Potential underestimation of ambient brown carbon absorption based on the methanol extraction method and its impacts on source analysis

Zhenqi Xu¹, Wei Feng¹, Yicheng Wang¹, Haoran Ye¹, Yuhang Wang², Hong Liao¹, and Mingjie Xie¹

¹Collaborative Innovation Center of Atmospheric Environment and Equipment Technology, Jiangsu Key Laboratory of Atmospheric Environment Monitoring and Pollution Control, School of Environmental Science and Engineering, Nanjing University of Information Science & Technology, 219 Ningliu Road, Nanjing 210044, China
²School of Earth and Atmospheric Sciences, Georgia Institute of Technology, Atlanta, GA 30332, United States

Correspondence: Mingjie Xie (mingjie.xie@colorado.edu, mingjie.xie@nuist.edu.cn)

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Abstract. The methanol extraction method was widely applied to isolate organic carbon (OC) from ambient aerosols, followed by measurements of brown carbon (BrC) absorption. However, undissolved OC fractions will lead to underestimated BrC absorption. In this work, water, methanol (MeOH), MeOH/dichloromethane (MeOH/DCM, 1:1, v/v), MeOH/DCM (1:2, v/v), tetrahydrofuran (THF), and N,N-dimethylformamide (DMF) were tested for extraction efficiencies of ambient OC, and the light absorption of individual solvent extracts was determined. Among the five solvents and solvent mixtures, DMF dissolved the highest fractions of ambient OC (up to ∼95 %), followed by MeOH and MeOH/DCM mixtures (<90 %), and the DMF extracts had significantly (p<0.05) higher light absorption than other solvent extracts. This is because the OC fractions evaporating at higher temperatures (>280 °C) are less soluble in MeOH (∼80 %) than in DMF (∼90 %) and contain stronger light-absorbing chromophores. Moreover, the light absorption of DMF and MeOH extracts of collocated aerosol samples in Nanjing showed consistent temporal variations in winter when biomass burning dominated BrC absorption, while the average light absorption of DMF extracts was more than 2 times greater than the MeOH extracts in late spring and summer. The average light absorption coefficient at 365 nm of DMF extracts was 30.7 % higher (p<0.01) than that of MeOH extracts. Source apportionment results indicated that the MeOH solubility of BrC associated with biomass burning, lubricating oil combustion, and coal combustion is similar to their DMF solubility. The BrC linked with unburned fossil fuels and polymerization processes of aerosol organics was less soluble in MeOH than in DMF, which was likely the main reason for the large difference in time series between MeOH and DMF extract absorption. These results highlight the importance of testing different solvents to investigate the structures and light absorption of BrC, particularly for the low-volatility fraction potentially originating from non-combustion sources.
1 Introduction

Besides black carbon (BC) and mineral dust, growing evidence shows that organic carbon (OC) aerosols derived from various combustion sources (e.g., biofuel and fossil fuel) and secondary processes (e.g., gas-phase oxidation, aqueous and in-cloud processes) can absorb sunlight at short visible and UV wavelengths (Laskin et al., 2015; Hems et al., 2021). The radiative forcing (RF) of the light-absorbing organic carbon, also termed “brown carbon” (BrC), is not well quantified due to the lack of its emission data, complex secondary formations, and large uncertainties in in situ BrC measurements (Wang et al., 2014, 2018; Saleh, 2020). The imaginary part of the refractive index \( k \) of BrC is required when modeling its influence on aerosols direct RF and is retrieved by the optical closure method combining online monitoring of aerosol absorption and size distributions with Mie theory calculations (Lack et al., 2012; Saleh et al., 2013, 2014). However, several assumptions must be made on aerosol morphology (spherical Mie model) and mixing states of BC and organic aerosols (OA), which might introduce large uncertainties in the estimation of \( k \) (Mack et al., 2010; Xu et al., 2021).

To improve the understanding of chemical composition and light-absorbing properties of BrC chromophores, organic matter (OM) in aerosols was isolated through solvent extraction using water and/or methanol, followed by filtration and a series of instrumental analysis (e.g., UV–Vis spectrometer, liquid chromatograph–mass spectrometer; Chen and Bond, 2010; Liu et al., 2013; Lin et al., 2016). Referring to existing studies, a larger fraction of the methanol extract absorption comes from water-insoluble OM containing conjugated structures (Chen and Bond, 2010; Huang et al., 2020); the light absorption of biomass burning OM is majorly contributed by large molecules (molecular weight (MW) >500–1000 Da; Di Lorenzo and Young, 2016; Di Lorenzo et al., 2017) and depends on burn conditions (Saleh et al., 2014); and polycyclic aromatic hydrocarbons (PAHs) and nitroaromatic compounds (NACs) are ubiquitous BrC chromophores in the atmosphere (Huang et al., 2018; Wang et al., 2019), but the identified species only explain a small percentage (<10 %) of total BrC absorption (Huang et al., 2018; Li et al., 2020).

Methanol can extract >90 % OM from biomass burning (Chen and Bond, 2010; Xie et al., 2017b), while the extraction efficiency \( \eta \) decreases to ~80 % for ambient organic aerosols (Xie et al., 2019b, 2022), possibly due to other sources emitting large hydrophobic molecules and the oligomerization of small molecules during the aging process (Cheng et al., 2021; Li et al., 2021). The light-absorbing properties and structures of methanol-insoluble OC (MIOC) are still unknown. By comparing BrC characterization results of offline and online methods, some studies conclude that the MIOC dominates BrC absorption in source and ambient aerosols (Bai et al., 2020; Atwi et al., 2022). However, the online-retrieval and offline-extraction methods are designed based on different instrumentation and purposes, and the online method depends largely on presumed and uncertain optical properties of BC (Wang et al., 2014). Even when the solvent extract absorption is converted to particulate absorption with Mie calculations, pH and solvent matrix effects, as well as the potential incomplete solubility of BrC in common solvents, should still be considered before comparing BrC absorption measured directly in particles versus that derived from solvent extracts. To reveal the absorption and composition of MIOC, it is necessary to find a new solvent or develop a new methodology to improve OC extraction efficiency (Shetty et al., 2019).

