

1 **Measurement report: Formation of tropospheric brown carbon in a**
2 **lifting air mass**

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30 **Abstract:** An enhanced formation of brown carbon (BrC) with a non-negligible warming effect
31 at the tropopause has recently been found. However, its formation mechanism is unclear. Here
32 we report a BrC formation process that happens during air mass upward transport by
33 conducting simultaneously a 4-hour time resolution of measurement on atmospheric BrC at the
34 mountain foot (MF, 400m a.s.l.) and mountainside (MS, 1120m a.s.l.) of Mt. Hua, China in
35 2016 summer. Our results showed that the daytime light-absorption (Abs_{365nm}) of BrC at MS is
36 approximately 60% lower than that at MF due to a dilution effect caused by the planetary
37 boundary layer expansion, but the daytime light-absorption of BrC relative to black carbon at
38 MS is about 30% higher than that at MF, suggesting a significant formation of secondary BrC
39 in the lifting process of air mass from MF to MS. Such a secondary formation accounted
40 for >50% of BrC at MS but only 27% of BrC at MF. Moreover, N:C elemental ratio of the
41 daytime BrC was 15% higher at MS than that at MF, mainly due to an aerosol aqueous phase
42 formation of water-soluble organic nitrogen (WSON) compounds. Stable nitrogen isotope
43 composition further indicated that such light-absorbing WSON compounds were produced
44 from the aerosol aqueous-phase reaction of carbonyls with NH_4^+ . Our work for the first time
45 revealed that ammonia-induced aerosol aqueous reactions can significantly promote BrC
46 formation during the air mass lifting process, which is probably responsible for an enhanced
47 light absorption of BrC in the upper boundary layer.

48 **Keywords:** Brown carbon; Ammonia and carbonyls; Nitrogen-containing organic compounds;
49 Air mass upward transport; Aerosol aqueous-phase reaction.

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51 **1. Introduction**

52 Light-absorbing organic aerosols, known as brown carbon (BrC), can efficiently absorb
53 solar radiation in the visible to near ultraviolet (UV) wavelength range (Laskin et al., 2015; Liu
54 et al., 2020a; Chakrabarty et al., 2023a) which is corresponding to 27~70% of black carbon
55 (BC) light-absorption in the lower troposphere (Saleh et al., 2015; Lin et al., 2014; Lin et al.,
56 2015) suggesting that BrC can perturb substantially the planetary radiation budget (Qian et al.,
57 2015; Lin et al., 2014; Liu et al., 2015). By absorbing solar radiation at short wavelengths, BrC
58 can strongly alter local gas-phase photochemistry and atmospheric oxidation through
59 decreasing the photolysis rates of OH radicals, NO₂ and O₃, leading to a reduction in
60 atmospheric oxidant concentration by up to ~30% (Hammer et al., 2016; Gligorovski et al.,
61 2015; Jo et al., 2016). BrC in the atmosphere also acts as photosensitizers and produces active
62 intermediates, and thus can promote sulfate formation (Liu et al., 2020b). In addition, BrC
63 comprises numerous organic species and can induce adverse human health effects, because
64 some of chromophores are toxic (Huang et al., 2018; Hsu et al., 2014; Yan et al., 2018).

65 Atmospheric BrC has both primary and secondary sources. Biomass burning is believed to
66 be the major source of primary BrC (Chakrabarty et al., 2023b), while emissions from fossil
67 fuel combustion is also an important source of primary BrC in the urban atmosphere (Yan et al.,
68 2017; Corbin et al., 2019), which accounts for even more than 40% of the total BrC in heating
69 season (Li et al., 2023). In the past decades numerous studies reported that BrC can also be
70 secondarily generated in the atmosphere, such as photooxidation of aromatics under high NO_x
71 conditions (Lin et al., 2015; Liu et al., 2021), NH₄⁺-initiated reactions with atmospherically
72 relevant carbonyls (Li et al., 2021b; Kampf et al., 2012; Laskin et al., 2014; Li et al., 2019b)

73 and $\cdot\text{OH}/\text{NO}_3\cdot$ radical oxidations of various VOCs (Sumlin et al., 2017; Gelencser et al., 2003;
74 Lu et al., 2011). BrC is chemically active, which may undergo photobleaching (Schnitzler et al.,
75 2022; Gilardoni et al., 2016), posing significant challenges for characterizing BrC molecular
76 composition and its links to optical properties.

77 Recently a aircraft measurement conducted over the continental United States observed an
78 enhanced short-wavelength optical absorption of BrC relative to BC at altitudes between 5 and
79 12 km (Zhang et al., 2017), indicating that secondary formation is one of crucial sources for
80 these high-altitude BrC. Numerical model studies reported that global radiative forcing caused
81 by BrC ranges from 0.1-0.6 Wm^{-2} (Zhang et al., 2020; Lin et al., 2014; Druge et al., 2022),
82 suggesting a non-negligible impact of BrC on the global climate change. Studies found that
83 such climate effects are highly sensitive to BrC and the sensitivity rapidly increases along with
84 an increase in altitude (Zhang et al., 2017; Nazarenko et al., 2017; Hodnebrog et al., 2014).
85 These abundant BrC at the tropopause would bring about prominent impacts on radiative
86 forcing, which is even twice of that induced by low-altitude BrC (Zhang et al., 2017). Due to
87 the limited number of field observations, however, the vertical distribution and formation
88 mechanism of tropospheric BrC are still unclear especially those in the upper troposphere,
89 where the direct radiative forcing of BrC is much stronger than that in the ground surface
90 atmosphere.

91 To elucidate the formation mechanism of BrC in the troposphere, synchronous
92 observations on atmospheric BrC were conducted on the mountainside and the mountain foot
93 of Mt. Hua, which is located closely to Guanzhong Basin, one of the areas with heaviest $\text{PM}_{2.5}$
94 pollution in China owing to intensive activities of fossil fuel combustion and the unfavorable

95 topography (Wang et al., 2022b; Wu et al., 2020; Wang et al., 2011; Wang et al., 2016). Our
96 previous study has shown that inorganic aerosol chemistry in the atmosphere over Mt. Hua is
97 dominated by the air mass transport from the Guanzhong Basin ground surface, in which
98 $(\text{NH}_4)_2\text{SO}_4$ is continuously produced during the air mass lifting process along with a decrease
99 in aerosol acidity (Wu et al., 2022). Here, we investigated the formation mechanism of
100 secondary BrC during the lifting process of air mass from the Guanzhong Basin to the
101 mountainous atmosphere of Mt. Hua. We firstly discussed the differences in vertical
102 distribution and optical absorption of water-soluble BrC between the ground surface and the
103 mountainous atmosphere, then explored their formation mechanism in the upper boundary
104 layer. To the best of our knowledge, we for the first time found that ammonia-induced aerosol
105 aqueous phase reaction with carbonyls is the dominant formation pathway of BrC in the air
106 mass lifting process, which is responsible for the high ratio of BrC to BC in the top
107 troposphere.

