¹ Fluorescence characteristics, absorption properties, and

radiative effects of water-soluble organic carbon in seasonal snow across northeastern China

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10 Abstract. Water-soluble organic carbon (WSOC) in the cryosphere can significantly influence the global 11 carbon cycle and radiation budget. However, WSOC in the snowpack has received little scientific 12 attention to date. This study reports the fluorescence characteristics, absorption properties, and radiative effects of WSOC based on 34 snow samples collected from sites in northeastern China. A significant 13 degree of regional WSOC variability is found, with concentrations ranging from $0.5 \pm 0.2 \ \mu g \ g^{-1}$ to 5.7 14 \pm 3.7 µg g⁻¹ (average concentration: 3.6 \pm 3.2 µg g⁻¹). The three principal fluorescent components of 15 16 WSOC are identified as: (1) the high-oxygenated humic-like substances (HULIS-1) of terrestrial origin, 17 (2) the low-oxygenated humic-like substances (HULIS-2) of mixed origin, and (3) the protein-like 18 substances (PRLIS) derived from autochthonous microbial activity. In southeastern Inner Mongolia 19 (SEIM), a region dominated by desert and exposed soils, the WSOC exhibits the highest humification 20 index (HIX) but the lowest fluorescence (FI) and biological (BIX) indices; the fluorescence signal is 21 mainly attributed to HULIS-1, and thus implicates soil as the primary source. By contrast, the HIX (FI 22 and BIX) value is the lowest (highest) and percentage of PRLIS is the highest in the remote area of the 23 northeastern Inner Mongolia (NEIM), suggesting a primarily biological source. For south and north of 24 northeastern China (SNC and NNC), both of which are characterized by intensive agriculture and 25 industrial activity, the fluorescence signal is dominated by HULIS-2, and the HIX, FI, and BIX values

are all moderate, indicating the mixed origins for WSOC (anthropogenic activity, microbial activity, and
soil). We also observe that, throughout northeastern China, the light absorption of WSOC is dominated
by HULIS-1, followed by HULIS-2 and PRLIS. The contribution of WSOC to albedo reduction (average
concentration: 3.6 µg g⁻¹) in the ultraviolet–visible (UV–vis) band is approximately half that of black
carbon (BC average concentration: 0.6 µg g⁻¹). Radiative forcing is 3.8 (0.8) W m⁻² in old (fresh) snow,
equating to 19 % (17 %) of the radiative forcing of BC. These results indicate that WSOC has a profound
impact on snow albedo and the solar radiation balance.

8 1 Introduction

9 Seasonal snow plays a significant role in Earth's solar radiation energy budget owing to its high 10 reflectivity (Beniston et al., 2017; Usha et al., 2020; Xie et al., 2018). In recent decades, however, the 11 extent of snow-covered areas has trended downward, partially as a result of the presence of lightabsorbing particles (LAPs) in the snowpack (Barnett et al., 2008; Dumont et al., 2014; Groisman et al., 12 13 1994). The LAPs in seasonal snow, such as black carbon (BC), organic carbon (OC), mineral dust (MD), 14 and biota (Beres et al., 2020; Di Mauro, 2020; Els et al., 2020; Qian et al., 2015; Wu et al., 2016), can 15 strongly absorb solar radiation, which serve to lower surface albedo and impose a positive radiative 16 forcing (Cui et al., 2021; Dumont et al., 2014; Hansen and Nazarenko, 2004; Shi et al., 2022b; Warren 17 and Wiscombe, 1980; Zhang et al., 2017). Ultimately, LAPs can accelerate snow melting (Li et al., 2021b) 18 and disturb the global radiative balance, therefore, have important implications for regional and global 19 climate change (Shi et al., 2022a; Skiles et al., 2018). 20 Snowpack BC and MD have been the focus of considerable research in snow-covered regions worldwide 21 (Antony et al., 2014; Doherty et al., 2014; Hegg et al., 2010; Li et al., 2021a; Wang et al., 2014b; Zhang

et al., 2018). As the most important LAP (Bond et al., 2013; Doherty et al., 2010; Shi et al., 2020; Wang