In this work, a series of single solvents and solvent blends were tested for extraction efficiencies of OC in ambient particulate matter with aerodynamic diameter <2.5 \( \mu \text{m} \) (PM\(_{2.5}\)), and the sample extract absorption of each solvent was compared. The solvent or solvent mixture with the highest \( \eta \) value was applied to extract a matrix of collocated PM\(_{2.5}\) samples, followed by light absorption measurements. In our previous work, the light absorption of methanol extracts of the same samples was measured, and source apportionment was performed using organic molecular marker data (Xie et al., 2022). Through comparison with the study results in Xie et al. (2022), this study evaluated potential underestimation of BrC absorption in methanol and its impacts on BrC source attributions. These results suggest that different solvents should be used in future investigations on the absorption, composition, sources, and formation pathways of low-volatility BrC.

2 Methods

2.1 Solvent selection

Five solvents and solvent mixtures including water, methanol (MeOH), MeOH/dichloromethane (MeOH/DCM, 1 : 1, \( v : v \)), MeOH/DCM (1 : 2, \( v : v \)), tetrahydrofuran (THF), and N,N-dimethylformamide (DMF) were selected to extract OC from identical PM\(_{2.5}\) samples to determine which solvent or solvent mixture has the highest \( \eta \) value. Water and methanol are the most commonly used solvents to extract BrC from source or ambient particles. Cheng et al. (2021) found that OC produced through the combustion of toluene, isoctane, and cyclohexane was more soluble in DCM than MeOH. Since a major part of BrC absorption is coming from unknown large molecules (Di Lorenzo and Young, 2016; Di Lorenzo et al., 2017), polar aprotic solvents THF and DMF were tested due to their high capacity for dissolving large polymers. Except for water and MeOH, DCM and THF were rarely used to extract OC for light absorption measurements (Cheng et al., 2021; Moschos et al., 2021), and DMF has not ever been tested for extracting BrC in literature.
2.2 Sampling

- **Sampling for solvent test.** To compare OC extraction efficiencies and extract absorption of the five selected solvents and solvent mixtures, 21 ambient PM$_{2.5}$ samples were collected on the rooftop of a seven-story library building in Nanjing University of Information Science and Technology (NUIST; 32.21° N, 118.71° E). Details of the sampling site and equipment were provided by Yang et al. (2021). Two identical mid-volume samplers (Sampler I and II; PM$_{2.5}$–PUF-300, Mingye Environmental, China) equipped with 2.5 µm cut-point impactors were used for ambient air sampling during daytime (8:00–19:00) and nighttime (20:00–7:00 the next day), respectively, in December 2019. After the impactor, PM$_{2.5}$ in the air stream was collected on a pre-baked (550 °C, 4 h) quartz filter (20.3 cm × 12.6 cm, Munktell Filter AB, Sweden) at a flow rate of 300 L min$^{-1}$. PM$_{2.5}$ filter and field blank samples were sealed and stored at −20 °C before chemical analysis. Information about PM$_{2.5}$ samples for the solvent test is provided in Table S1 in the Supplement.

- **Ambient sampling for BrC analysis.** Details of the ambient sampling were described in previous work (Qin et al., 2021; Yang et al., 2021; Xie et al., 2022). Briefly, Sampler I and II were equipped with two quartz filters in series (quartz behind quartz (QBQ) method; Q$_b$ and Q$_f$) followed by adsorbents. Collocated filter and adsorbent samples were collected every sixth day during daytime and nighttime from 28 September 2018 to 28 September 2019. Field blank sampling was performed every 10th sample to address contamination. Q$_f$ samples loaded with PM$_{2.5}$ were.speciated and extracted for light absorption measurements. The OC adsorbed on Q$_b$ and its light absorption were analyzed to determine positive sampling artifacts. The adsorbents in Sampler I (a polyurethane foam (PUF)–XAD-4 resin–PUF sandwich) and II (a PUF plug) were used to collect gas-phase nonpolar and polar organic compounds, respectively. The measurement results of gas- and particle-phase organic compounds were provided by Gou et al. (2021) and Qin et al. (2021).

2.3 Solvent test for light absorption and extraction efficiency

An aliquot (~6 cm$^2$) of each filter sample was extracted ultrasonically in 10 mL of each solvent or solvent mixture (HPLC grade) for 30 min (one-time extraction procedure, N = 11; Table S1). After filtration, the light absorbance ($A_k$) of individual solvent extracts was measured over 200–900 nm using a UV–Vis spectrometer (UV-1900, Shimadzu Corporation, Japan) and was converted to a light absorption coefficient (Abs$_k$, Mm$^{-1}$) by

$$\text{Abs}_k = (A_k - A_{700}) \times \frac{V_l}{V_a} \times L \times \ln(10),$$  

where $A_{700}$ is subtracted to correct baseline drift, $V_l$ (m$^3$) is the air volume of the extracted sample, $L$ (0.01 m) is the optical path length, and ln (10) was multiplied to transform Abs$_k$ from a common to a natural logarithm (Hecobian et al., 2010). To understand if multiple extractions could draw out more BrC, a two-time extraction procedure was applied for another 10 ambient PM$_{2.5}$ samples in the same manner (Table S1). The $A_k$ of the first and second extractions (10 mL each) was measured separately for Abs$_k$ calculations.