108 **2. Materials and Methods**

109 **2.1 Sample Collection**

110 Offline $\text{PM}_{2.5}$ samples with a 4-hour interval were synchronously collected onto prebaked
111 quartz filters (at 450 °C for 6 h) at two locations of Mt. Hua from 27 August to 17 September
112 2016. One sampling site locates at the mountain foot (MF, 34°32'N, 110°5'E; 400 m a.s.l.) and
113 another one is situated on the mountainside (MS; 34°29'N, 110°3'E; 1120 m a.s.l.) with little
114 anthropogenic activities due to its steep terrain in the mountain region. The horizontal distance
115 between the two sites is ~8 km and the vertical distance is about 1km (Figure S1). As revealed in
116 our previous study (Wu et al., 2022), vertical divergence simulated by WRF-Chem model

117 decreased gradually as enhanced elevation, along with the prevailing southerly winds, indicating
118 a feasibility of vertical transport of air parcel from MF to MS. And we also note that the change
119 of emission sources among two site was insignificant in a lifting air mass as indicated by
120 indistinctive divergences of diagnostic ratios and proportion of organic tracers from emission
121 sources. Such conditions can avoid the interferences caused by the emission sources change when
122 exploring aging process of BrC. More descriptions on the two sites have been documented by
123 our previous study along with the details on the sampling instrument setup (Wu et al., 2022).
124 Mass concentrations of PM_{2.5}, NO₂ and O₃ at MS site were directly quantified by E-BAM (Met
125 One Instruments, USA) and NO_x and O₃ Analyzer (Thermo, Model 42i, USA; Thermo, Model
126 49i, USA), respectively. At MF site, the data of above species apart from PM_{2.5} monitored by
127 another E-BAM were downloaded from the Weinan Ecological Environment Bureau
128 (<http://sthjj.weinan.gov.cn/>, last access: 8 July 2021). Meteorological parameters of both
129 sampling sites were downloaded from the Shaanxi Meteorological Bureau website
130 (<http://sn.cma.gov.cn/>, last access: 8 July 2021).

131 **2.2 Chemical Analysis**

132 The organic carbon (OC) and element carbon (EC) of PM_{2.5} filter samples were quantified
133 by a DRI model 2001 thermal–optical carbon analyzer following the IMPROVE-A temperature
134 protocol (Chow et al., 2007). Water-soluble organic carbon (WSOC) and water-soluble total
135 nitrogen (WSTN) of PM_{2.5} were extracted using Milli-Q pure water (18.2 MΩ) and determined
136 using a total organic carbon (TOC) analyzer (Model TOC-L CPH, Shimadzu, Japan). Water-
137 soluble organic nitrogen (WSON) is calculated by deducting the water-soluble inorganic
138 nitrogen (WSIN) from WSTN (i.e., WSON=WSTN-WSIN). Molecular compositions (e.g.,

139 nitrophenols, PAHs and other organic tracers) in the PM_{2.5} filter samples were quantified by a
140 gas chromatography (HP 7890A, Agilent Co., USA) coupled with mass spectroscopy detector
141 (GC/MS) (HP 5975, Agilent Co., USA) after the sample extraction and derivatization. Th
142 details of the extraction and derivatization can be found elsewhere (Li et al., 2023; Li et al.,
143 2020; Wang et al., 2006). Briefly, one-fourth of the filter sample was extracted with a mixture
144 of methanol and dichloromethane (2:1, v/v). Then the extracts were derivatized with N,O-bis-
145 (trimethylsilyl) trifluoroacetamide (BSTFA).

146 The stable nitrogen isotope compositions of NH₄⁺ ($\delta^{15}\text{N-NH}_4^+$) were determined by the
147 isotopic analysis of nitrous oxide (N₂O) derived from chemical conversion of NH₄⁺, and finally
148 quantified by a Precon-GasBench-IRMS system. This is a reliable method for nitrogen isotope
149 analysis of the sample with low NH₄⁺ concentration, of which precision can be up to 0.2‰.
150 More details upon the analytical artifact and quality control protocols can be found in our
151 previous studies. Furthermore, only the daytime samples were analyzed for the $\delta^{15}\text{N-NH}_4^+$ here,
152 and a merging pretreatment was applied for the daily samples to meet for the analysis
153 requirements.

154 Additionally, a high-resolution time-of-flight aerosol mass spectrometer (Aerodyne
155 Research Inc., Billerica, MA, USA) was employed to determine the chemical compositions of
156 water-soluble organic matter (WSOM) in PM_{2.5}, of which the method is similar to the report by
157 Daellenbach et al. (2016). The offline analytical procedure has been reported previously, here
158 we only give a brief description (Ge et al., 2017; Sun et al., 2011). One-eighth of the PM_{2.5}
159 filter samples was extracted with pure water. Then, the water-extracts were atomized using
160 argon as carrier gas, dried by a diffusion drier, and ultimately quantified by the aerosol mass

161 spectrometer. Purified water was also treated in the same manner prior to each sample running,
162 which was deemed as an analytical blank. As we mainly focused on the WSOM chemical
163 composition, a deep post-processing was conducted for the V-mode data in this study using the
164 Igor-based Aerosol Mass Spectrometer Analysis Toolkit. Element ratios of WSOM including
165 oxygen-to-carbon (O/C), hydrogen-to-carbon (H/C), nitrogen-to-carbon (N/C) , and organic
166 mass-to-organic carbon (OM/OC) ratios were determined according to the Improved Aiken (I-
167 A) method (Canagaratna et al., 2015). The mass load of WSOM in ambient air can be
168 accurately estimated using Eq.1, since the chemical species concentration in atomized aerosols
169 depends on the flow rate of the carrier gas and extract concentration.

$$\text{WSOM}=\text{WSOC}\times\text{OM}/\text{OC}_{\text{WSOM}} \quad \text{Eq. 1}$$

170 Where WSOM is water-soluble organic matter (WSOM) in the atmosphere ($\mu\text{g m}^{-3}$),
171 WSOC is water-soluble organic carbon (WSOC, μgCm^{-3}) in the atmosphere and measured by
172 the TOC analyzer, and $\text{OM}/\text{OC}_{\text{WSOM}}$ is the mass ratio of WSOM and OC determined by the
173 aerosol mass spectrometer. To obtain reliable data, the ionization efficiencies of HR-AMS was
174 calibrated with 300 nm (D_m) ammonium nitrate and ammonium sulfate particle following the
175 standard protocols (Sun et al., 2020; Jayne et al., 2000); and the relative ionization efficiencies
176 (RIEs) of 4.1 and 0.8 were used for ammonium and sulfate. While, the default RIEs were
177 applied for organics, nitrate and chloride.

178 **2.3 Optical Absorption of BrC**

179 Measurement of UV–vis absorption spectra of water-soluble BrC in $\text{PM}_{2.5}$ was performed
180 using a liquid waveguide capillary UV–vis spectrometer with a long effective path length (1
181 m). The extracted solution of BrC were prepared by a similar treatment to WSOC (Text S1), of

182 which absorption spectra were converted into the absorption coefficient at a given wavelength λ
183 (abs_λ , M/m, eq S1). The mass absorption efficiency (MAE_λ , m^2/gC) corresponding to water-
184 soluble BrC at a given wavelength λ can be calculated as follows:

$$\text{MAE}_\lambda = \frac{\text{abs}_\lambda}{M} \quad \text{Eq. 2}$$

185 Where M ($\mu\text{gC}/\text{m}^3$) is the mass concentration for water-soluble organic carbon (WSOC).
186 Absorption Ångström exponent (AAE) indicates the spectral dependence of a species, which
187 was quantified by a linear regression of $\log(\text{abs}_\lambda)$ versus $\log(\lambda)$ over a wavelength range of 300-
188 500 nm (Wu et al., 2020).

