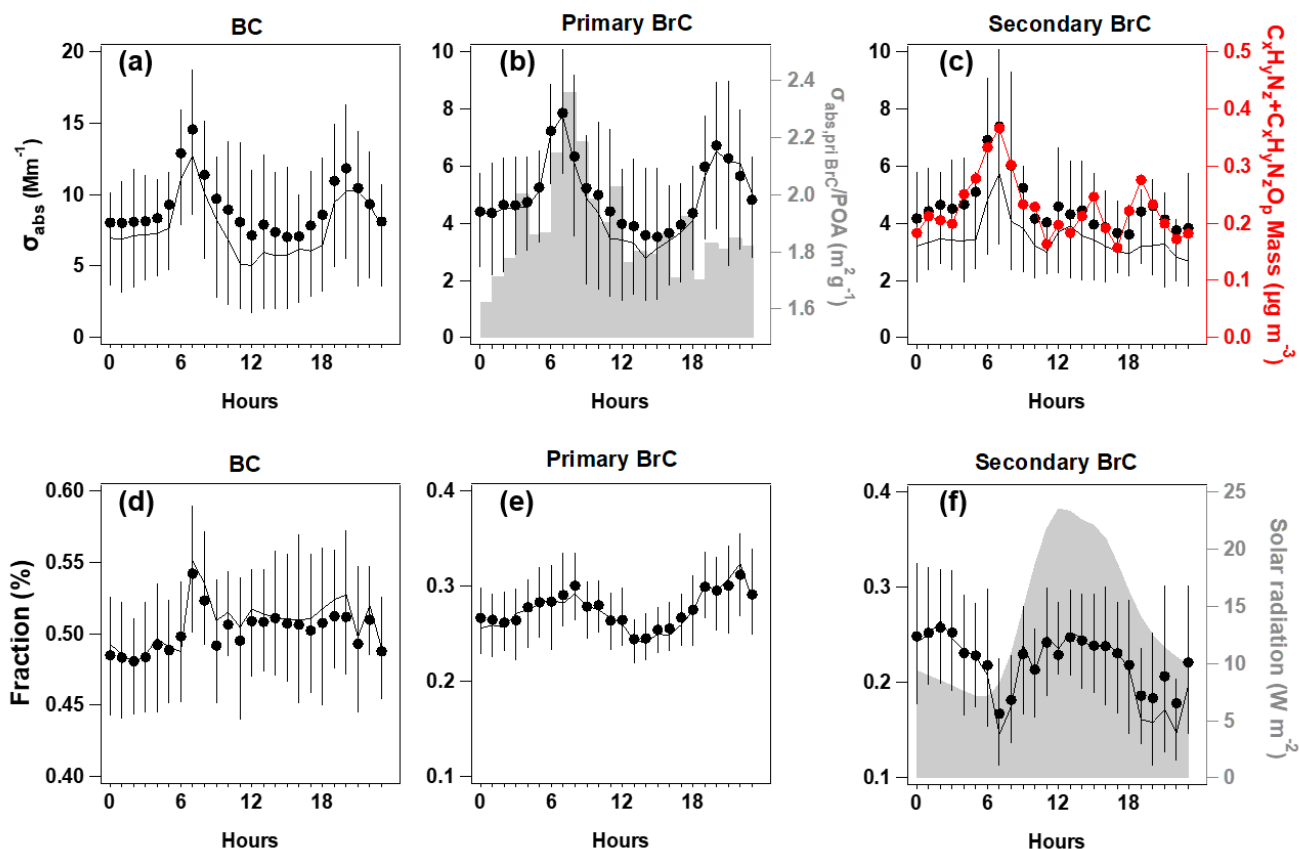
265 3.4 Simultaneous whitening and darkening process of BrC

266 The diurnal variation of $\sigma_{\text{abs},375}$ for BC and primary BrC and their fractions showed consistent morning rush-hour peaks at
267 6:00-8:00 and the night-time enhancement due to reduced boundary layer (Fig. 4a-b). This was in line with the morning peak
268 of HOA and night peak of BBOA. The traffic source in this region, in particular the diesel vehicles, was reported to emit
269 considerable OA with certain chromophores, such as aromatics (Yao et al., 2015) and heterocyclic organic compounds (Gentner
270 et al., 2017; Schuetzle, 1983). In the morning rush-hour, BC and primary BrC accounted for $51 \pm 4\%$ and $29 \pm 4\%$ in the total
271 $\sigma_{\text{abs},375}$ respectively, with the remaining $20 \pm 2\%$ classified as secondary BrC. The morning peak coinciding with the primary
272 BrC may result from the rapid formation of BrC from sources when emitted gases condensed and formed aerosols. These may
273 lead to high cooccurrence between primary and secondary BrC. Previous studies in urban environment also observed

274 concurrent peaks of primary and secondary BrC, which usually occurred at morning rush hour (Zhang et al., 2020). The night
275 had contributions from BC and primary BrC at $50\pm 2\%$ and $30\pm 3\%$ respectively, with $20\pm 3\%$ as secondary BrC. Fig. 4b showed
276 the decrease of primary BrC absorption tended to be more rapid than the HOA and BBOA mass (even a slight increase for
277 HOA Fig. 1m and Fig. 1o), leading to decreased absorption coefficient per unit mass of primary BrC (shade in Fig. 4b), which
278 indicates the photobleaching process. In this context, a recent chamber study reported that the primary BrC from biomass
279 burning plumes could be bleached to half of the initial absorptivity in 2-3 hours (Liu et al., 2021).