1	et al., 2014b), the radiative efficiency of snowpack BC can be more than three times greater than that of
2	carbon dioxide (Flanner et al., 2007). MD, another important snowpack LAP, is also known to alter the
3	cryospheric environment owing to its light-absorbing properties (Di Mauro et al., 2015; Painter et al.,
4	2007; Sarangi et al., 2020; Shi et al., 2021). Recently, researchers have also begun evaluating the
5	influence of biomes on global snow albedo (Hotaling et al., 2021). In contrast, however, the role of OC
6	remains poorly understood because of its complex composition and a lack of OC-focused research.
7	Consequently, substantial uncertainty surrounds the origins, optical properties, and radiative effects of
8	snowpack OC.
9	A recent study has reported that the storage of OC in mountain glaciers and ice caps (~11 % of Earth's
10	land surface) could be as high as 6 petagrams (Pg, Hood et al., 2015), the majority of which is water-
11	soluble organic carbon (WSOC) (Yan et al., 2016). The substantial part of WSOC in the glacier is highly
12	bioavailable and can be a source of labile organic matter for downstream ecosystems (Singer et al., 2012).
13	The physical and photochemical processes can occur within various WSOC in snow cover and glaciers,
14	and therefore have a great effect on atmospheric and glacier chemistry (Antony et al., 2011; Domine,
15	2002; Grannas et al., 2007). Moreover, WSOC has important influences on the energy budget and
16	radiative forcing of snow cover and glaciers (Kirillova et al., 2014; Ram et al., 2010; Yan et al., 2016).
17	As the chief absorber of WSOC, water-soluble brown carbon (WS-BrC) can absorb significant measures
18	of solar radiation in the UV-vis wavelengths (Murphy et al., 2008). For instance, in their analysis of 21
19	snow samples from the Arctic and Antarctic, Anastasio and Robles (2007) attributed 50 % of the total
20	light absorption coefficients at wavelengths of > 280 nm to organic chromophores of WSOC. Beine et
21	al. (2011) reported that WSOC occupies almost the entire absorption spectrum of the photochemically
22	active region (300-450 nm) in surficial snow samples from Barrow, Alaska. Feng et al. (2016) observed

1	that absorption in cryoconite samples from the central Tibetan Plateau was dominated by WSOC
2	components in the 300-350 nm range. Similarly, Yan et al. (2016) measured WSOC in glacial snow
3	from Laohugou glacier, northern Tibetan Plateau, where they found that the radiative forcing was ~ 10 %
4	that of BC. Together, these studies indicate that WSOC plays a key role in global snowpack energy
5	absorption (Niu et al., 2018; Zhang et al., 2020). We note that recent research on cryospheric WSOC
6	mainly focused on alpine glaciers and polar regions (Guo et al., 2022; Li et al., 2022), while the extensive
7	mid-latitude regions impacted by seasonal snowpack remain relatively understudied.
8	The composition of WSOC is typically complex, and characteristics of fluorescence and absorption can
9	vary widely among the different components. Nonetheless, recent studies have tended to treat WSOC as
10	a single entity and focus on the overall impacts (Barrett and Sheesley, 2017; D'Sa et al., 2014; Niu et al.,
11	2018; Wu et al., 2019), such that the specific roles of individual components are poorly constrained. One
12	commonly used analytical method for distinguishing the components and properties of fluorescence is
13	the fluorescence excitation-emission matrix (EEM), which has the advantage of high sensitivity and
14	small sample size (Coble, 1996; Kowalczuk et al., 2005). First applied in oceanic contexts (Coble et al.,
15	1990), EEM has been gradually extended to lakes, fog water, and rainwater (Birdwell and Valsaraj, 2010;
16	Huguet et al., 2009; McKnight et al., 2001). At present, the application of EEM on atmospheric aerosols
17	has entered a mature stage. EEM has been used to identify fluorescent WSOC components in aerosols
18	from polar regions or urban backgrounds, and it has been found that different structures of WSOC
19	fractions exhibit different oxidation properties, which may provide a clue to understand the chemical
20	formation or loss of organic chromophores in atmospheric aerosols (Chen et al., 2016; Fu et al., 2015).
21	Recently, this method has been gradually extended to the analysis of glacier samples and snow samples
22	(Feng et al., 2018; Guo et al., 2022; Zhou et al., 2019b). Concurrently, parallel factor analysis