189 **2.4 Positive matrix factorization (PMF) source apportionment**

190 To quantitatively determine the fractional contribution of specific sources to BrC, a PMF
191 receptor model (EPA PMF 5.0 version) coupled with a bootstrap technique was applied here, of
192 which principle has been documented in previous studies (Brinkman et al., 2006; Paatero and
193 Tapper, 1994). Briefly, WSOC, WSON, SO_4^{2-} , NH_4^+ , Mg^{2+} , Ca^{2+} , $\text{abs}_{365 \text{ nm}}$ and organic tracers
194 (BbF, Bghip and levo.) are the input variables in the present work, all of which are regarded as
195 strong variable expect for Bghip with a low S/N ratio (0.6). Another input dataset is uncertainty
196 matrix that is calculated according to the following equations (Eq.3). The uncertainties of each
197 factor profile are also evaluated by a bootstrap analysis, of which result showed that
198 reproducibility of each source factor was $>80\%$ (Table S1), indicating a well robustness. In our
199 previous study on Mt. Hua (Wu et al., 2022), insignificant change in the corresponding emission
200 sources was revealed during air mass lifting process. Thus, the daytime samples from both sites
201 were added together as one data matrix. Considering Q values and interpretability, four factors
202 were obtained as the optimal solution after numerous testes with three to seven factors, and the

203 input species matched well with simulated ones with significant correlations ($R^2 > 0.88$).

$$\text{uncertainty} = \begin{cases} \frac{5}{6} \times \text{MDL} & (\text{concentration} < \text{MDL}) \\ \sqrt{(\text{error fraction} \times \text{concentration})^2 + (0.5 \times \text{MDL})^2} & (\text{concentration} > \text{MDL}) \end{cases} \text{ Eq.3}$$

204 Where MDL is the method detection limit. And the error fraction is set to 5% for $\text{PM}_{2.5}$ (Gao
205 et al., 2018); WSOC , WSON , SO_4^{2-} , NH_4^+ , Mg^{2+} , Ca^{2+} are estimated to be 7 %, those of other
206 species are 12%. To reduce the error, the sample with missing data for individual species would
207 be excluded rather than replace by the mean value of whole campaign.

208 **2.5 Random forest analysis for WSON**

209 Random forest (RF), as a powerful tool, has been used widely in the regression and prediction
210 problems upon atmospheric pollutions, even the data have complex nonlinear relationships and
211 interactions (Hu et al., 2017; Vu et al., 2019). To reveal the key factors that may affect the WSON
212 formation during the air mass lifting process, a RF regression model was applied for the daytime
213 samples at MF and MS sites, respectively. And the potential factors, including pH, ALWC, T, RH,
214 NH_4^+ , $\text{NH}_3(\text{aq})$, NO_2 , nitrophenols, O_3 and organic matter (OM), herein were regarded as the
215 predictors for WSON. In the RF model design, about 70% of these original data were randomly
216 divided into the training dataset to construct the RF model, and the rest was deemed as the testing
217 data for testing the model performance. There are two important parameters being constantly
218 optimized in the model construction process, including the number of trees grown (n_{tree}) and
219 number of variables split at each node (n_{mtry}); After numerous tests, n_{tree} and n_{mtry} were set as 100
220 and 10 for MF data, and 128 and 9 for MS data, respectively, to achieve the best prediction
221 accuracy. Furthermore, a 10-fold cross-validation technique was employed here to
222 simultaneously tune model parameters and estimate model performance. And the statistical
223 metrics including coefficient of determination (R^2), mean square error (MSE) or root-mean-

224 square error (RMSE) and mean absolute error (MAE), were established to evaluate prediction
225 accuracy of the model. As shown in Table S2, the predicted data for the testing dataset has strong
226 correlativity with observed ones, with small values for those error metrics; These results indicated
227 a satisfactory performance of the RF model for explain the importance of these factors to daytime
228 WSON formation.

229 **3. Results and Discussion**

230 **3.1 Enhanced Light Absorption of BrC in the Mountainous Atmosphere**

231 Figure 1 shows the temporal variations in light absorption (abs_{365nm}) and concentrations of
232 fine particulate WSOC simultaneously observed at the mountain foot (MF) and mountainside
233 (MS) sites. The variation patterns of water-soluble BrC (i.e., abs_{365nm}) at both sites were closely
234 followed by WSOC ($R^2 > 0.70$, Figure 1 and Figure S2); this indicated that BrC is an important
235 part of WSOC, of which light absorption of BrC markedly increased with a decrease in light
236 wavelengths. As summarized in Table 1, the averaged abs_{365nm} of BrC was $2.1 \pm 1.4 M m^{-1}$ at
237 MS, approximately corresponding to 40% of that ($5.1 \pm 2.4 M m^{-1}$) at MF. The light-absorbance
238 level of BrC at the high altitude MS site is in the same range as those reported from Chinese
239 megacities such as Beijing (Cheng et al., 2016) and Xi'an (Wu et al., 2020), indicating a strong
240 light-absorption of BrC in the upper boundary layer over Guanzhong Basin, inland China.
241 Absorption Ångström exponent (AAE) at MS is 5.7 ± 1.3 (Table 1), slightly lower than that at
242 the ground MF site (6.0 ± 0.5). Such a difference in AAE ($p < 0.05$) is most likely related to the
243 difference in chemical composition of the chromophores between the two sites with different
244 altitudes. The averaged mass absorption efficiency (MAE) at MS (MAE_{365nm} , $0.67 \pm 0.2 m^2 g^{-1}$)
245 was almost equal to that at MF ($0.69 \pm 0.2 m^2 g^{-1}$) but 30-40% higher than those observed in

246 Chinese megacities such as Beijing (Du et al., 2014) and Nanjing (Chen et al., 2018) ($\sim 0.5 \text{ m}^2$
247 g^{-1} , in summertime), further demonstrating a strong light-absorption nature of BrC in the upper
248 boundary layer of Guanzhong Basin, inland China.

249 Figure 2 shows the diurnal variations of $\text{abs}_{365 \text{ nm}}$ and $\text{MAE}_{365 \text{ nm}}$ at both sites during the
250 campaign. At MF site a morning peak of $\text{abs}_{365 \text{ nm}}$ driven by enhanced traffic emissions
251 occurred at 8:00~12:00 (local time, thereafter), and then gradually decreased and reached a
252 minimum at 12:00~16:00 with the lowest MAE at 365 nm wavelength ($\text{MAE}_{365 \text{ nm}}, 0.57 \pm 0.14$
253 $\text{m}^2 \text{ g}^{-1}$ (Figures 2a and 2b). Such a ground surface decrease in light absorption of BrC at early
254 afternoon can be attributed to the daytime boundary layer growth and photobleaching. This can
255 be verified by the oxidation state of carbon (OSc) measured by the aerosol mass spectrometer,
256 of which higher value is indicative of a deeper degree of atmospheric oxidation (Li et al.,
257 2019a). As seen in Figure 3a, $\text{abs}_{365 \text{ nm}}$ negatively correlated with OSc, which is consistent with
258 those reported by previous laboratory experiments (Lee et al., 2014; Zhao et al., 2015; Sunlin
259 et al., 2017) and suggests that atmospheric aging can significantly diminish the light-absorption
260 of BrC. On the contrary, $\text{abs}_{365 \text{ nm}}$ at MS site remarkably enhanced with the boundary layer
261 growth and peaked at 12:00~16:00 (Figure 2a), despite the fact that the aerosol was further
262 oxidized during the upward transport as indicated by the a higher OSc value at MS site (Figure
263 3b); The OSc variation among both sites coincided with that of BaP/BeP being a known proxy
264 of whether aerosols are freshly emitted (> 1) or aged (< 1) (Figure S3). Moreover, a moderate-
265 increased $\text{MAE}_{365 \text{ nm}}$ was also observed in this process (Figure 2b). As shown in Table 1, the
266 light absorption of BrC at 365 nm relative to BC at 550 nm ($\text{abs}_{365 \text{ nm}}\text{-BrC}/\text{abs}_{550 \text{ nm}}\text{-BC}$) during the
267 daytime at MS was 0.28 ± 0.08 , which is approximately 30% higher than that (0.22 ± 0.08 , Table

268 1) at MF. Our previous study at Mt. Hua found that changes in sources of primary organic
269 aerosols in the air mass transported from MF to MS were insignificant (Wu et al., 2022),
270 indicating that there was no additional emission of BrC during the air mass upward transport.
271 Thus, the enhanced light-absorption of BrC relative to BC at MS is solely ascribed to a
272 secondary formation of absorbing BrC (Figure 2c); and these secondary BrC were highly light-
273 absorbing despite more aged atmosphere aloft, as verified by a strongly positive correlation
274 between MAE_{365nm} and OSc values at MS site (Figure S4).