Prior to solvent extractions, the concentrations of OC and elemental carbon (EC) in each filter sample were analyzed using a thermal-optical carbon analyzer (DRI, 2001A, Atmoslytic, United States) following the IMPROVE-A protocol. OC and EC were converted to CO$_2$ step by step during two separate heating cycles (OC1 (140 °C), OC2 (280 °C), OC3 (480 °C), and OC4 (580 °C) in pure He and EC1 (580 °C), EC2 (740 °C), and EC3 (840 °C) in 98 % He/2 % O$_2$), and the emitted CO$_2$ during each heating step was converted to CH$_4$ and measured using a flame ionization detector (FID).

After extractions, filters extracted by MeOH, MeOH/DCM (1:1), MeOH/DCM (1:2), and THF were air-dried in a fume hood and analyzed for residual OC (rOC; µg m$^{-3}$) using the identical method. Filters extracted in water and DMF cannot be air-dried in the short term due to the low volatility of solvents, and their rOC was measured after baking at 100 °C for 2 h. The total amount of OC dissolved in water for each sample was also measured as water-soluble OC (WSOC) by a total organic carbon analyzer (TOC-L; Shimadzu, Japan; Yang et al., 2021). To examine if the baking process would influence rOC measurements, the rOC of filters extracted in MeOH, MeOH/DCM mixtures, and THF was also measured after the baking process and compared to that determined after being air-dried. The pyrolytic carbon (PC) was used to correct for sample charring and was determined when the filter transmittance or reflectance returned to its initial value during the analysis (Schauer et al., 2003), but the formation of PC is very scarce when analyzing extracted filters. In this study, solvent-extractable OC (SEOC; µg m$^{-3}$) was determined by the difference in OC1–OC4 between pre- and post-extraction samples. The extraction efficiency ($\eta$, %) of each solvent was expressed as

$$\eta = \frac{\text{SEOC}}{\text{OC}} \times 100\%,$$  

Here, SEOC denotes WSOC when the solvent is water. For the ambient samples extracted twice, rOC was measured only after the two-extraction procedure was completed.

The solution mass absorption efficiency (MAE$_k$; m$^2$ g$^{-1}$ C) was calculated by dividing Abs$_k$ by the concentration of SEOC,
MAE₂ = \frac{Abs_2}{SEOC}, \quad (3)

and the solution absorption Ångström exponent (Å), a parameter showing the wavelength dependence of solvent extract absorption, was obtained from the regression slope of \( \log(\text{Abs}_b) \) versus \( \log(\lambda) \) over 300–550 nm.

The solvent effect is not uncommon when measuring aerosol extract absorbance in difference solvents (Chen and Bond, 2010; Mo et al., 2017; Moschos et al., 2021) but is rarely accounted for in previous studies. To evaluate the influence of solvent effects on light absorption of different solvent extracts of the same sample, solutions of 4-nitrophenol at 1.90 mg L\(^{-1}\), 4-nitrocatechol at 1.84 mg L\(^{-1}\), and 25-PAH mixtures (Table S2) at 0.0080 and 0.024 mg L\(^{-1}\) (each species) in the five solvents and solvent mixtures were made up five times and analyzed for UV–Vis spectra. The absorbance of PAH mixtures in water was not provided due to their low solubility.

2.4 Measurements and analysis of ambient BrC absorption

Collocated \( Q_f \) and \( Q_b \) samples were extracted using the solvent with the highest \( \eta \) value once followed by light absorbance measurement. OC concentrations in \( Q_f \) and \( Q_b \) samples were obtained from Yang et al. (2021), and SEOC values were estimated from OC concentrations and the average \( \eta \) value determined in Sect. 2.1 for one-time extraction. In this work, \( Q_b \) measurements were used to correct Abs\(_b\), MAE\(_b\), and Å of BrC in ambient PM\(_{2.5}\) in the same manner as those for water and methanol extracts in Xie et al. (2022):

\[
\text{artifact-corrected Abs}_b = Abs_b^{Q_f} - Abs_b^{Q_b}, \quad (4)
\]

\[
\text{artifact-corrected MAE}_b = \frac{Abs_b^{Q_f} - Abs_b^{Q_b}}{SEOC_{Q_f} - OC_{Q_b}}, \quad (5)
\]

where \( Abs_b^{Q_f} \) and \( Abs_b^{Q_b} \) are Abs\(_b\) values of \( Q_f \) and \( Q_b \) samples, respectively; \( SEOC_{Q_f} \) represents SEOC concentrations in \( Q_f \) samples; and \( OC_{Q_b} \) denotes OC concentrations in \( Q_b \) samples, assuming that OC in \( Q_b \) is completely dissolved (Xie et al., 2022). Artifact-corrected Å values were generated from the regression slope of \( \log(\text{Abs}_b^{Q_f} - \text{Abs}_b^{Q_b}) \) versus \( \log(\lambda) \) over 300–550 nm. Artifact-corrected Abs\(_b\), MAE\(_b\), and Å during each sampling interval were determined by averaging each pair of collocated measurements. If one of the two numbers in a pair is missed, the other number will be directly used for the specific sampling interval. To compare with previous studies based on water and/or methanol extraction methods, Abs\(_b\) and MAE\(_b\) at 365 nm were shown and discussed in this work.

Pearson’s correlation coefficient \( (r) \) was used to show how collocated measurements of BrC in ambient PM\(_{2.5}\) vary together. The coefficient of divergence \( (\text{COD}) \) was calculated to indicate consistency between collocated measurements. The relative uncertainty of BrC absorption derived from duplicate data was depicted using the average relative percent difference \( (\text{APRD}; \%) \), which was used as the uncertainty fraction for BrC measurements. Calculation methods of COD and ARPD are provided in Sect. S1 in the Supplement. To examine the influence of potential BrC underestimation based on the methanol extraction method on source apportionment, positive matrix factorization (PMF) version 5.0 (U.S. Environmental Protection Agency) was applied to attribute the light absorption of aerosol extracts in methanol and solvent with the highest \( \eta \) to sources. The total concentration data \( (Q_f + Q_b + \text{adsorbent}) \) of organic compounds have been used to apportion the light absorption of MeOH-soluble OC to specific sources (Xie et al., 2022), so as to avoid the impacts of gas–particle partitioning. In this work, the input particulate bulk components and total organic molecular marker \( (\text{OMM}) \) data for PMF analysis were obtained from Xie et al. (2022) and are summarized in Table S3. Solutions with 4 to 10 factors were tested to retrieve a final factor number with the most physically interpretable base-case solution. More information on input data preparation and the factor number determination is provided in the Supplement (Sect. S2 and Table S4).