1 (PARAFAC) is an effective approach to extracting the individual fluorescence components and their 2 corresponding fluorescence information from complex EEMs, thus making EEM-PARAFAC a direct 3 and viable means for exploring sources of WSOC. For example, Zhou et al., (2019b) used EEM-4 PARAFAC to identify the multiple sources of WSOC measured in seasonal snow in northwestern China. 5 Accordingly, we have applied EEM–PARAFAC for our analysis of snow samples in this study. 6 EEM-PARAFAC can only provide plausible information about the component-specific influence of 7 WSOC on fluorescent properties, and the quantitative fractional contributions of specific components to 8 light absorption are still unknown. Recently, Chen et al. (2019a) collected atmospheric aerosol samples 9 in PM_{2.5} over Xi'an, China, and successfully attributed the dithiothreitol (DTT) activity levels to various 10 BrC components by coupling DDT and BrC datasets. A similar attribution method has been applied to 11 various research areas, including climate change, extreme weather, and atmospheric environments (Cao 12 et al., 2015; Pokrovsky, 2019; Xin et al., 2016; Zhao et al., 2019). In this study, we applied a multiple 13 linear regression method comparable to that of Chen et al. (2019a) to derive the fractional contribution 14 of each WSOC component to light absorption. Despite this method having been used elsewhere (Wu et 15 al., 2022; Wu et al., 2021), it remains a highly innovative approach to evaluating the light absorption of 16 snowpack WSOC.

Northeastern China supports an extensive snowpack during winter and spring. As a major industrial and agricultural center, this region is also the principal source of heavy airborne pollutants that are incorporated into seasonal snow via wet and dry deposition (Wang et al., 2017). Coupled with intensive tilling of farmland, the geographical proximity of northeastern China to the neighboring desert regions also provides a source of soil organic matter that becomes entrained into the snowpack (Wang et al., 2013b). Compared with research on BC-snow mixing ratios and their radiative impact in northeastern

1	China (Dang et al., 2017; Huang et al., 2011; Pu et al., 2019), studies of WSOC are still in their infancy.
2	To address this deficiency, we made the first investigation of the fluorescence characteristics, absorption
3	properties, and radiative effects of WSOC in seasonal snow samples in northeastern China. Specifically,
4	we applied EEM-PARAFAC to identify the origins and fluorescence characteristics of snowpack WSOC,
5	after which we derived individual absorption contributions for each WSOC component using
6	fluorescence data, an absorption data series, and an attribution method. Finally, we estimated the
7	reduction of snow albedo and radiative forcing caused by WSOC and BC via the Spectral Albedo Model
8	for Dirty Snow (SAMDS) radiative transfer model.
9	2 Methods
10	2.1 Sample collection
11	During January and December 2020 and January 2021, we collected 34 snow samples from sites across
12	northeastern China, including the eastern part of Inner Mongolia and Heilongjiang and Jilin provinces.
13	Sample numbers were set following previous campaigns (Pu et al., 2017; Wang et al., 2013b, 2017), with
14	the exception that samples from the Changbai Mountain area are numbered individually. The
15	geographical distribution of sampling sites and respective land-cover types are shown in Figure 1a. The
16	sites are characterized by five land-cover types, including forest, grassland, desert, cropland, and frozen
17	lake/river (Fig. 1b-g). On the basis of these geographical and environmental classifications, we divided
18	the sampling sites into five broad regions: southeastern Inner Mongolia (SEIM; Q494-495, Q497-499),
19	the south of northeast China (SNC; Q470-471, Q473-474, Q477, Q484, Q486-Q489, Q491-Q493,
20	Q501), the north of northeast China (NNC; Q480-483), the Changbai Mountain area (CBM; CM1-CM2,
21	CM5, CM11, CM13-CM14), and northeastern Inner Mongolia (NEIM; Q440, Q443, Q447, Q449,
22	Q454).





Figure1: (a) Information on sampling site distributions in northeastern China, including the land cover type,
site number, and grouping. Land cover types are derived from Collection 5.1 of the MODIS global land cover
type dataset (MCD12Q1: https://lpdaac.usgs.gov/products/mcd12q1v006/) and are indicated by specific
colors and symbols relative to sampling sites. Sampling sites are divided into five groups defined by black
rectangles. (b–g) Photographs depicting the typical snow and ground-cover conditions of our various sampling
sites.