275 To further elucidate the above hypothesis, a PMF analysis was applied for the source
276 apportionment of the daytime abs_{365 nm} at both sites. As seen in Figure S5, four types of BrC
277 sources were identified. In brief, fossil fuel combustion and biomass burning influenced by
278 local-related emissions were primary sources for the surface BrC, consistent with observations
279 in other cities (Li et al., 2023; Wu et al., 2020; Wang et al., 2022a). However, BrC at MS site
280 was produced dominantly from secondary formation, of which the contribution to the total BrC
281 is 53% and about twice of that at MF (Figure 2d) further corroborating a substantial formation
282 of BrC with relatively stronger light-absorptivity during the air mass lifting process. These
283 secondarily formed BrC chromophores engender a more light-absorption of high-altitude BrC
284 compared with that of BC (or EC, Figure 2c); Similar vertical profile of BrC in the upper
285 troposphere (5~12 km) of the continental US was also observed by in-situ aircraft
286 measurements (Zhang et al., 2017).

287 **3.2 Secondary Formation of BrC in the Air Mass Lifting Process**

288 Figure 4a illustrates the diurnal cycles of N:C ratio of the water-soluble organic matter in
289 PM_{2.5} measured by the high-resolution time-of-flight aerosol mass spectrometer. At the MF site

290 N:C ratio did not vary much with time and was even leveling off in the daytime, which
291 indicates that the compositions of light-absorbing chromophores are similar throughout the day.
292 Nonetheless, the diurnal pattern of N:C ratio at MS was analogous to that of $\text{abs}_{365\text{ nm}}$ and
293 $\text{MAE}_{365\text{ nm}}$ with a daily peak at 12:00~16:00 and a moderate positive correlation was also
294 observed between N:C ratio and $\text{abs}_{365\text{ nm}}$ ($R^2=0.38$, $P<0.01$), suggesting that nitrogen-
295 containing organic compounds (NOCs) have significant contributions to the BrC light-
296 absorption in the upper boundary layer. Such a results is consistent with the laboratory
297 simulation, in which NOCs have been reported to contribute up to 60% of the absorbance of
298 secondary BrC over a wavelength rang of 300–400 nm (Lin et al., 2015). Moreover, the
299 daytime N:C ratios were 20% higher at MS (0.066 ± 0.014) than those at MF (Figure 4a),
300 indicating that additional NOCs were produced in the air mas lifting process. In fact, numerous
301 N-containing organic fragments including $\text{C}_x\text{H}_y\text{N}$ and $\text{C}_x\text{H}_y\text{O}_z\text{N}$ at the MS site were detected by
302 the aerosol mass spectrometer, accounting for ~13% of the total water-soluble OM; And
303 fractional contribution of above fragments at MS enhanced by approximately 10% compared to
304 that at MF site, even up to ~25% at the day with low $\text{PM}_{2.5}$ load ($<75\ \mu\text{g}/\text{m}^3$) (Figure S6). This
305 suggested an enhanced formation of WSON during the air mass transport from the lower
306 mountain foot site to the upper mountainside site. Since WSON at MS moderately was
307 positively correlated with light absorption of BrC at $\lambda=365\text{ nm}$ (Figure S7, $R^2=0.45$, $p<0.05$),
308 the enhancement in BrC light-absorption at MS can largely be attributed to secondary
309 formation of NOCs during the air mass transport from the ground surface to the upper
310 boundary layer.

311 Light-absorbing NOCs including reduced nitrogen species (e.g., imidazoles and pyrazines)

312 and oxidized ones (e.g., nitroaromatics) can be generated via various types of gas- and particle-
313 phase reactions, such as, NH₃-mediated carbonyl-to-imine reactions, nitration of aromatic
314 compounds, and heterogeneous reactions of ·OH and NO₂· radicals with phenolic compounds
315 (Moise et al., 2015; Laskin et al., 2015). The potential pathways and dominating factors for
316 NOCs formation at MS site will be explored in the following sections.

317 **3.3 Gas-Phase Formation of BrC in the Air Mass Lifting Process**

318 Nitroaromatic compounds (NACs) are strong light-absorbing compounds and are
319 ubiquitous in the atmosphere. In this study, a total of six NACs in the PM_{2.5} samples were
320 detected (Table S3), which exhibited a significant correlation with abs_{365 nm} at both sampling
321 sites (Figure S2c and S2d), indicating an important impact of NACs on the aerosol light-
322 absorption. As seen in Figure 5a, both NACs concentration and NACs/OC ratio decreased
323 gradually at MF, reaching the daily minimum at 12:00-16:00. Such an abatement in NACs was
324 mainly attributed to the boundary layer expansion and an enhanced photooxidation (Figure 3).
325 Furthermore, the daytime NACs at MF well correlated with (BbF+levoglucosan) being known
326 as tracers for combustion emissions ($R^2 > 0.76$, Figure 5c), but not correlated with gaseous NO₂
327 (Figure 5e), suggesting that most of NACs at the ground surface site were directly emitted from
328 combustion sources. This can be further verified by a strong positive correlation of NACs and
329 CO prevailing in combustion exhausts (Figure S8). As partial NACs at the MF site can be
330 transported aloft by anabatic valley winds, thus a moderate correlation ($R^2 = 0.49$, Figure 5d)
331 between NACs and (BbF+levoglucosan) was observed at MS site. However, a moderate
332 correlation between NO₂ and NACs ($R^2 = 0.57$, $p < 0.01$, Figure 5f) observed at MS suggests a
333 non-negligible formation of secondary NACs under the transport process. As revealed by

334 previous studies, the $\text{NO}_3\cdot/\cdot\text{OH}$ -oxidation of phenolic volatile organic in the presence of NO_x
335 can form numerous NACs that followed by partitioning to the condensed phase (Wang and Li,
336 2021; Li et al., 2021a; Finewax et al., 2018); and the photolysis of nitrite in aerosol aqueous
337 phase can also lead to the nitration of phenol/catechol to generated NACs (Vione et al., 2005).
338 But a poor relationship between NACs and aerosol liquid water content (ALWC; $R^2=0.34$,
339 $P>0.05$) and particulate phenol ($R^2<0.1$), indicating a minor contribution of aqueous-phase
340 formation to NACs aloft.

341 The preceding discussion provided reliable evidences that partial NACs could be formed
342 by gas-phase reactions, but they only accounted for a very small fraction of OC (Figure 5a-b),
343 suggesting that gas-phase formation is probably not the major formation pathway of secondary
344 NOCs during the air mass vertical transport. Further evidence for this hypothesis was provided
345 by a random forest (RF) analysis being used as a metric for the degree of correlation between
346 these influencing factors (ALWC, pH, T, NO_2 , NH_4^+ , etc.) and WSON at both sampling sites
347 (Figure 4b and Figure S9). As revealed by RF model results, nitrophenols and gaseous NO_2
348 were important influencing factors for the daytime WSON at MF site (Figure S9), of which
349 importance was explained up to 35% but only~15% of that aloft (Figure 4b), confirming a less
350 importance of the gas-phase reactions for the light-absorbing NOCs formation in the vertical
351 transport process.