3 Results and discussion

3.1 Solvent test

3.1.1 Extraction efficiency of different solvents

The concentrations of OC and EC fractions in each sample prior to solvent extractions are listed in Table S1. SEOC concentrations and extraction efficiencies of individual solvents and solvent mixtures are detailed in Table 1. Generally, DMF presented the highest extraction efficiency of total OC whenever filter samples were extracted once (89.0 ± 7.96 %) or twice (95.6 ± 3.67 %), followed by MeOH (one-time extraction 82.3 ± 8.68 %, two-time extraction 86.6 ± 7.86 %) and MeOH/DCM mixtures (∼75 %, ∼85 %). Although THF and DMF are frequently used to dissolve polymers (e.g., polystyrene) for characterization, THF had the lowest \( \eta \) values (64.2 ± 8.08 %, 70.1 ± 8.01 %) comparable to water (66.7 ± 8.58 %, 69.9 ± 5.88 %). Compared with one-time extraction, the extraction efficiencies of selected solvents were improved by a few percent when filter samples were extracted twice, and \( \eta \) values of MeOH/DCM mixtures became closer to those of MeOH (Table 1). These results showed that solvents can reach more than 80 % of their dissolving capacity with the one-time extraction, and the ambient OC in Nanjing is more soluble in MeOH than in DCM.

From OC1 to OC4, the volatility of OC fractions is expected to decrease continuously, and the molecules in OC fractions evolving at higher temperatures should be larger

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than those in OC1 with similar functional groups. In Table 1, MeOH and MeOH/DCM mixtures had comparable or even higher η values (82.6 ± 25.9 %–97.9 ± 5.02 %) of OC1 and OC2 than DMF (88.8 ± 4.98 %–97.2 ± 2.12 %). But OC3 and OC4 accounted for more than 60 % of OC concentrations, and DMF exhibited significant (p < 0.05) larger η values than other solvents, indicating that DMF had stronger dissolving capacity for large organic molecules than MeOH.

Concentrations of extracted OC fractions in MeOH, MeOH/DCM mixtures, and THF based on the two methods for rOC measurements (Sect. 2.2) are compared in Figs. S1 and S2. The total SEOC concentrations derived from the two methods are compared in Fig. S3. All the scatter data of SEOC fell along the 1:1 line with significant correlations (r > 0.85, p < 0.01). Because the measurement uncertainty of dominant species is lower than minor ones (Hyslop and White, 2008; Yang et al., 2021), the slightly greater relative difference between the two methods for extractable OC1 was likely attributed to its low concentrations (<1 μg m⁻³; Tables 1 and S1). Thus, baking extracted filters to dryness was expected to have little influence on SEOC measurements, particularly for low-volatility OC fractions (OC2-OC4).

Although water dissolves less OC than MeOH, WSOC is intensively extracted and analyzed for its composition and light absorption (Hecobian et al., 2010; Liu et al., 2013; Wagenfelder et al., 2015). WSOC can play a significant role in changing the radiative and cloud-nucleating properties of atmospheric aerosols (Haller et al., 2013; Taylor et al., 2017). It also served as a proxy measurement for oxygenated (OOA) or secondary organic aerosol (SOA) in some regions (Kondo et al., 2007; Weber et al., 2007). In previous work, MeOH was commonly used as the most efficient solvent in extracting OC from biomass burning (η > 90 %; Chen and Bond, 2010; Xie et al., 2017b) and ambient particles (η ~ 80 %; Xie et al., 2019b, 2022). MeOH-insoluble OC has rarely been investigated through direct solvent ex-