8 The sampling sites were chosen at random but had to be located ≥ 20 km from cities and villages and at 9 least 1 km upwind of roads or railroads to minimize the influence of single-point pollution sources and 10 to ensure the broadest regional representation. Furthermore, we performed sample collection oriented 11 toward the wind to avoid contamination from personnel. At each site, we used a sterile disposable shovel 12 to collect 0-5 cm-thick samples of surface snow, which were subsequently stored in sterile Whirlpak 13 (Nasco, WI, USA) bags. For snow depths of < 5 cm, we determined the sampling depth according to the 14 actual conditions to avoid inducting significant soil impurities during sampling. Snow samples were 15 melted at room temperature (25 °C) and stored in a freezer at -20 °C until analysis in the laboratory. For 16 more operational details, we refer the reader to Wang et al. (2013b).

1 2.2 Chemical species analysis

2 All collected snow samples were stored in a freezer at -20 °C until analysis in the laboratory. In the lab 3 process, the samples were first melted at room temperature (25 °C). Then 30 mL meltwater was taken 4 for each sample with the clean disposable syringe (Jiangnan, Anhui, China) and injected into the pre-5 baked (4h, 450 °C) glass bottle passing through 0.45 µm pore-sized polytetrafluoroethylene filters 6 (Jinteng, Tianjin, China). Finally, the concentration of WSOC was measured by the total organic carbon 7 analyzer (Aurora 1030W, OI Analytical, TX, USA), and measurement detection limits and relative 8 standard deviations are 2 μ g L⁻¹ and 1 %, respectively. The concentration of WSOC for ultrapure water 9 blank is 0.35 mg L⁻¹, and the value of each sample after blank subtraction is presented in Table S1. 10 We used 0.4 µm pore-sized polycarbonate filter membranes (Whatman, USA) to isolate BC and other 11 insoluble particles, following the protocols outlined by Wang et al. (2020), after which we employed a 12 custom-developed two-sphere integrating-sandwich (TSI) filter-based spectrophotometer to measure 13 particle absorption. Coupled with the mass of filtered meltwater, these optical measurements were then 14 converted to snowpack BC concentrations. To make these calculations, we applied a BC mass-absorption coefficient (MAC) and absorption Ångström exponent (AAE) of 6.3 m² g⁻¹ (550 nm) and 1.1, 15 16 respectively, after Pu et al. (2017). We note that TSI provides greater accuracy and smaller overall 17 uncertainties in the quantification of seasonal snow BC than do thermo-optical carbon analysis (Wang et 18 al., 2020), and thus it has been applied widely in this type of research (Shi et al., 2020). For more detailed 19 information, see Wang et al. (2013b).

20 2.3 Fluorescence and absorption measurement

We obtained absorbance and fluorescence EEMs for filtered meltwater samples via synchronous
 absorption-3D fluorescence scanning spectrometry (Aqualog, Horiba Scientific) with the following

1	measurement parameters: fluorescence spectra excitation range = 240-800 nm in 3 nm intervals,
2	emission range = 152.25–929.92 nm in 5.04 nm (8 pixels) intervals, scanning interval = 0.3 seconds.
3	Prior to sample measurement, the aliquots of filtered ultra-pure water were analyzed (18.2 M ω cm, Milli-
4	q Purification System, Millipore) as analytical blanks. The fluorescence intensity was normalized to that
5	of the water Raman unit (RU), which exhibited a peak excitation wavelength of 350 nm, and deducted
6	this Raman signal from all subsequent sample tests (Lawaetz and Stedmon, 2009). The inner filtration
7	effect correction was based on the absorbance-based approach (Kothawala et al., 2013), using the
8	measured absorbance at each pair of excitation and emission wavelengths across the EEMs to convert
9	the observed fluorescence intensity into the corrected fluorescence intensity. Rayleigh scattering peaks
10	were processed by interpolation algorithm in the EEMscat MATLAB toolbox (Bahram et al. 2006). As
11	fluorescence signals with wavelengths greater than 600 nm are primarily noise (Zhou et al., 2019b), they
12	are not considered further in this study.
13	We used version 0.6.3 of the MATLAB drEEM toolbox (http://dreem.openfluor.org/; Murphy et al.,
14	2013) to perform PARAFAC analysis on EEMs. Comprising the consistency index, residuals, and visual
15	inspections, the 3-component model was considered more reliable and representative than the 2-7-
16	component models (Fig. S1) and passes the S4C6T3 split scheme (Fig. S2; Murphy et al., 2013). The
17	contributions of these three components to the overall fluorescence signal were expressed as relative
18	percentages of F_{max} in RU, and the total fluorescence volume (TFV, RU nm ²) was calculated from the
19	EEMs (Song et al., 2019). Normalized TFV equates to NFV (RU nm ² (mg $L^{-1})^{-1}$), using NFV =
20	TFV·c(WSOC) ⁻¹ , where c(WSOC) is the concentration of WSOC in the snow (mg L^{-1})), and represents
21	the fluorescence ability of WSOC (Chen et al., 2019a).