352 **3.4 Aerosol Aqueous Formation of BrC in the Air Mass Lifting Process**

353 As shown in Figure 4b, RF analysis showed that the variation in concentration of WSON
354 in $\text{PM}_{2.5}$ at MS was largely affected by NH_4^+ (23.0%) and ALWC (17.3%). Given a relatively
355 strong correlation between WSON and NH_4^+ ($R^2=0.70$, Figure 4c), we proposed that aqueous

356 phase reactions induced by ammonium is the major formation pathway for water-soluble NOCs
357 at MS. For further demonstrating such a hypothesis, we analyzed nitrogen isotope composition
358 ($\delta^{15}\text{N-NH}_4^+$) of ammonium in the $\text{PM}_{2.5}$ samples at both sites, of which the analytical details
359 has been described in our previous study (Wu et al., 2022). As seen in Figure 4d, WSON
360 showed a strong negative correlation with $\delta^{15}\text{N-NH}_4^+$, probably due to the irreversible reactions
361 involving ammonia favored ^{15}N depletion in the particle form as revealed by Heaton et al.
362 (1997). In contrast, WSON at MF presented a similar correlation with NH_4^+ as that at MS but
363 did not correlate with $\delta^{15}\text{N-NH}_4^+$ (Figures S10 (a) and (b)).

364 Previous studies have demonstrated the importance of $\text{NH}_4^+/\text{NH}_3$ in the formation of light-
365 absorbing imidazoles and N-heterocycles from the carbonyls (e.g., glyoxal (Gly) and
366 methylglyoxal (mGly) generated from oxidation of VOCs) (Li et al., 2021b; Moise et al., 2015;
367 Kampf et al., 2012; Liu et al., 2023). Figure S11 depicts a simple reaction pathway for above
368 aqueous reactions, in which the chromophore products contain less amounts of O and H atoms.
369 Such a phenomenon was found for the daytime NOCs at MS. As shown in Figure 6, the H/N
370 and O/N ratios of BrC in $\text{PM}_{2.5}$ at MS exhibited a strongly negative correlation with N/C ratio,
371 respectively ($R^2=0.92$ for H/N and $R^2=0.84$ for O/N). Considering the fact that Gly and mGly
372 are abundant at the daytime atmosphere of Mt. Hua (Qi et al., 2023), the aqueous reactions of
373 dicarbonyls with $\text{NH}_4^+/\text{NH}_3$ are probably the major pathway to yield NOCs during the vertical
374 transport.

375 Above aqueous reactions could also occur at MF site as depicted in Figure S11, but it was
376 insignificant compared to that at MS site, attributing the disparity in chemical compositions.
377 Our previous study found that the ground surface MF aerosols were more acidic ($\text{pH}=2.9$) and

378 dominated by NH_4HSO_4 , while the upper boundary layer MS aerosols were less acidic
379 ($\text{pH}=3.4$) and dominated by abundant $(\text{NH}_4)_2\text{SO}_4$. Such differences in aerosol acidity and
380 chemical compositions between two sites can favor the formation of NOCs at the MS site, as
381 evident from a recent experimental observation by Li et al. (2021b), who found that NOCs
382 yield on the $(\text{NH}_4)_2\text{SO}_4$ seeds exposing to Gly or mGly vapor was relatively higher than that on
383 NH_4HSO_4 seeds. Also, they found that mGly has a larger uptake coefficient on $(\text{NH}_4)_2\text{SO}_4$
384 particles with a relatively higher NOCs yield compared to Gly, because mGly has a stronger
385 interfacial attraction and thus has a more efficient nucleophilic addition involving the
386 carbenium ions (Li et al., 2021b). Our previous study showed that the summertime atmosphere
387 of Mt. Hua is dominated by biogenic VOCs and the concentration of fine particulate mGly is
388 about five times that of Gly (Meng et al., 2014). Such a predominance of mGly over Gly and a
389 less acidic aerosol aqueous aerosol phase at the MF site are favorable for light-absorbing NOCs
390 formation on $(\text{NH}_4)_2\text{SO}_4$ particles, which can mainly be responsible for the enhanced light
391 absorption of BrC at the mountainous site with the ratio of light absorption of BrC to BC higher
392 in the upper boundary layer than that in the ground surface.

393 **3.5 Atmospheric implications**

394 Our work provides an evolution profile of BrC during air mass vertical transport, and
395 highlights a secondary formation of BrC in this process, which can be responsible for the
396 enhancement of BrC relative to BC in the upper boundary layer. As the longer lifetime of high-
397 altitude BrC, they can disperse rapidly into a large area (Zhang et al., 2017), exerting
398 significant influence on regional climate that is even comparable to BC in the upper tropical
399 troposphere (Jo et al., 2016). Moreover, we also revealed a vital role of aqueous-phase

400 reactions for the secondary formation of BrC in the air mass lifting process, specifically the
401 $\text{NH}_4^+/\text{NH}_3$ -induced reactions (e.g., Maillard reaction) that can form NOCs with stronger light-
402 absorptivity. As ammonia and carbonyls such as glyoxal and methylglyoxal are ubiquitous in
403 the troposphere, thus our work suggests that the above formation mechanism on the light-
404 absorbing NOC aerosols could extensively occur in the troposphere.

405 In the past decade, the haze pollution in China has changed from previous sulfate-
406 dominated environment (SD) to the current nitrate-dominated environment (ND) due to the
407 effective sulfur emission control, which would significantly enhance the aerosol ALWC since
408 nitrate is more hygroscopic than sulfate at a given RH and aerosol loading. While, as indicated
409 by our previous observational evidences (Lv et al., 2023), high ALWC load induced by
410 abundant nitrate would efficiently promote more WSOC partitioning into the aerosol phase
411 compared with that in SD ones, and thus may increase the BrC yield because WSOC contains
412 numerous BrC precursors. With the increase in relative abundance of nitrate to sulfate, nitrate-
413 enhanced gas-to-particle partitioning of WSOC will become highly efficient in China in the
414 near future, meaning that the BrC formation will be more active hereafter. Additionally, the
415 national VOCs and NH_3 emissions have remained at high levels, and even have shown a slight
416 increasing trend. The abundant NH_3 can not only participates in the formation of BrC but also
417 affects BrC optical properties by regulating the aerosol acidity. To further reveal the impact on
418 the BrC, the NH_3 concentration and $\text{MAE}_{365\text{ nm}}$ value of water-soluble BrC in different region
419 of China were statistically explored on a national scale (Figure 7a). As depicted in Figure 7a,
420 the spatial pattern of $\text{MAE}_{365\text{ nm}}$ is closely coincident with NH_3 levels with a robust positive
421 correlation ($R^2=0.87$). Such a spatial distribution pattern indicates that NH_3 -rich conditions are

422 favorable for formation of BrC with strong light-absorptivity. China is one of the countries with
423 strongest NH₃ emissions in the world due to huge demand for N-fertilizer (Van Damme et al.,
424 2018), and thus atmospheric NH₃ in China is much higher than that in Europe and the United
425 States. This is probably one of the factors causing the higher concentrations of BrC with
426 stronger light-absorptivity in China compared with developed countries (Figure 7b).

427 **4. Conclusion**

428 Synchronous observations upon optical properties and chemical compositions of
429 atmospheric BrC were conducted at MF and MS of Mt. Hua, and revealed that light-absorption
430 of BrC aloft was only ~40% of that at surface owing to a dilution effect caused by planetary
431 boundary layer upliftment. And the light-absorption of BrC relative to black carbon moderately
432 enhanced in the lifting process of air mass from MF to MS coincide with the variation of the
433 daytime MAE_{365nm} aloft, indicating a secondary formation of BrC; and these secondary BrC
434 accounted for >50% of total at MS site. While, the surface BrC was mainly originated from
435 direct combustion emission, with a 55% of fractional contribution to the total.

