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Unraveling the complexity of atmospheric brown carbon produced by smoldering boreal peat using size-exclusion chromatography with selective mobile phases†

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Boreal wildfires are a significant source of atmospheric brown carbon (BrC), a complex mixture of thousands of light-absorbing organic compounds that contributes to the warming effects of combustion particulate matter. Here, we use size-exclusion chromatography (SEC) coupled with photodiode array detection to characterize BrC collected from the controlled combustion of boreal peat. Importantly, rather than attempting to estimate the molecular weight of BrC chromophores through the minimization and correction of secondary interactions, we instead exploit these interactions to systematically explore BrC hydrophobicity, lability, and size-dependent light absorption properties. Using this new approach, which we corroborate using independent asymmetric flow field-flow fractionation (AF4) analysis, we show that the components of fresh wildfire BrC span a wide range of sizes, polarities, and light absorption characteristics. Unlike atmospherically aged wildfire BrC, which has previously been shown to resemble terrestrial humic substances in both its absorption profile and its retention behaviour, the fresh BrC sample studied here contains both higher-MW chromophores with “humic-like” featureless absorption and smaller-MW chromophores with structured absorption, and is more susceptible to hydrophobic interactions with the column matrix. Interestingly, we find that the contribution of the low-MW fraction to overall BrC absorption increases with increasing mobile phase acetonitrile content, which suggests that the high-MW fraction consists of metastable aggregates held together by easily disrupted intermolecular forces. Together, these results highlight the compositional diversity of atmospheric BrC and the challenge and potential of SEC for the characterization of complex, and poorly defined, environmental matrices.

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Environmental significance

Atmospheric brown carbon (BrC) is the dominant light-absorbing component of wildfire particulate matter. Our ability to quantify the climate impact of BrC emissions is currently limited by gaps in our understanding of its composition, properties, and atmospheric fate. Here, we use size-exclusion chromatography to examine the size, polarity, and lability of fresh BrC produced by the combustion of boreal peat. Through the careful examination of solvent-dependent elution behaviour, we show that this BrC sample is more diverse in both size and polarity than atmospherically aged BrC examined in previous studies, and that its large-MW components consist largely of smaller units aggregated together by weak intermolecular interactions. These results demonstrate the importance of studying the composition and light-absorbing properties of BrC over its entire atmospheric lifetime, as well as the need for careful consideration of extraction and analysis conditions when studying this complex class of atmospheric particulate matter.

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† Electronic supplementary information (ESI) available: A comprehensive discussion of our SEC column characterization results; selection/design of quality control experiments; details regarding the collection and extraction of a complementary BrC sample collected on quartz fiber filters (QFF); a description of our integration procedure for SEC absorption density plots;

and a summary of asymmetric flow field-flow fractionation (AF4) analysis conditions. Eight supporting figures, which present SEC calibration curves; supplementary characterization information for SRHA; SEC absorption density plots for quality control samples, including the QFF sample described above; and complementary AF4 characterization of our BrC sample. Two supporting tables, which present calibration curve equations (and associated SRHA MW estimates) and retention time reproducibility data for acetone and SRHA. See DOI: 10.1039/d1ea00011j

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1 Introduction

Combustion processes release large amounts of carbonaceous particulate matter (PM) into the atmosphere,^{1,2} which both absorbs and scatters solar radiation and influences global climate in complex ways.^{3,4} Light-absorbing carbonaceous PM can be divided into two categories: soot, also known as black carbon (BC), which absorbs light efficiently from the UV region to the infrared;⁵ and a complex mixture of organic compounds known as brown carbon (BrC), which exhibits a sharp decrease in absorption with increasing wavelength from the UV region to the visible.^{6,7} Although measurements and models have suggested that the warming effect of BrC is substantial,^{8,9} inclusion of BrC light absorption in climate simulations is challenging,¹⁰ primarily because of the complexity of BrC formation^{11,12} and light absorption mechanisms,^{13,14} chemical composition,¹⁵ and atmospheric fate.^{16,17}

One major BrC source is wildfires, which directly release BrC as a result of incomplete combustion and indirectly lead to BrC formation *via* atmospheric photolysis, oxidation, and gas-to-particle partitioning of co-emitted volatile organic compounds (VOCs).¹⁸ An understanding of the molecular characteristics of BrC chromophores in wildfire PM, as well as their susceptibility to chemical transformations in the atmosphere,^{19,20} is necessary for the accurate prediction of the overall climate impact of wildfires. Measured increases in the frequency and intensity of boreal wildfire activity,²¹ coupled with projected increases in extreme fire weather²² and overall wildfire risk,²³ lend particular urgency to these investigations.