| Table 1. SEOC concentrations and extraction efficiencies (η, %) of total OC and OC fractions for different solvents. |
|-----------------------------------------------|
| OC prior to extractions | Water⁹ | MeOH⁶ | MeOH/DCM (1:1)⁶ | MeOH/DCM (1:2)⁶ | THF⁶ | DMF⁸ |
|-----------------------------------------------|
| One-time extraction (N = 11) |
| SEOC (μg m⁻³) | 9.36 ± 2.27 | 6.38 ± 2.03 | 7.85 ± 2.40 | 7.08 ± 1.32 | 6.99 ± 1.71 | 6.14 ± 2.01 | 8.49 ± 2.52 |
| Total OC | 0.66 ± 0.21 | 0.61 ± 0.20 | 0.64 ± 0.21 | 0.65 ± 0.20 | 0.64 ± 0.22 | 0.59 ± 0.18 | 0.59 ± 0.24 |
| OC1 | 2.69 ± 0.55 | 2.20 ± 0.60 | 2.50 ± 0.55 | 2.34 ± 0.41 | 2.37 ± 0.46 | 2.09 ± 0.55 | 2.48 ± 0.60 |
| OC2 | 3.35 ± 0.93 | 1.82 ± 0.80 | 2.48 ± 0.96 | 2.23 ± 0.49 | 2.18 ± 0.70 | 1.98 ± 0.93 | 2.86 ± 1.01 |
| OC3 | 2.75 ± 0.81 | 1.76 ± 0.65 | 2.23 ± 0.84 | 1.86 ± 0.51 | 1.78 ± 0.61 | 1.48 ± 0.61 | 2.56 ± 0.87 |
| OC4 | 66.7 ± 8.58 | 82.3 ± 8.68 | 76.0 ± 7.70 | 74.3 ± 7.83 | 64.2 ± 8.08 | 89.0 ± 7.96 |
| 91.7 ± 4.85 | 96.1 ± 6.73 | 97.9 ± 5.02 | 97.4 ± 4.35 | 89.6 ± 9.55 | 88.8 ± 4.98 |
| OC2 | 80.8 ± 8.11 | 92.7 ± 3.69 | 87.7 ± 5.87 | 88.5 ± 7.21 | 76.9 ± 7.62 | 91.4 ± 6.17 |
| OC3 | 52.4 ± 11.8 | 73.0 ± 11.5 | 68.1 ± 8.64 | 65.2 ± 10.2 | 57.6 ± 12.0 | 84.3 ± 9.79 |
| OC4 | 63.3 ± 9.13 | 80.3 ± 11.4 | 69.0 ± 9.26 | 64.5 ± 8.11 | 52.7 ± 5.86 | 92.8 ± 9.69 |
| Two-time extraction (N = 10) |
| SEOC (μg m⁻³) | 10.9 ± 4.93 | 7.74 ± 4.01 | 9.33 ± 4.11 | 9.34 ± 4.19 | 9.11 ± 4.04 | 7.56 ± 3.38 | 10.4 ± 4.80 |
| Total OC | 0.66 ± 0.47 | 0.62 ± 0.45 | 0.62 ± 0.49 | 0.59 ± 0.50 | 0.60 ± 0.51 | 0.59 ± 0.49 | 0.60 ± 0.47 |
| OC1 | 2.76 ± 0.77 | 2.20 ± 0.59 | 2.60 ± 0.66 | 2.57 ± 0.65 | 2.60 ± 0.68 | 2.28 ± 0.53 | 2.69 ± 0.78 |
| OC2 | 4.11 ± 2.01 | 2.55 ± 1.62 | 3.26 ± 1.62 | 3.37 ± 1.68 | 3.20 ± 1.58 | 2.62 ± 1.39 | 3.88 ± 1.95 |
| OC3 | 3.36 ± 1.77 | 2.38 ± 1.42 | 2.84 ± 1.42 | 2.81 ± 1.47 | 2.71 ± 1.39 | 2.08 ± 1.06 | 3.23 ± 1.70 |
| OC4 | 69.9 ± 5.88 | 86.6 ± 7.86 | 86.2 ± 8.73 | 84.8 ± 7.76 | 70.1 ± 8.01 | 95.6 ± 3.67 |
| OC1 | 93.6 ± 4.08 | 90.3 ± 13.9 | 82.6 ± 25.9 | 83.8 ± 22.4 | 82.9 ± 15.1 | 92.2 ± 13.9 |
| OC2 | 80.1 ± 5.01 | 94.8 ± 4.20 | 93.6 ± 4.94 | 94.7 ± 2.51 | 83.5 ± 6.86 | 97.2 ± 2.12 |
| OC3 | 59.0 ± 10.6 | 80.0 ± 10.2 | 82.3 ± 9.86 | 79.1 ± 10.6 | 63.9 ± 10.7 | 94.2 ± 4.15 |
| OC4 | 69.3 ± 6.46 | 86.3 ± 12.0 | 84.3 ± 12.0 | 82.7 ± 13.3 | 62.9 ± 7.76 | 96.9 ± 5.18 |

⁹ Concentrations of rOC in extracted filters were measured after the baking process (100°, 2 h). ¹⁰ rOC was measured when extracted filters were air-dried.
Table 2. Light-absorbing properties of SEOC following one-time and two-time extraction procedures.

| Solvent | Water | MeOH | MeOH/DCM (1:1) | MeOH/DCM (1:2) | THF | DMF |
|---------|-------|------|----------------|----------------|-----|-----|
|         |       |      |                |                |     |     |
| One-time extraction |       |      |                |                |     |     |
| Abs$^{365}$ (Mm$^{-1}$) | 5.13 ± 2.04 | 11.9 ± 5.83 | 10.3 ± 4.42 | 8.12 ± 3.38 | 5.48 ± 3.01 | 17.5 ± 8.05 |
| Abs$^{550}$ (Mm$^{-1}$) | 0.35 ± 0.12 | 1.28 ± 0.87 | 0.97 ± 0.55 | 0.35 ± 0.47 | 0.42 ± 0.47 | 4.40 ± 2.34 |
| MAE$^{365}$ (m$^2$ g$^{-1}$ C) | 0.87 ± 0.19 | 1.46 ± 0.41 | 1.41 ± 0.36 | 1.13 ± 0.22 | 0.87 ± 0.25 | 2.02 ± 0.58 |
| MAE$^{550}$ (m$^2$ g$^{-1}$ C) | 0.062 ± 0.028 | 0.15 ± 0.084 | 0.13 ± 0.054 | 0.042 ± 0.52 | 0.059 ± 0.56 | 0.30 ± 0.12 |
| Å | 6.63 ± 0.49 | 5.44 ± 0.75 | 5.65 ± 0.54 | 6.59 ± 0.66 | 6.17 ± 0.69 | 4.52 ± 0.41 |