436 The N:C ratio of WSOM was measured by offline AMS, of which diurnal pattern was
437 analogous to that of abs_{365 nm} and MAE_{365 nm} aloft, substantiating a considerable contribution of
438 nitrogen-containing organic compounds (NOCs) to BrC light-absorption. And daytime N:C
439 ratio site was approximately 15% higher at MS than that at MF, mainly due to a significant
440 formation of secondary NOCs produced by NH₄⁺/NH₃-induced reactions (e.g., Maillard
441 reaction). Moreover, a robust positive relationship between MAE_{365 nm} and NH₃ load was
442 statistically explored in nationwide, strongly manifested that abundant NH₃ maybe one of key
443 factors for the high BrC load with strong light-absorptivity in China compared to that in

444 developed countries. Therefore, NH₃ emission control in China is indispensable for further
445 alleviating haze and BrC pollutions in the country.

446 **Data availability.** The data used in this study are freely available at
447 <https://doi.org/10.5281/zenodo.10926469> (Wu, 2024). And Meteorological data and hourly
448 PM_{2.5}, NO₂, O₃ concentrations can be obtained from <https://doi.org/10.5281/zenodo.7413640>
449 (Wu, 2022a).

450 **Author contributions.** G.W. designed research and contributed analytic tools. C.W., C.C. and
451 J.L. collected the samples. C.W., X.L., K.Z. and G.W. conducted the sample analysis. C.W.,
452 S.Z. and G.W. performed the data interpretation. C.W. and G.W. wrote the paper. All authors
453 contributed to the paper with useful scientific discussions.

454 **Competing interests.** The authors declare no competing interest.

455 **Acknowledgements.** This work was financially supported by the National Natural Science
456 Foundation of China (No. 42130704, 42007202) and ECNU Happiness Flower program.

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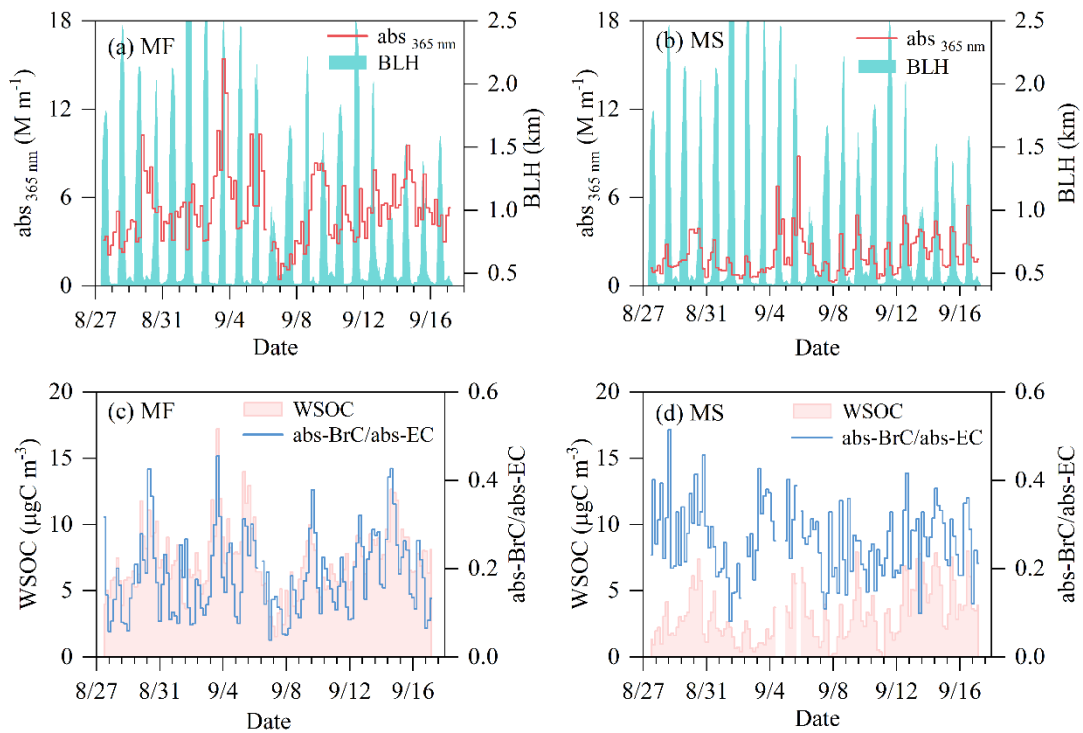
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Table 1 Optical properties of BrC and mass concentrations organic carbon/nitrogen in PM_{2.5} and meteorological parameters at the two sampling sites.

	Mountain foot (MF)			Mountainside (MS)		
	Average	Daytime	Nighttime	Average	Daytime	Nighttime
(i) Optical properties of BrC and acidity of PM_{2.5}						
abs ₃₆₅ (M m ⁻¹)	5.1±2.4	5.0±2.5	5.2±2.2	2.1±1.4	2.6±1.3	1.6±1.3
MAE ₃₆₅ (m ² g ⁻¹)	0.69±0.2	0.66±0.18	0.73±0.18	0.67±0.21	0.67±0.15	0.68±0.26
AAE	6.0±0.5	6.1±0.51	6.0±0.51	5.7±1.3	5.5±0.9	5.8±1.7
abs ₃₆₅ -BrC/abs ₅₅₀ -BC ^a	0.18±0.09	0.22±0.08	0.17±0.09	0.26±0.08	0.28±0.08	0.25±0.07
pH	2.9±2.0	2.3±1.6	3.6±2.1	3.4±2.2	3.5±2.2	3.3±2.2
(ii) Concentrations of carbonaceous PM_{2.5} and aerosol liquid water content (ALWC)						
WSOC (µgC m ⁻³)	7.3±2.5	7.6±2.8	7.0±2.1	3.2±2.1	4.0±2.1	2.4±1.7
WSON (µgN m ⁻³)	2.3±1.6	2.5±1.7	2.0±1.4	1.2±0.9	1.5±1.1	0.8±0.7
OC (µgC m ⁻³)	14.0±4.7	12.4±4.6	15.4±4.4	5.0±2.8	6.3±2.8	3.8±2.3
EC (µgC m ⁻³)	4.3±2.0	3.1±1.0	5.4±1.9	1.1±0.7	1.3±0.7	0.8±0.4
Nitrophenols (ng m ⁻³)	16±13	12±10	19±15	2.5±1.9	3.2±2.2	1.7±1.1
ALWC (µg m ⁻³)	28±64	11±15	44±86	27±71	18±24	35±95
WSOC/OC	0.54±0.15	0.62±0.13	0.47±0.11	0.62±0.21	0.62±0.16	0.61±0.25
(iii) Meteorological parameters						
T (°C)	23±4.2	27±3.0	20±2.4	15±2.5	16±2.3	14±2.3
RH (%)	69±18	56±14	81±14	63±20	62±19	63±21
Wind speed (m s ⁻¹)	1.3±1.1	1.5±0.93	1.2±1.2	3.2±2.0	2.7±1.5	3.8±2.3
Visibility (km)	14±9.5	16±9.6	12±9.0	22±12.1	21±12	24±12.0