Molecular-level BrC characterization is usually accomplished *via* offline analysis of PM extracts using high-performance liquid chromatography (HPLC) coupled with photodiode array (PDA) and high-resolution mass spectrometric (HRMS) detection; however, summing the contribution of chromophores identified by MS (*e.g.*, N-containing aromatic compounds, aromatic carbonyls, and aromatic carboxylic acids) typically cannot account for all BrC absorption.^{24–29} This discrepancy likely reflects qualitative and quantitative challenges associated with the application of HPLC-PDA-HRMS to the analysis of this highly complex yet poorly characterized substrate, including the need to account for (largely unknown) variations in component ionization efficiency,^{25,30} distinguish which of a set of co-eluting compounds is responsible for HPLC-PDA absorption signals in complex chromatograms,²⁴ and apply multiple ionization methods to achieve more comprehensive analyte coverage.²⁵ In addition, the analysis of the high-molecular-weight (MW) component of BrC (often referred to as “humic-like substances”, or HULIS), which consists of macromolecular aggregates composed of highly conjugated aromatic structures with polar functional groups containing oxygen and nitrogen (*e.g.*, carboxyl and nitro groups),³¹ may be complicated by fragmentation in the MS³² and by multiple charging effects.³³

A promising technique for the analysis of the high-MW component of BrC is size-exclusion chromatography (SEC),³⁴ which has been widely employed to estimate the MW and MW distributions of dissolved organic matter (DOM),³⁵ humic substances,³⁶ and the HULIS fraction of ambient atmospheric

PM.^{37,38} In recent years, this technique has also been coupled with PDA detection to estimate the MW and characterize the light absorbing properties of BrC chromophores in ambient^{39–41} and laboratory-generated^{42,43} PM samples. One major challenge associated with SEC separation is the hydrophobic and/or electrostatic interaction of analytes with the column matrix, which can lead to changes in elution profiles and thus introduce large uncertainties to MW estimates.³⁴ Additional uncertainties arise from intra- and intermolecular interactions, which can influence analyte hydrodynamic volume (*e.g.*, *via* coiling, expansion, and aggregation/agglomeration), thereby leading to either early or delayed elution and corresponding overestimates or underestimates in MW.^{36,44} Although the challenges associated with these secondary effects have been acknowledged in the broader atmospheric literature,^{45,46} studies of BrC using this technique have largely assumed that size exclusion is the dominant separation mechanism. However, as BrC consists of a complex mixture of compounds with a wide range of polarities and sizes,⁷ and with the potential to undergo aggregation and/or dissociation with changes in solvent environment,^{47,48} this assumption warrants critical assessment.

In this study, we used HPLC-SEC-PDA to characterize BrC produced by combustion of boreal peat, an important yet poorly understood source of biomass burning PM.^{49,50} Importantly, rather than attempting to obtain accurate absolute MW estimates for BrC samples through the use of mobile phase solvents chosen to minimize secondary effects, we instead exploited these effects to systematically investigate the hydrophobicity and lability of the samples. In particular, we examined their elution behaviour as a function of mobile phase composition, including organic solvent content and ionic strength, and compared this behaviour to that of Suwannee River Humic Acid (SRHA), which has been previously used as a proxy for biomass burning BrC.³⁹ In addition, we compared these results to those obtained using a complementary technique, asymmetric flow field-flow fractionation (AF4).^{51–53} Our results highlight the complex nature of secondary interactions in the analysis of fresh wildfire BrC using SEC and, at the same time, show that a sound understanding of these interactions can be used to provide new insights into the properties of this important PM class, including distributions of molecular size and polarity. Using this new approach, this study proceeds to reveal the aggregation and dissociation of water-soluble BrC components for the first time.

2 Methods

2.1 Boreal peat sampling

Boreal peat was sampled in the traditional territory of the Bigstone Cree Nation (Calling Lake, northern Alberta, Canada; 55.092, –113.272) in June 2018. To capture the typical depth of burn (~10 cm) during wildfires,⁵⁴ the bulk peat was sampled from the ground surface to ~40 cm. A BrC sample generated from the combustion of the peat surface layer (0–5 cm) was used in this study; the remaining samples were used for a comprehensive study of the influence of sample depth and moisture content on the light absorption properties of BrC generated from peat combustion.⁵⁵