| Two-time extraction |       |      |                |                |     |     |
|---------------------|-------|------|----------------|----------------|-----|-----|
| Abs$^{365,1st^{a}}$ (Mm$^{-1}$) | 6.64 ± 4.25 | 14.1 ± 7.09 | 14.6 ± 8.05 | 11.6 ± 6.78 | 7.17 ± 4.26 | 20.5 ± 10.6 |
| Abs$^{550,1st^{a}}$ (Mm$^{-1}$) | 0.42 ± 0.12 | 1.34 ± 0.70 | 1.34 ± 0.83 | 0.84 ± 0.50 | 0.53 ± 0.27 | 2.82 ± 1.44 |
| Abs$^{365^{b}}$ (Mm$^{-1}$) | 8.26 ± 5.21 | 15.5 ± 7.76 | 16.8 ± 8.82 | 14.0 ± 8.91 | 8.35 ± 4.81 | 21.9 ± 11.2 |
| Abs$^{550^{b}}$ (Mm$^{-1}$) | 0.50 ± 0.18 | 1.60 ± 0.78 | 1.64 ± 0.99 | 1.22 ± 0.98 | 0.69 ± 0.43 | 3.01 ± 1.49 |
| MAE$^{365}$ (m$^2$ g$^{-1}$ C) | 1.19 ± 0.26 | 1.70 ± 0.60 | 1.80 ± 0.52 | 1.50 ± 0.51 | 1.10 ± 0.40 | 2.11 ± 0.49 |
| MAE$^{550}$ (m$^2$ g$^{-1}$ C) | 0.082 ± 0.30 | 0.19 ± 0.11 | 0.17 ± 0.083 | 0.13 ± 0.069 | 0.094 ± 0.054 | 0.29 ± 0.075 |
| Å | 6.32 ± 0.58 | 5.37 ± 0.57 | 5.47 ± 0.67 | 5.57 ± 0.39 | 6.06 ± 0.54 | 4.53 ± 0.21 |

$^{a}$ Light absorption coefficient of SEOC after the first extraction. $^{b}$ Sum of SEOC absorption in first and second extracts.

Light absorption of different solvent extracts

Table 2 shows the average Abs$_{365}$ and MAE$_{365}$ values of different solvent extracts at 365 and 550 nm. The Abs$_{365}$ and MAE$_{365}$ spectra of selected samples are illustrated in Fig. S4. Not including DMF, MeOH extracts exhibited the strongest light absorption. Since MeOH can dissolve more OC3 and OC4 than DCM (Table 1), the Abs$_{365}$ and MAE$_{365}$ of MeOH/DCM extracts decreased as the fraction of DCM increased in solvent mixtures (Table 2 and Fig. S4). Water and THF extracts had the smallest Abs$_{365}$ and MAE$_{365}$ due to their low extraction efficiencies for low-volatility OC (OC2-OC4; Table 1). In comparison to MeOH extracts, Abs$_{365}$ and MAE$_{365}$ of DMF extracts were at least more than 40% higher ($p<0.05$). Given that the relative difference in extraction efficiency of total OC between MeOH and DMF was less than 10%, and DMF dissolved more OC3 and OC4 than other solvents (Table 1), low-volatility OC should contain stronger light-absorbing chromophores (Saleh et al., 2014), and its mass fraction might determine the difference in BrC absorption across solvent extraction methods. Moreover, the relative difference in Abs$_{365}$ and MAE$_{365}$ between MeOH and DMF extracts increased with wavelength (Table 2 and Fig. S4). This is because the light absorption of DMF extracts that contain stronger BrC chromophores depends less on wavelengths than other solvent extracts ($\lambda \sim 4.5$, Table 2). As shown in Fig. S5, average Å and MAE$_{365,550}$ values of individual solvent extracts in Table 2 are negatively correlated.

In this work, insoluble organic particles coming off the filter during sonication might lead to underestimated SEOC concentrations and $\eta$ values, and then the MAE$_{365}$ of solvent extracts would be underestimated. Previous studies rarely considered the loss of insoluble OC during the extraction process (Yan et al., 2020), of which the impact on MAE$_{365}$ calculation was still inconclusive. But Abs$_{365}$ measurements would never be influenced, as the light absorbance of solvent extracts was analyzed after filtration. In Table 2, the second extraction only increases the average Abs$_{365}$ and Abs$_{550}$ values of DMF extracts by 6.70% ($p = 0.78$) and 6.76% ($p = 0.77$), respectively. We suspected that the difference in $\eta$ values of DMF between one-time and two-time extraction procedures was mainly ascribed to the detachment of insoluble OC particles.

In Fig. S6, the absorbance spectra of 4-nitrophenol and 4-nitrocatechol in water shift toward longer wavelengths compared to their MeOH solution. This is because neutral and deprotonated forms of 4-nitrophenol and 4-nitrocatechol may have different absorbance spectra, and these two compounds are deprotonated at pH $\approx 7$ (Lin et al., 2015b, 2017). The strong light absorption of 4-nitrophenol and 4-nitrocatechol in DMF at 450 nm was not observed in other solvents and was likely caused by unknown reactions. Then the solvent effect introduced by DMF might overestimate the light absorption of low-molecular-weight (LMW) nitrophenol-like species at $>400$ nm in source or ambient aerosols. Evidence shows that BrC absorption is dominated by large molecules.
with extremely low volatility (Saleh et al., 2014; Di Lorenzo and Young, 2016; Di Lorenzo et al., 2017), and LMW nitrophenol-like species have very low contributions to particulate OM (e.g., <1 %) and aerosol extract absorption (e.g., <10 %) (Mohr et al., 2013; Zhang et al., 2013; Teich et al., 2017; Xie et al., 2019a, 2020; Li et al., 2020). The shapes of the light absorption spectra of aerosol extracts in DMF were similar to other solvents (Fig. S4) and PAH solutions (Fig. S6g–l), and no elevation in light absorption appeared at 400–500 nm. Thus, the overestimated absorption of LMW nitrophenol-like species in DMF might not substantially impact the overall BrC absorption of aerosol extracts. Furthermore, the absorbance of 4-nitrophenol and 4-nitrocatechol in DMF at 365 nm (A$_{365}$) was lower than that in MeOH, and PAH solutions showed very similar absorbance spectra across the five solvents (Fig. S6g–l and Table S5). Considering the fact that low-volatility OC fractions (e.g., OC3 and OC4) in the ambient are less water-soluble (Table 1) and have a high degree of conjugation (Chen and Bond, 2010; Lin et al., 2014), their structures probably feature a PAH skeleton. Therefore, the large difference in Abs$_{365}$ between DMF and MeOH extracts (Table 2) was primarily ascribed to the fact that DMF can dissolve more OC3 and OC4 than methanol (Table 1). However, we cannot rule out the impact of solvent effects on the comparison of light absorption spectra between MeOH and DMF extracts (Fig. S4), and more work is warranted in identifying the structures more soluble in DMF than in MeOH.