700 ^aThe abs₅₅₀-BC was calculated according to the mass absorption efficiency (MAE) of BC (black
 701 carbon) reported by Bosch et al. (2014), and the light wavelengths for the abs₃₆₅-BrC and abs₅₅₀-
 702 BC are 365 nm and 550 nm, respectively.
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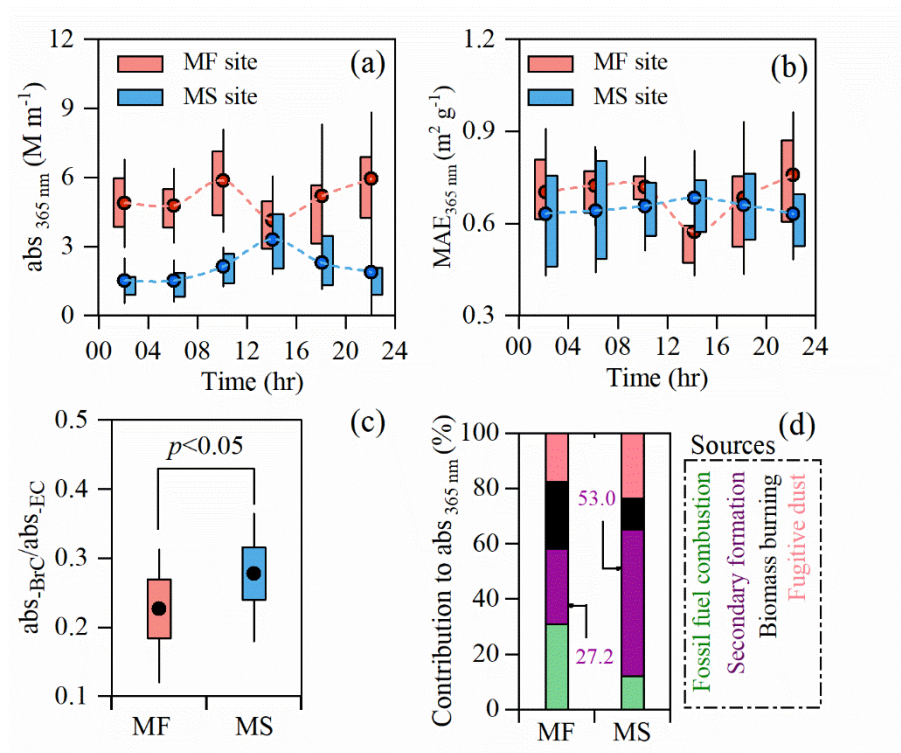
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Figure 1. Temporal variations of light absorption of BrC in the ground surface (mountain foot site, MF) and the upper boundary layer (mountainside site, MS) atmospheres in inland China. (a, b) Abs_{365} and boundary layer height (BLH). (c, d) Concentration of WSOC and the ratio of light absorption of BrC at $\lambda=365 \text{ nm}$ to BC at $\lambda= 550 \text{ nm}$.

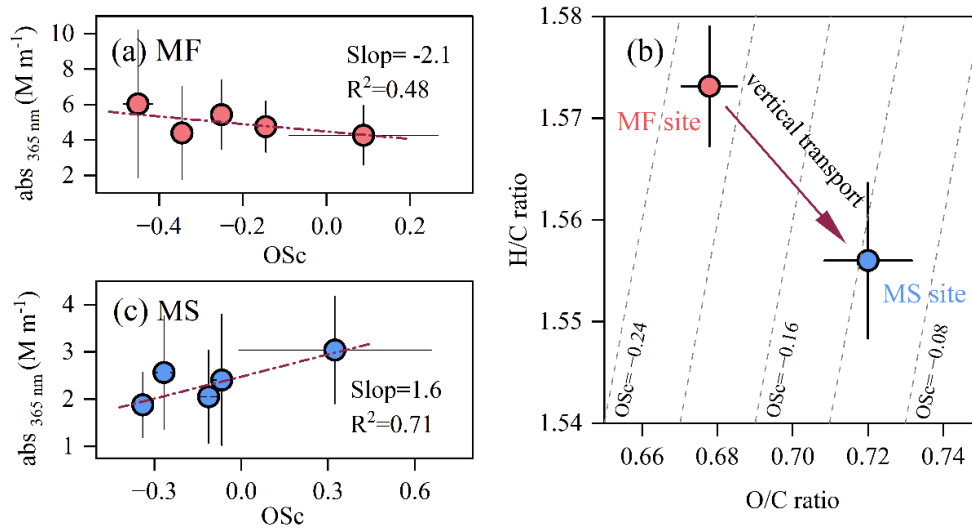
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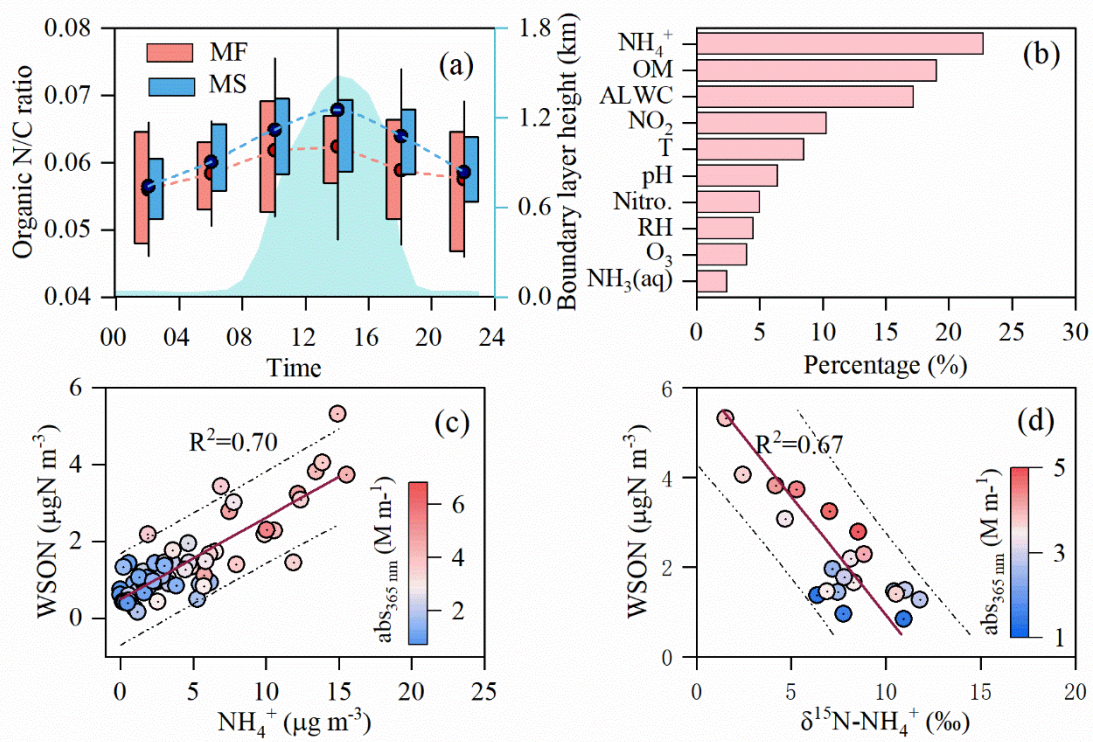
716 **Figure 2. BrC formation in air mass lifting process.** (a) and (b) Diurnal variations in $abs_{365\text{ nm}}$
717 and MAE at the mountain foot (MF) and mountainside (MS) sites. (c) Ratio of light absorption
718 of BrC at $\lambda=365\text{ nm}$ to that of BC (abs_{BrC}/abs_{BC}) at $\lambda=550\text{ nm}$ in daytime at both sites (The
719 abs_{BC} at $\lambda=550\text{ nm}$ was calculated according to mass absorption efficiency of EC reported by
720 Bosch et al. (2014)). (d) Source apportionment for the daytime BrC at the two sites. The whisker
721 boxes show mean (dot), 25th–75th percentile ranges (box), and standard deviation values
722 (whiskers).

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Figure 3. Evolution in chemical composition of daytime water-soluble BrC in the air mass transport from the mountain foot (MF site, red dots) to the mountainside (MS site, blue dots). (a) and (c) Light absorption (abs_{365}) of daytime water-soluble BrC as a function of their oxidation state ($\text{OSc}=2\text{O/C}-\text{H/C}$) at MF and MS sites, respectively. (b) The VK-triangle diagram of water-soluble BrC at the two sites.