2.2 Peat combustion and BrC collection

Biomass burning PM was generated from the peat sample described above at the laboratory combustion facility in the Northern Forestry Centre (Edmonton, Alberta). The peat sample was ignited by placing a quartz tube infrared electric heater (210 V, 1000 W; Re-Verber-Ray BAH-25) operating at ~ 800 °C over it for 1 min. The combustion phase (*i.e.*, flaming *versus* smoldering) was estimated using the modified combustion efficiency ($MCE = \Delta CO_2 / (\Delta CO_2 + \Delta CO)$),⁵⁶ which was calculated using concentrations of CO (ULTRAMAT 23, Siemens) and CO₂ (ULTRAMAT 6, Siemens) measured using an inlet at the intake of the overhead exhaust, ~ 3 m above the sample; here, this value was 0.84, which is consistent with smoldering combustion.⁵⁷

Water-soluble BrC was collected ~ 1 m above the combustion site using a particle-into-liquid sampler (PILS; Model 4001, Brechtel) system equipped with an inlet for the selective collection of particles with diameters smaller than 2.5 μm (PM_{2.5}), an activated carbon VOC denuder, and an auto-collection system (3.5 min per sample).⁵⁸ The aerosol sampling volumetric flow rate was 13.5–13.7 L min⁻¹; the quartz impactor plate wash flow rate was 0.42–0.46 mL min⁻¹. PILS sample vials were stored at 277 K until analysis, at which time samples were diluted ($\times 3$) with deionized water (18 M Ω) and filtered (PTFE, 0.2 μm , 13 mm, Fisherbrand Basix).

2.3 HPLC-SEC-PDA analysis

The system adopted in this paper is based on the previous work of Di Lorenzo *et al.*^{39–41} A HPLC (Agilent 1100) system equipped with an aqueous gel filtration column (300 \times 7.8 mm, 75 000–250 Da, Polysep GFC P-3000, Phenomenex) and coupled to a PDA (Agilent G1315B) detector was employed for the separation and detection of samples. The mobile phase consisted of phosphate buffer (NaH₂PO₄ + Na₂HPO₄, pH = 6.8, 20–100 mM) mixed with varying concentrations of acetonitrile (ACN) and/or methanol (MeOH) as organic modifier; all mixing ratios of buffer and organic phase are reported here as volume percentages, and all separations were performed under isocratic conditions. The flow rates and column temperatures employed were either 1 mL min⁻¹ and 40 °C (50% and 25% ACN) or 0.8 mL min⁻¹ and 55 °C (50% MeOH, 40% MeOH/10% ACN), with conditions selected to avoid overpressure of the column. Sample-to-sample reproducibility was assessed using the retention times of acetone and SRHA (Table S2[†]); on a given day, the variation in retention time for each marker was < 0.015 min. The sample injection volume was 15 μL .

2.4 Asymmetric flow field-flow fractionation (AF4) analysis

To measure the size-resolved optical properties of BrC in the absence of a stationary phase and thereby verify that our results were not specific to SEC analysis, BrC and Suwannee River natural organic matter (SRNOM; International Humic Substances Society) were also analyzed using AF4 (AF2000 Multiflow, Postnova Analytics) with UV-Vis absorbance and fluorescence detection (G4212 DAD and G1321B FLD, Agilent). Fluorescence emission (280–450 nm, excitation at 230 nm) was

collected with < 1 s time resolution over the course of the separation. The relationship between retention time and molecular mass was calibrated before and after the sample separations using a mixture of poly(styrene sulfonate) sodium salts (Polymer Standards) and a molecular standard (bromophenol blue; Sigma Aldrich) that together spanned the range of 0.67–20.7 kDa. This relationship was used to determine the molecular mass at peak maximum (M_p) of the BrC and SRNOM, measured at an absorbance wavelength of 254 nm. The AF4 flow program was optimized to maximize resolution in the lower size range, and a polyethersulfone (PES) membrane with the smallest available pore size (300 Da) was used to maximize retention of the smallest molecules. The carrier fluid consisted of an ammonium carbonate buffer (2 mM, pH 8). Additional details regarding the analysis method are published elsewhere.⁵³

2.5 Materials and chemicals

Suwannee River humic acid (SRHA; analyzed as a saturated aqueous solution) was obtained from the International Humic Substances Society. MeOH (Optima grade, 99.9%), ACN (Optima grade, 99.9%), acetone (ACS grade), and monosodium phosphate (ACS grade, 100.2%) were purchased from Fisher Chemical. Dibasic sodium phosphate (ACS grade, 99.0%) was purchased from Sigma Aldrich. Sodium poly(styrene sulfonate) standards for SEC analysis (PSS; M_w : 1690 Da, 5580 Da, 7540 Da, 16 kDa, 33.4 kDa, 68.3 kDa, 78.4 kDa) were purchased from Scientific Polymer Products, Inc.; aqueous standard solutions (~ 50 mg L⁻¹) were stored at 277 K until use. Deionized water (18.2 M Ω) was obtained from a Millipore Synergy UV ultrapure water system.