### Table 3. Comparisons of light-absorbing properties of ambient PM$_{2.5}$ extracts in DMF and MeOH derived from duplicate Q$_f$–Q$_b$ data ($N = 109$).

|                | DMF          | MeOH*         |
|----------------|--------------|---------------|
|                | Median | Mean ± SD | Range | Median | Mean ± SD | Range |
| Abs$_{365}$ (Mm$^{-1}$) | 6.99  | 8.42 ± 5.40 | 1.14–30.8 | 5.59  | 6.43 ± 4.66 | 0.38–29.6 |
| MAE$_{365}$ (m$^2$ g$^{-1}$ C$^{-1}$) | 1.13  | 1.20 ± 0.49 | 0.34–2.45 | 0.91  | 1.03 ± 0.58 | 0.089–2.49 |
| Å              | 5.21   | 5.25 ± 0.64 | 3.21–6.82 | 6.49  | 6.81 ± 1.64 | 4.34–11.3  |

* Data for MeOH extracts were obtained from Xie et al. (2022).

### Figure 1. Comparisons between collocated measurements for light-absorbing properties of PM$_{2.5}$ extracts in DMF after Q$_b$ corrections.

3.2 Collocated measurements and temporal variability

Abs$_{365}$ values of collocated Q$_f$ and Q$_b$ extracts in DMF are summarized in Table S6. No significant difference was observed (Q$_f$ p = 0.96; Q$_b$ p = 0.42) between the two samplers. After Q$_b$ corrections, Abs$_{365}$, MAE$_{365}$, and Å of DMF-extractable OC (Abs$_{365,d}$, MAE$_{365,d}$, and Å$_{d}$) in PM$_{2.5}$ were calculated by averaging each pair of duplicate Q$_f$–Q$_b$ data and are compared with those of methanol extracts (Abs$_{365,m}$, MAE$_{365,m}$, and Å$_{m}$) in Table 3. Fig. 1 shows comparisons between collocated measurements of Abs$_{365,d}$, MAE$_{365,d}$, and Å$_{d}$. Generally, all comparisons indicated good agreement with COD <0.20 (0.094–0.15). Abs$_{365,d}$ and MAE$_{365,d}$ had comparable uncertainty fractions (ARPD, 22.7 % and 24.5 %, Fig. 1) to Abs$_{365,m}$ and MAE$_{365,m}$ (28.4 % and 28.8 %; Xie et al., 2022). Since different primary combustion sources can have similar spectral dependence for BrC absorption (Chen and Bond, 2010; Xie et al., 2017b, 2018, 2019a), most Å$_{d}$ data clustered on the identity line, with much lower variability than Abs$_{365,d}$ and MAE$_{365,d}$. As shown in Table 3, average Abs$_{365,d}$ and MAE$_{365,d}$ values were 30.7 % (p<0.01) and 17.3 % (p<0.05) larger than average Abs$_{365,m}$ and MAE$_{365,m}$. Because the k value of BrC in bulk solution is directly estimated from Abs$_{3}$ or MAE$_{3}$ (Liu et al., 2013, 2016; Lu et al., 2015), the estimation method needs to be revised when ambient BrC is extracted using DMF instead of MeOH. Both MAE$_{365,d}$ and MAE$_{365,m}$ were negatively correlated (p<0.01) with their corresponding Å values (Fig. S7), and the lower average Å$_{d}$ (5.25 ± 0.64,
Figure 2. Time series comparisons of light-absorbing properties of DMF and MeOH extracts using artifact-corrected data. MeOH extract data were obtained from Xie et al. (2022).

3.3 Sources of DMF- and MeOH-extractable BrC

A final factor number of eight was determined based on the interpretability of different base-case solutions (4 to 10 factors), the change in $Q/Q_{\text{exp}}$ with factor numbers, and robustness analysis (Sect. S2 and Table S4). Normalized factor profiles of seven- to nine-factor solutions are compared in Fig. S8. The seven-factor solution failed to resolve the lubricating oil combustion factor characterized by hopanes and steranes (Fig. S8c). An unknown factor containing various source tracers related to crustal dust ($\text{Ca}^{2+}$ and $\text{Mg}^{2+}$), lubricating oil (hopanes and steranes), and soil microbiota (sugar and sugar alcohols) was identified in the nine-factor solution (Fig. S8i). Median and mean values of input $\text{Abs}_{365,\text{d}}$, $\text{Abs}_{365,\text{m}}$, and bulk component concentrations agreed well with PMF estimations (Table S7), and the strong correlations ($r = 0.86–0.99$) between observations and PMF estimations indicated that the eight-factor solution simulated the time series of input species well. In comparison to Xie et al. (2022), where $\text{Abs}_{365}$ of MeOH and water extracts were apportioned to nine sources using the same speciation data, this work lumped secondary nitrate and sulfate into the same factor (termed “secondary inorganics”; Fig. S8h), and the other seven factors had similar factor profiles linked with biomass burning, non-combustion fossil, lubricating oil combustion, coal combustion, dust resuspension, biogenic emission, and isoprene oxidation. Interpretations of individual factors based on characteristic source tracers and contribu-
The average relative contributions of the identified factors to $\text{Abs}_{365,d}$, $\text{Abs}_{365,m}$, and bulk components are listed in Table S8. Consistent contribution distributions of $\text{Abs}_{365,m}$ were observed between Xie et al. (2022) and this study, indicating that the PMF results were robust to the inclusion of $\text{Abs}_{365,d}$ data. Figure 3 compares the time series of factor contributions to $\text{Abs}_{365,d}$ and $\text{Abs}_{365,m}$. ARPD and COD values between factor contributions to $\text{Abs}_{365,d}$ and $\text{Abs}_{365,m}$ and the absolute difference are exhibited in Fig. S9. $\text{Abs}_{365,d}$ and $\text{Abs}_{365,m}$ had comparable contributions from biomass burning, lubricating oil combustion, and coal combustion (Fig. 3a, c, d). The small COD values of these three factors (0.0041–0.17) indicated no significant divergence. The biogenic emission and isoprene oxidation factors exhibited complete difference (ARPD = 200 %, COD = 1; Fig. S9f, g) as they had no contribution to $\text{Abs}_{365,m}$. Among the eight factors, the non-combustion fossil, dust resuspension, and isoprene oxidation factors had the largest median difference in factor contributions to $\text{Abs}_{365,d}$ and $\text{Abs}_{365,m}$ (0.63–0.67 Mm$^{-1}$) with substantial heterogeneity (COD > 0.20).
followed by the secondary inorganics factor (0.20 Mm$^{-1}$, COD = 0.41). The temporal variations of the absolute difference shown in Fig. S9 are identical to the contributions of individual factors to $\text{Abs}_{365,d}$ or $\text{Abs}_{365,m}$ (Fig. 3).