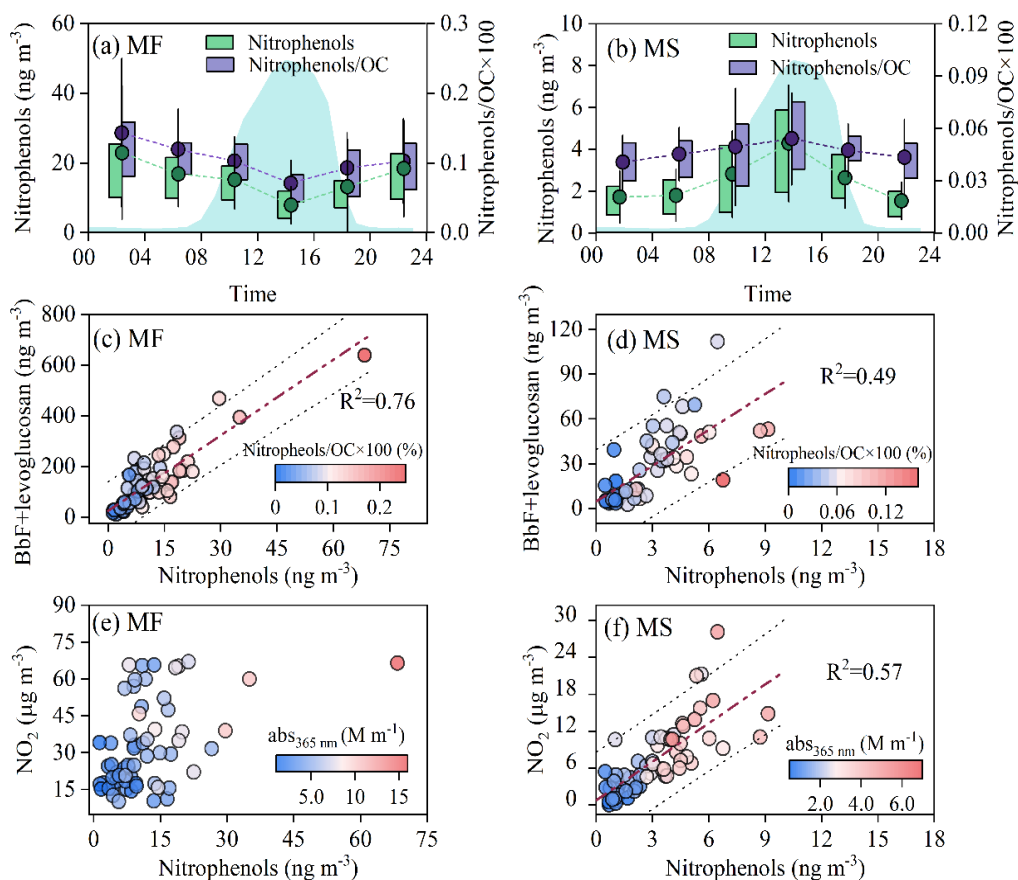
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744 **Figure 4. Formation of water-soluble organic nitrogen compounds (WSOs) in the air**
 745 **mass lifting process.** (a) Diurnal variations in elemental ratio of N/C of fine particulate water-
 746 soluble organics at the mountain foot (MF) and mountainside (MS) sites, the whisker boxes
 747 show mean (dot), 25th–75th percentile ranges (box), and standard deviation values (whiskers).
 748 (b) Importance assessment for the key factors affecting the daytime WSON at MS site. (c) and
 749 (d) Linear fit regressions for WSONs with NH₄⁺ and δ¹⁵N-NH₄⁺ in the daytime PM_{2.5} aerosols
 750 at MS site, respectively.

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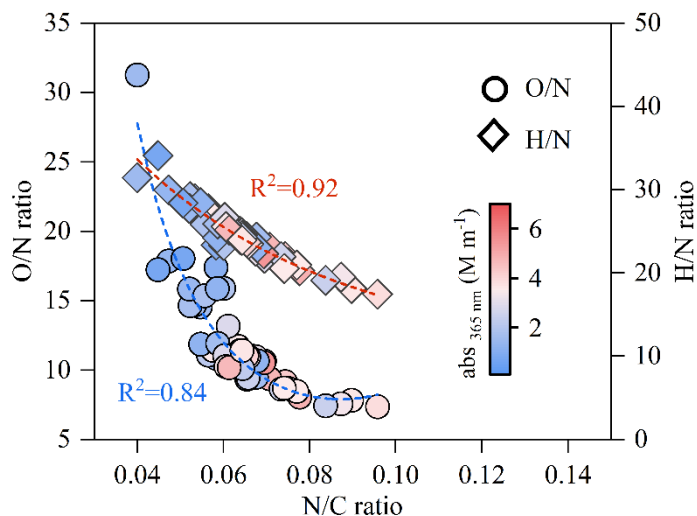
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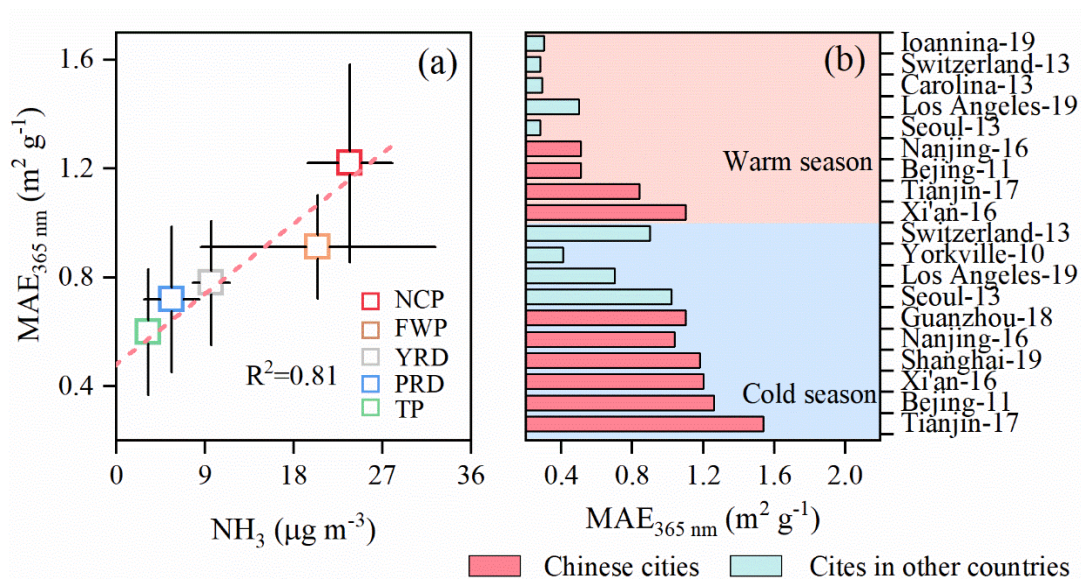
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756 **Figure 5.** The source and secondary formation for nitrophenols at MF and MS sites. (a and b)
757 Diurnal variations in nitrophenols and mass ratio of nitrophenols to OC (nitrophenols/OC); blue
758 sky shade indicates diurnal variation of boundary layer height, and the whisker boxes show
759 mean (dot), 25th–75th percentile ranges (box), and standard deviation values (whiskers). Linear
760 fit regression for nitrophenols with BbF+levoglucosan (c and d) and NO₂ (e and f),
761 respectively.
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Figure 6. Elemental composition of daytime WSOC at MS site.



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Figure 7. Impact of NH_3 on atmospheric BrC over China. (a) Linear correlation between NH_3 and MAE of BrC in different regions of China (NCP, north China Plain; FWP: Fenwei Plain; YRD, Yangtze River delta; PRD, Pearl River delta; TP: Tibetan Plain). (b) MAE₃₆₅ of BrC in China and other countries (The details of datasets from literatures, including specific sites, time periods, and sources, etc., are given in Table S4 and S5).