2.6 Safety statement

The primary hazard associated with this work was the peat combustion process itself; however, this was carried out in a dedicated combustion facility, which minimized risk.

3 Results and discussion

3.1 Challenges associated with application of SEC to absolute MW determinations

Given the chemical complexity of BrC, we first characterized our size-exclusion column using SRHA as a BrC proxy;³⁹ PSS (M_w : 1690 Da to 78.4 kDa), a linear anionic polyelectrolyte that has a similar charge density to NOM,⁵⁹ as calibrant; and acetone as a marker for the total permeation volume (V_t). Retention times for SRHA, PSS standards, and acetone are presented in Table S1.[†]

As discussed in the Introduction, SEC MW estimates are subject to biases from hydrophobic and/or electrostatic interactions of analytes with the column matrix; these biases can be mitigated using organic and/or ionic mobile phase modifiers,³⁴ which suppress analyte–column interactions. In order to assess the importance of secondary interactions in this study, therefore, we prepared PSS calibration curves at a range of phosphate buffer (20–100 mM, pH 6.8) and organic modifier (0–50% ACN, 0–10% MeOH) concentrations. As shown in Fig. S1 and Table S1,[†] the elution behaviors of the PSS standards and SRHA



responded differently to modifications in mobile phase composition; as a result, the estimated MW values of SRHA over our range of mobile phase compositions ranged from ~ 1000 to $>10\,000$ Da, despite being obtained using mobile phase-specific PSS calibration curves.

In summary, because SEC separation is based on analyte hydrodynamic volume rather than MW and ideal analyte behaviour is often compromised by non-size effects, accurate estimation of MW using SEC requires the selection of calibrants that are chemically and structurally similar to the analytes of interest.^{34,38} Owing to their different molecular properties, PSS and SRHA are subject to different non-size effects and undergo different conformational changes in response to mobile phase compositional changes; we discuss this in detail in the ESI.† As a result, although analysis conditions can be optimized for calibrant and analyte separately, conditions that are near-ideal for both are challenging to find. This challenge is magnified for BrC, which is a complex mixture of hundreds to thousands of organic compounds⁷ that differ in their three-dimensional structure, charge density, and hydrophobicity. Given these limitations, we do not use our SEC results to provide BrC MW estimates; instead, as described in the following sections, we use the identity and magnitudes of non-size effects observed under different mobile phase conditions to gain insight into BrC composition and properties.

3.2 Insights into BrC composition and properties from changes in mobile phase composition

3.2.1 Fresh peat combustion BrC differs from Suwannee River humic acid and ambient biomass burning BrC.

Laboratory studies have shown that fresh PM emitted by the combustion of Alaskan duff core⁶⁰ and Indonesian peat⁶¹ is weakly to non-hygroscopic. In this context, we hypothesized that our BrC sample, which was collected immediately above the combustion source, would be susceptible to hydrophobic interactions with the column matrix. To test this hypothesis, we examined its elution behaviour as a function of mobile phase organic modifier content.

As shown in Fig. 1b, the BrC chromatogram exhibited a “smeared” light absorption at 25% ACN, which persisted >15 min after the total permeation volume marker. Similar column retention was observed by Wong *et al.*,⁴² who subjected fresh BrC from laboratory pyrolysis of wood to SEC analysis using a 25 mM ammonium acetate buffer with 10% MeOH as mobile phase organic modifier. In our study, this smearing was significantly reduced at 50% ACN (Fig. 1d), which suggests that it arose from hydrophobic adsorption of BrC components at the surface of the column matrix. As illustrated in Fig. 1a and c, SRHA did not exhibit this smeared absorption, and was not influenced by these changes in mobile phase composition, which clearly highlights its limitations as a proxy for fresh biomass burning BrC.

Further comparison of the elution behaviour of our BrC sample to that of SRHA reveals that these samples also differ in their molecular size, compositional complexity, and light absorption profiles. At 50% ACN, for example, SRHA eluted as a single component, with featureless light absorption that decreased sharply with increasing wavelength (Fig. 1c). By contrast, under these conditions, our BrC sample eluted as two distinct fractions, both of which eluted later than SRHA: a high-



Fig. 1 Absorption density plots of SRHA and fresh biomass burning BrC (sampled *via* PILS, as described in the main text) as a function of mobile phase ACN content: (a) SRHA, 25% ACN; (b) BrC, 25% ACN; (c) SRHA, 50% ACN; (d) BrC, 50% ACN. In all cases, the remainder of the mobile phase was 20 mM phosphate buffer (pH 6.8).