The non-combustion fossil factor represents unburned fossil-fuel emissions (e.g., petroleum products), which contain substantial large organic molecules (e.g., high MW PAHs; Simoneit and Fetzer, 1996; Mi et al., 2000). This might explain why the non-combustion fossil factor contributed more $\text{Abs}_{365,d}$ than $\text{Abs}_{365,m}$ throughout the year (Fig. S9b). Dust resuspension and isoprene oxidation factors show prominent contributions to $\text{Abs}_{365,d}$ in spring and summer, respectively (Fig. 3e, g). The dust resuspension factor had the highest average contributions to both crustal materials (Ca$^{2+}$ and Mg$^{2+}$) and carbonaceous species (OC and EC; Table S8 and Fig. S8) and was considered a mixed source of crustal dust and motor vehicle emissions (Yu et al., 2020; Xie et al., 2022). Besides the influences from primary emissions, aging processes of organic components in dust aerosols can induce the formation of BrC through iron-catalyzed polymerization (Link et al., 2020; Al-Abadleh, 2021; Chin et al., 2021). It was demonstrated that the isoprene-derived polymerization products through aerosol-phase reactions are light-absorbing chromophores (Lin et al., 2014; Nakayama et al., 2015). This might explain the elevated difference between $\text{Abs}_{365,d}$ and $\text{Abs}_{365,m}$ contributions of the isoprene oxidation factor in summer (Fig. S9g). The biogenic emission factor was characterized by tracers related to microbiota activities (sugar and sugar alcohols) and decomposition of high plant materials (odd-numbered alkanes) in soil (Rogge et al., 1993; Simoneit et al., 2004) and had negligible contributions (<0.1%) to $\text{Abs}_{365,d}$ and $\text{Abs}_{365,m}$. Evidence shows that secondary BrC can be generated through gas-phase reactions of anthropogenic volatile organic compounds with NO$_x$ (Nakayama et al., 2010; Liu et al., 2016; Xie et al., 2017a), aqueous reactions of SOA with reduced nitrogen-containing species (e.g., NH$_3^+$; Updyke et al., 2012; Powelson et al., 2014; Lin et al., 2015a), and evaporation of water from droplets in the atmosphere containing soluble organics (Nguyen et al., 2012; Kasthuriarachchi et al., 2020). These processes can also lead to the formation of low-volatility oligomers (Nguyen et al., 2012; Song et al., 2013), and their contributions might be lumped into the secondary inorganics factor due to the lack of OMMs. According to these results, one possible explanation for the difference in time series between $\text{Abs}_{365,d}$ and $\text{Abs}_{365,m}$ (Fig. 2) is that large BrC molecules from unburned fossil fuels and atmospheric processes are less soluble in MeOH than in DMF.

4 Conclusions and implications

Comparisons of extraction efficiencies and light absorption of ambient aerosol extracts across selected solvents and solvent mixtures indicate that MeOH may sometimes be replaced with DMF for measuring BrC absorption, as low-volatility OC fractions containing strong chromophores are less soluble in MeOH than in DMF. Existing modeling studies on the radiative forcing of BrC (Feng et al., 2013; Wang et al., 2014; Zhang et al., 2020) often retrieved or estimated its optical properties from laboratory or ambient measurements based on water/methanol extraction methods (Chen and Bond, 2010; Hecobian et al., 2010; Liu et al., 2013; Zhang et al., 2013) and had a potential to underestimate the contribution of BrC to total aerosol absorption. However, the influence of the solvent effect was not accounted for in this work when comparing the light absorption of different solvent extracts. The difference between MeOH and DMF extract absorption might change with time and location due to the variations in BrC sources. The results of this work also imply the necessity of applying different solvents or combinations of solvents with broad polarity and dissolving capability to study BrC composition and absorption, particularly for low-volatility fractions.

Although light-absorbing properties of DMF and MeOH extracts had good agreement in cold periods, when biomass and coal burning sources dominated BrC emissions, their distinct time series in spring and summer implies that the contributions of certain BrC sources were underestimated or missed when the MeOH extraction method was used. Source apportionment results of $\text{Abs}_{365,d}$ and $\text{Abs}_{365,m}$ based on organic molecular marker data indicated that large and methanol-insoluble BrC molecules are likely coming from unburned fossil fuels and polymerization of aerosol organics. Laboratory studies have observed the polymerization process through heterogeneous reactions of several precursors (e.g., catechol; Lin et al., 2014; Link et al., 2020), but the structures and light-absorbing properties of potential polymerization products in ambient aerosols (Fig. 3e, g) are less understood and warrant further study.

Data availability. Data used in the writing of this paper are available in the Harvard Dataverse (https://doi.org/10.7910/DVN/CGHPXB, Xu et al., 2022)

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