280 Besides the morning rush-hour peak, there was an early afternoon peak for the absorption coefficient of secondary BrC,
281 prevailing the dilution effect of daytime boundary layer (Fig. 4c-S5). The night and morning peak of OOA2 and the morning
282 peak of $\sigma_{\text{abs,secBrC}}$ may result from primarily emitted moderately oxygenated OA, which was reported from some diesel sources
283 (Dewitt et al., 2015; Gentner et al., 2012). The fraction of secondary BrC thus had a pronounced early afternoon peak soon
284 after the peak solar radiation (Fig. 4f) and a peak after midnight soon after the nighttime peak of primary BrC (Fig. 4e). Fig
285 4e-f shows the photooxidation led to an enhanced contribution of secondary BrC by 30% but reduced contribution of primary
286 BrC about 20%. This shift of peaking time from primary to secondary BrC demonstrates the likely process of SOA formation
287 from gases, and these SOA compounds containing nitrogen (i.e., the OOA2) considerably contributed to the light absorption.
288 This ageing or oxidation likely occurred through photooxidation during early afternoon and aqueous processes (high RH
289 conditions prevail during nighttime) during nighttime. The oxidized volatile organic compounds (VOCs) with nitrogen
290 chemistry involved could condense to produce additional mass in particle phase (Ehn et al., 2014; Finewax et al., 2018). Due
291 to the high NO_x emission, photooxidation of traffic VOCs may have largely involved nitrogen chemistry. Previous studies
292 found the NO_x -involved SOA could produce considerable chromophores (Lin et al., 2015; Siemens et al., 2022), such as the
293 traffic VOCs may produce SOA in a time scale of hours, containing nitro-aromatics (Wang et al., 2019b; Keyte et al., 2016).
294 The daytime formation of organic nitrate may follow the gas-phase photooxidation mechanism, in which the excess NO could
295 add to the peroxy radical to produce organic nitrate (Liebmann et al., 2019). The nighttime chemistry involving NO_3 radical
296 through the oxidation of NO_2 by O_3 , contributed to the important formation of organic nitrate by initializing the production of
297 nitrooxy peroxy radicals (Ng et al., 2008; Rollins et al., 2012). Laboratory studies (Nakayama et al., 2013; Liu et al., 2015c)
298 also widely observed the rapid production of nitrogen-containing OA involving NO_x chemistry could contribute to light
299 absorption of aerosols.

300 Overall, by apportioning the absorption of primary and secondary BrC, we found the photooxidation led to an enhanced
301 contribution of secondary BrC by 30% but reduced contribution of primary BrC about 20% in the semi-urban environment.
302 This revealed that the whitening and darkening of BrC occurred simultaneously, and the secondary BrC produced by
303 photooxidation may compensate some bleaching effect of primary BrC. The dominance of both competing processes may
304 depend on the timescale and altitude in the atmosphere. For example, the enhanced BrC fraction observed above the planetary
305 boundary layer may be explained by the enhanced secondary BrC (Tian et al., 2020), while further ageing may bleach the
306 produced chromophores of these SOA.



307
 308 **Figure 4.** Diurnal variations of absorption coefficient at $\lambda=375\text{nm}$ ($\sigma_{\text{abs},375}$) for BC (a), primary BrC and absorption efficiency of
 309 primary BrC ($\sigma_{\text{abs,priBrC}}/\text{POA}$) is shown in shade (b), and secondary BrC, along with the $\text{C}_x\text{H}_y\text{N}_z$ and $\text{C}_x\text{H}_y\text{N}_z\text{O}_p$ fragments (c); the
 310 respective fraction in total for the segregated $\sigma_{\text{abs},375}$ (d-f), with direct radiation shown in shade. In each plot, the lines, dots and
 311 whiskers denote the median, mean and the 25th/75th percentiles at each hour respectively.

312 4. Conclusion

313 This study apportioned the shortwave absorption of BC, primary and secondary BrC, through concurrent measurements of BC
 314 microphysical properties and OA mass spectra. The apportioned primary BrC absorption was linked with traffic and biomass
 315 burning emissions, while secondary BrC was found to be associated with an oxygenated secondary OA factor with higher
 316 nitrogen content. The enhancement of secondary BrC and decrease of primary BrC simultaneously occurred via daytime
 317 photooxidation. The results emphasize the importance of nitrogen-containing OA in contributing to BrC. These OA could
 318 primarily emit as aerosol phase, or in gas phase which requires further oxidation to be in aerosol phase to serve as BrC. The
 319 NO_x -involved chemistry is prone to add nitrogen element to the existing OA and enhance the absorptivity of chromophores.
 320 The anthropogenic NO_x emission could be therefore an important source in producing shortwave absorbing components in the
 321 atmosphere, which may offset some of the conventionally-thought photobleaching of BrC.

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325 **Author contribution**

326 D.L., X.J. and Qian L. prepared and designed the observation. D.L., Qian L., X.J and P.T. initiated the field campaign and
327 conducted the measurements. Qian L., D.L. P.T., Y.W., S.L. and K.H. contributed to the data analysis. Qian L., H.M., L.R.,
328 B.K., D.D. and S.K. provided technical support and assistance. Qian L. and D.L. wrote the manuscript. All authors read and
329 approved the final manuscript.

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