MW fraction, the absorption profile of which resembled SRHA, and a low-MW fraction, which exhibited a structured absorption profile in the UV region (Fig. 1d). Since, as discussed above, this mobile phase composition effectively mitigated hydrophobic interactions between BrC and the column matrix, these observations suggest that the MW of our BrC sample is smaller than that of SRHA, and again underscores that SRHA is an unsuitable calibrant for fresh BrC.

To measure the size-resolved optical properties of BrC in the absence of a stationary phase and thereby verify that our conclusions were not specific to SEC analysis, we also analyzed BrC and Suwannee River natural organic matter (SRNOM) standard using a complementary technique, AF4. As shown in Fig. 2, although performed using a different mobile phase and with SRNOM rather than SRHA as comparative standard, AF4 shows similar results to those obtained using SEC. In particular, the measured molecular size of BrC (0.81 ± 0.02 kDa) determined *via* AF4 analysis was significantly smaller than that of SRNOM (1.20 ± 0.03 kDa). In addition, whereas the SRNOM eluted as a single peak with a relatively broad and featureless fluorescence emission maximum near 440 nm, the BrC sample again eluted as two distinct fractions. The first fraction, which eluted over a narrow size range, had a comparably lower MW and a narrow fluorescence maximum near 305 nm. In terrestrial/marine environments, this emission profile is typically attributed to tyrosine-like⁶² and/or polyphenolic⁶³ fluorophores, but this may not be the case for atmospheric samples.⁶⁴ The second fraction, which eluted over a broad size range, had a comparably larger MW and a broad fluorescence emission profile centred at 390 nm, the

latter of which is consistent with emission by HULIS in ambient PM samples collected in Malaysia during periods of intense haze resulting from peatland fire emissions.⁶⁵ Although further investigation of BrC using AF4 is certainly warranted, these results broadly verify the intrinsic complexity of BrC, as revealed by our SEC results.

All of these results differ from those of Di Lorenzo and co-workers,^{39,40} who used HPLC-SEC-PDA to study light absorption by ambient BrC sampled from atmospherically aged biomass burning plumes (~ 10 h to >3 days). Specifically, their BrC samples displayed only minor evidence of hydrophobic interactions with the column matrix, even in the absence of organic mobile phase modifiers. In addition, the light absorption of their samples was similar to SRHA, in that humic-like absorption was observed over the entire chromatogram, without the structured absorption we observed for our fresh BrC; finally, their samples eluted at a similar time as SRHA.

These differences could reflect fundamental differences in sample types: whereas our BrC sample is solely from peat combustion, the BrC samples studied by Di Lorenzo *et al.* were collected from a variety of locations, and would therefore be expected to include contributions from multiple fuel types—and, in some cases, urban and industrial emissions.⁴⁰ These differences could also reflect changes in BrC properties during atmospheric transport: for example, previous aircraft⁶⁶ and laboratory⁶⁷ studies of water uptake by biomass burning PM have shown an increase in particle hygroscopicity with increasing particle oxidation. Additional evidence for the role of atmospheric aging is provided in the Di Lorenzo *et al.* study itself, which showed that



Fig. 2 Asymmetric flow field-flow fractionation (AF4; UV-Vis and fluorescence detection) characterization of (a) fresh peat BrC and (b) Suwannee River natural organic matter (SRNOM) standard. For each sample, the leftmost plot shows the fluorescence emission spectrum (excitation at 230 nm) as a function of AF4 elution time, and the rightmost plot shows the corresponding single-wavelength absorption chromatogram (254 nm).



the contribution of high-MW chromophores to overall BrC absorption increased with plume age; therein, the authors attributed this change to the in-plume (photo)oxidation and/or oligomerization of low-MW species.⁴⁰ Since our BrC sample was collected immediately after its release from the combustion source, we would expect it to be less oxidized, more hydrophobic, and less depleted in low-MW chromophores than even the “freshest” BrC sample in the aforementioned work, all of which is consistent with our observations. To fully understand these differences, however, investigation of the aging behaviour of our specific BrC sample would be required.

3.2.2 The apparent size of fresh BrC changes with mobile phase composition. As shown in Fig. 1, the elution profile of our fresh BrC sample is highly sensitive to changes in mobile phase composition; in addition, at 50% ACN, our sample exhibits two distinct size fractions, each with different light-absorbing properties. This compositional complexity is consistent with recent work by Spranger *et al.*,³⁷ who used SEC coupled with reversed-phase HPLC to show that aqueous extracts of ambient PM collected from biomass burning-influenced regions vary in both molecular size and polarity.

To further explore the compositional diversity of our BrC sample, we examined its elution behaviour as a function of mobile phase solvent strength. In these experiments, we employed a 50 : 50 mixture of 20 mM phosphate buffer and an organic modifier blend consisting of varying ratios of MeOH and ACN; the resulting BrC elution profiles are shown in Fig. 3a. To quantify the relative contribution of the high-MW and low-MW fractions described above to the total BrC absorption for each mobile phase composition, we integrated the absorbance values obtained for each fraction (see the ESI† for details); these results are shown in Fig. 3b.