1	We calculated three fluorescence-derived indices-the fluorescence index (FI), biological index (E	3IX),
2	and humification index (HIX)-from the ratio of fluorescence intensity at specific excitation	and
3	emission wavelengths. As demonstrated by previous studies (Birdwell and Valsaraj, 2010; Huguet e	t al.,
4	2009; McKnight et al., 2001), these ratios can help characterize potential sources of WSOC. Specific	ally,
5	the FI is taken to represent the relative amount of DOM derived from terrestrial and microbial/a	ılgae
6	sources (McKnight et al., 2001); high values correspond to terrestrially derived organics, and low va	alues
7	reflect microbial sources. The HIX describes the degree of humification of soluble organic m	atter
8	(Zsolnay et al., 1999). During humification, the aromaticity of organic matter increases as micro	obial
9	availability decreases, such that higher HIX values correspond to more strongly humified and/or hi	gher
10	aromaticity organics (principally of terrestrial origin), whereas lower values indicate autochthonor	ıs or
11	microbial origins. As a measure of autochthonous productivity (Huguet et al., 2009), elevated BIX va	alues
12	are associated with increased contributions of microbial-derived fluorescent organic matter. The	three
13	indices are calculated by the following equations (Feng et al., 2016; Huguet et al., 2009; McKnight et al., 2009;	rt al.,
14	2001; Ohno, 2002):	
15	$FI = \frac{I(Ex = 370, Em = 470)}{I(Ex = 370, Em = 520)},$	(1)
16	$BIX = \frac{I(Ex = 310, Em = 380)}{I(Ex = 310, Em = 430)},$	(2)

17
$$HIX = \frac{I(Ex = 254, Em = 435 - 480)}{I(Ex = 254, Em = 300 - 345) + I(Ex = 254, Em = 435 - 480)},$$
(3)

18	where I is the fluorescence intensity, and Ex and Em represent the excitation and emission wavelengths,
19	respectively. To ensure a direct comparison with prior results, we recalculated published HIX data using
20	the same calculation methods as in our analyses. According to a previous study, FI values of ≤ 1.4
21	correspond to terrestrial sources, and values of \geq 1.9 denote a primarily microbial origin. The values of
22	1.4–1.9 suggest a mixed origin (McKnight et al., 2001).

21	2.4 Snow albedo modeling and radiative forcing calculations
20	concentration of WSOC.
19	where a_{WSOC} is the absorption coefficient derived from Equation (4) and c(WSOC) (mg L ⁻¹) is the
18	$MAC_{\lambda} = a_{WSOC}(\lambda) / c(WSOC), \qquad (6)$
17	et al., 2019b; Yan et al., 2016):
16	We calculated the mass absorption coefficient (MAC $_{\lambda}$, m ² g ⁻¹) of our samples using the equation (Chen
15	where <i>K</i> is a constant related to WSOC concentration.
14	$\mathbf{a}_{\mathrm{WSOC}}(\lambda) = K \cdot \lambda^{-\mathrm{AAE}},\tag{5}$
13	Yan et al., 2016):
12	exponent (AAE) via the following equation (Doherty et al., 2010; Niu et al., 2018; Wang et al., 2013b;
11	To investigate the wavelength dependence of WSOC absorption, we obtained the Absorption Ångström
10	et al., 2010).
9	$(a_{WSOC}(280))$ to characterize the absorption intensity of WSOC for comparison with other studies (Zhang
8	Owing to the absorption characteristics of WSOC, we selected the absorption coefficient at 280 nm
7	the absorption coefficient (m^{-1}) .
6	where Abs is absorbance, λ is the wavelength, <i>L</i> is the path length of the cuvette (0.01 m), and a_{WSOC} is
5	$a_{WSOC}(\lambda) = \ln(10) \cdot Abs(\lambda) \cdot L^{-1}, \qquad (4)$
4	absorbance to an absorption coefficient using the following equation:
3	average absorbance above 600 nm from the whole spectrum (Chen et al., 2019b). We converted sample
2	and scattering effects of the measurement for the absorption spectra were corrected by subtracting the
1	The absorption spectra of WSOC were derived from 240 to 800 nm in 3 nm intervals. The baseline shifts