As the composition of the organic blend shifted from MeOH to ACN, the elution profile of BrC changed substantially. At 40% MeOH/10% ACN, absorption by the low-MW fraction was barely visible, and ~75% of the total absorption was contributed by the high-MW fraction. As the ACN content of the organic blend increased, however, the absorption of the low-MW fraction became more apparent, especially around 250–300 nm, and its contribution to the integrated absorbance increased significantly: at 0% MeOH/50% ACN, for example, the low-MW fraction contributed nearly half of the total absorption. We note that the results obtained at 50% MeOH/0% ACN diverged from this trend: under these conditions, the BrC sample exhibited the same delayed elution and “smeared” absorption profile observed at 25% ACN (see Section 3.2.1); as a result, the absorption contributed by the low-MW fraction was biased high. These results, which are unsurprising given the relative elution strengths of these solvents in reversed-phase partition chromatography (*i.e.*, ACN > MeOH), suggest that MeOH is less effective than ACN at disrupting hydrophobic interactions between the sample and the column matrix.

In principle, the increased contribution of the low-MW fraction to the total BrC absorption at higher ACN contents could reflect a change in column–analyte interactions. However, if these interactions were dominant here, we would have expected to see an increase in the high-MW fraction at elevated ACN/MeOH ratios as a result of suppression of hydrophobic adsorption, which is the opposite of what we observed. Instead, we attribute our observations to the disruption of weak intermolecular forces present in the large-MW BrC fraction, presumably resulting from changes in the relative strengths of solute–solute and solute–solvent interactions in mobile phases with different amounts of ACN and MeOH. Specifically, in the case of MeOH, which is both a hydrogen bond donor and



Fig. 3 BrC elution behaviour as a function of mobile phase solvent strength: (a) absorption density plots and (b) absorption contributed by high-MW and low-MW BrC fractions, with the latter calculated as described in the ESI† Experiments were conducted under the following mobile phase organic modifier conditions: (i) 50% MeOH/0% ACN, (ii) 40% MeOH/10% ACN, (iii) 25% MeOH/25% ACN, (iv) 0% MeOH/50% ACN. In all cases, the remainder of the mobile phase was 20 mM phosphate buffer (pH 6.8).



acceptor, we suggest that the attractive forces between the BrC and the solvent are not strong enough to overcome the solvent-solvent hydrogen bonding network and the intermolecular forces between the BrC components. By contrast, because ACN is only a hydrogen bond acceptor, it weakens the solvent-solvent hydrogen bonding network; in addition, it can interact with π -electrons, thereby disrupting π - π interactions between BrC components.⁶⁸

Our observations are consistent with two possible roles for ACN: first, promoting the liberation of adsorbed smaller molecules from large molecules and/or stable aggregates; second, mediating the partial dissociation of labile aggregates. To investigate the importance of the second pathway, we examined our sample's elution behavior at even higher ACN contents (>50%), which we anticipated would further disrupt the intermolecular forces between BrC components. As shown in Fig. 4, at higher ACN contents, the BrC elution profile became narrower: specifically, as the ACN content increased and the ionic strength of the mobile phase correspondingly decreased, the high-MW fraction tended to elute later and the low-MW fraction tended to elute earlier. We attribute the early elution of the low-

MW fraction at elevated ACN contents to further suppression of hydrophobic interactions between the sample and the column matrix. If the elution behaviour of the high-MW fraction were dominated by hydrophobic interactions (or, alternatively, by either of ionic exclusion or intramolecular electrostatic repulsions), the high-MW fraction should also elute earlier at higher ACN contents; however, we observed the opposite trend. At 70% ACN, all BrC components eluted together with acetone at the total permeation volume (*i.e.*, the lower-MW limit of the column separation), with the structured absorption still remaining as an overlay on the broad, featureless absorption. These observations suggest that the high-MW fraction of the fresh BrC sample consists of aggregates susceptible to disruption by ACN; in addition, the preservation of this fraction's broad, featureless absorption extending into the visible region indicates that this disaggregation process does not affect the sample's chromophoric properties.

To further verify this conclusion, we subjected SRHA to the same mobile phase protocol (Fig. S2†). Previous studies³⁶ have shown that humic materials consist of supramolecular associations of relatively small fractions connected by nonbonding



Fig. 4 BrC and SRHA elution behaviour as a function of mobile phase solvent strength: (a–d) BrC absorption density plots and (e–h) single-wavelength (300 nm) chromatograms (— SRHA, — acetone, — BrC). Experiments were conducted under the following mobile phase organic modifier conditions: 25% ACN (a and e), 50% ACN (b and f), 60% ACN (c and g), and 70% ACN (d and h). In all cases, the remainder of the mobile phase was 20 mM phosphate buffer (pH 6.8).



intermolecular forces. Similar to the high-MW BrC fraction, SRHA eluted later and with a broader size distribution at higher ACN contents (Fig. 4e–g), and co-eluted with acetone at 70% ACN (Fig. 4h), indicating dissociation of SRHA aggregates into smaller subunits. These observations support our characterization of the high-MW component of BrC as small BrC components held together *via* hydrophobic interactions.

3.2.3 The apparent size of fresh BrC changes with mobile phase ionic strength. Atmospheric PM spans a wide range of ionic strengths.⁶⁹ In this context, we examined the elution behaviour of our BrC sample as a function of ionic strength of the phosphate buffer component of the SEC mobile phase. In these experiments, we used a 50 : 50 mixture of buffer and an organic modifier (25% ACN/25% MeOH), the latter of which we chose to minimize the influence of hydrophobic interactions between our samples and the column matrix, and varied the concentration of the buffer from 10 to 40 mM.

As shown in Fig. 5, the high-MW BrC fraction eluted later at 20 mM *versus* 10 mM; the same delay was also observed for SRHA (Fig. S3†). In principle, ionic strength may have influenced the elution in two ways here: first, *via* suppressing electrostatic interactions between the column matrix and polar and charged functional groups (*e.g.*, hydroxyl, carbonyl, amide)⁷ present in the BrC sample (*i.e.*, ionic exclusion); second, *via*

reducing intramolecular repulsions between these same charged functional groups, resulting in a more “compacted” molecule. Because both of these effects would delay elution, we cannot definitively distinguish between them. However, evidence supporting the existence of the second effect is provided by previous studies of humic substance conformation, which reported appreciable decreases in the size of humic aggregates at elevated ionic strengths.^{44,47} We also observed an enhanced absorption for the low-MW fraction at higher ionic strengths: in particular, as shown in Fig. 5, the absorption at ~300 nm was higher in the presence of 20 mM buffer than in the presence of 10 mM buffer. As well, at 40 mM, the high-MW and low-MW fractions were no longer resolved. These results could be explained by a decrease in the hydrophobic sorption capacities of BrC aggregates at higher ionic strengths,⁷⁰ leading to liberation of some absorbed low-MW BrC subunits.

4 Conclusions and implications for analysis of complex atmospheric samples

This study represents the first systematic investigation of the challenges and benefits associated with the application of HPLC-SEC-PDA to the characterization of atmospheric brown carbon (BrC), an important class of light-absorbing PM and the dominant contributor to the absorption profile of wildfire PM. In the following paragraphs, we highlight the implications of our results for SEC analysis of atmospheric samples and provide recommendations for future work in this area.

A major conclusion of this study is that careful consideration of solvent–analyte interactions is crucial when identifying appropriate extraction and chromatographic/spectroscopic analysis conditions for natural samples. Previous work has shown that the composition of PM extracts can be influenced by reactions of specific organic functional groups with sample extraction solvents (*e.g.*, MeOH).⁷¹ Here, we show that organic solvents can influence BrC properties even without reacting directly with BrC components: in particular, we show that ACN not only mitigates hydrophobic interactions between BrC and the SEC column matrix but also disrupts metastable BrC aggregates, thereby leading to changes in measured BrC size distributions. This aggregation and dissociation behaviour could potentially contribute to discrepancies between MW values inferred from SEC and those determined using mass spectrometry for the same sample.^{38,39,46,72}

Importantly, and more broadly, our findings imply a need for caution when using chromatographic elution behaviour (*e.g.*, in reversed-phase HPLC) to make inferences regarding the hydrophobicity and other properties of BrC, as the relationship between retention time and mobile phase identity for a given BrC component may reflect both changes in its distribution constant and changes in its intrinsic properties. For example, increases in ACN mobile phase content (*e.g.*, in gradient methods) could potentially lead not only to changes in partitioning behaviour but also to dissociation of BrC aggregates, with concomitant indirect changes in retention behaviour.

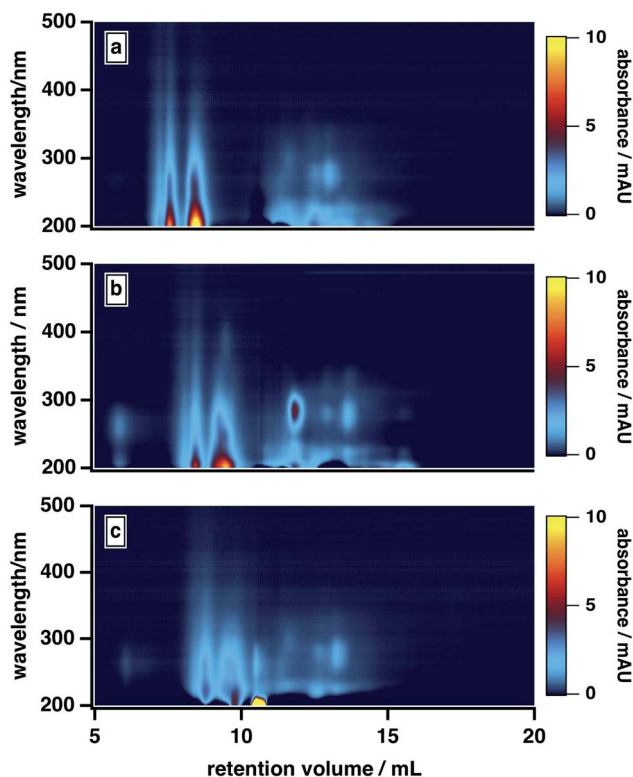


Fig. 5 Absorption density plots for fresh peat BrC as a function of mobile phase phosphate buffer concentration (*i.e.*, ionic strength). Mobile phases were prepared using three concentrations of phosphate buffer (pH 6.8): (a) 10 mM, (b) 20 mM, (c) 40 mM. In all cases, the mobile phase composition was 50% phosphate buffer, 25% MeOH, and 25% ACN.



Successful application of SEC to MW estimations of complex mixtures is a challenging task, as it requires both selection of calibration standards with comparable chemical and structural properties to the analytes of interest and careful consideration of biases resulting from differences in the magnitudes of secondary effects experienced by mixture components compared to the chosen calibrant suite. For example, previous studies applying SEC to analysis of atmospheric PM have employed both commercial macromolecules and atmospherically relevant small molecules (*e.g.*, phenols and aromatic acids) for MW calibration.^{38,43,73,74} However, given that we would expect these calibrants to be differently susceptible to secondary effects than BrC, this strategy has the potential to lead to dramatic changes in estimated MWs with changing solvent composition. In the case of humic substances, these issues have occupied analytical chemists for decades.⁷⁵ In the case of fresh BrC, which we show here is made up of components with a wide range of sizes, polarities, and susceptibility to conformational changes—and therefore subject to different secondary effects—achieving size-based separation of all BrC components at any single mobile phase composition, no matter how optimized, is likely an impossible task. This challenge has the potential to be especially important in cases where SEC is coupled with complementary techniques to investigate the chemical properties of specific analyte size fractions,^{37,73,74} as a given elution volume could potentially contain contributions not only from “ideal” analytes (*i.e.*, those for which secondary interactions have been minimized) but also from many other analyte classes (*e.g.*, higher-MW but more hydrophobic, smaller-MW but susceptible to intramolecular electrostatic repulsions).

In this context, we suggest a more expansive approach to the SEC characterization of complex atmospheric mixtures, in which its apparent limitations are instead reconceptualized as advantages. For example, in this study, by subjecting our BrC sample to different mobile phase conditions, we were able to explore not only its molecular size but also its hydrophobicity, conformation, and intermolecular associations.³⁶ This approach, which is analogous in some respects to the use of chromatographic retention behaviour to estimate the properties of single analytes (*e.g.*, vapour pressure⁷⁶ and K_{OW} ⁷⁷), could also be applied in other ways. For example, if coupled with refractive index detection, this strategy would enable the characterization of non-chromophoric PM components. In addition, the fluorescence detection approach described here could be expanded through the use of excitation–emission matrix analysis,⁶⁴ which has the potential to aid in the chemical characterization of the organic fluorophores present in BrC fractions. More broadly, the investigation of mobile phase-dependent sample behaviour has the potential to facilitate the prediction of analyte properties under the elevated organic content and ionic strength conditions characteristic of atmospheric PM, which are challenging and/or impossible to explore directly in the laboratory.

Author contributions

M. L., D. K. T., and S. A. S. designed the combustion experiments with assistance from C. J. Y., M. L., N. Z., S. A. S., and D.

K. T. collected the boreal peat samples and performed the combustion experiments. M. L. performed all HPLC-SEC-PDA analyses; C. W. C. conducted the AF4 analysis. M. L. and S. A. S. interpreted experimental data, with assistance from C. J. Y., and wrote the manuscript, with critical comments from D. K. T., C. W. C., and C. J. Y.

Conflicts of interest

There are no conflicts of interest to declare.

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