To reveal the radiative effect impact of snowpack WSOC in northeastern China, we used SAMDS to

1	simulate spectral snow albedo. This model is based on the asymptotic radiative transfer theory, which
2	has been verified in previous studies (Li et al., 2021b; Wang et al., 2017). As described in detail by Wang
3	et al. (2017), the model involves parameters including solar zenith angle, impurity concentrations, the
4	mass absorption coefficient of impurities, and snow grain radius. Measured values include the
5	concentration of BC and the absorption coefficients of WSOC. To quantify the influence of pollutants
6	on snow albedo, we assumed a semi-infinite snow layer and a uniform snow grain radius of 100 μ m for
7	fresh snow and 1000 µm for old snow, consistent with previous studies (Pu et al., 2021). With the solar
8	zenith angle fixed at 60°, consistent with our sampling dates and locations, we calculated the reduction
9	$(\Delta \alpha_i, i represents BC only, WSOC only, and BC + WSOC, similarly hereinafter.) in spectral snow albedo$
10	derived from different types of impurities for the UV-vis (280-400 nm) and ultraviolet-near infrared
11	(UV–NIR, 280–1500 nm) bands. The radiative forcing resulting from either BC or WSOC in snow (RF_i)
12	was then derived by multiplying the albedo reduction value by the incident solar radiation (Painter et al.,
13	2013):
14	$\mathrm{RF}_{i} = E \cdot (\alpha_{pure} - \alpha_{i}) = E \cdot \Delta \alpha_{i}, \tag{7}$
15	where α_{pure} is snow albedo for pure snow and E is the average daily downward shortwave solar radiation
16	flux acquired from NASA's Clouds and the Earth's Radiant Energy System (CERES) product "CERES

17 SYN1deg" (https://ceres.larc.nasa.gov/products.php?product=SYN1deg).

18 3 Results and discussion

19 **3.1** Characteristics of chemical species

20 Figure 2a shows the spatial distribution of measured WSOC in seasonal snow across northeastern China.

21 The averaged WSOC concentration (arithmetic mean \pm standard deviation) is 3.6 \pm 3.2 µg g⁻¹, with a

1	maximum of 18 μ g g ⁻¹ and a minimum of 0.3 μ g g ⁻¹ . Among all five regions, WSOC concentrations are
2	highest in SNC (average $5.7 \pm 3.7 \ \mu g \ g^{-1}$), likely reflecting the greater degree of agricultural and industrial
3	activity there compared with other regions (Lu et al., 2011; Wang et al., 2013b). We highlight that both
4	agricultural and industrial sources are considered anthropogenic. In contrast, the second highest
5	measured concentrations (3.4 \pm 1.5 μg g^-1) are from SEIM, where desertification occurs (Fang et al.,
6	2007) and that is considered a natural source of WSOC. For most sites, the underlying surface is desert
7	(Fig. 1a) that was incompletely covered by seasonal snow during the sampling period (Fig. 1d).
8	Consequently, the exposure of natural sandy soils is a potentially significant contributor to WSOC
9	through aeolian erosion and dry deposition. In NNC, where both the population density and agricultural
10	intensity are lower than in SNC (Choi et al., 2020), the contribution of anthropogenic pollution to the
11	snowpack is correspondingly lower, resulting in a moderate WSOC concentration of $2.7\pm0.8~\mu g~g^{-1}$
12	Meanwhile, being far from intensive human activity, both CBM and NEIM (Fig. 1a, b, and g) return the
13	low WSOC concentrations (CBM: $2.0 \pm 1.3 \ \mu g \ g^{-1}$; NEIM: $0.5 \pm 0.2 \ \mu g \ g^{-1}$). Nonetheless, the higher
14	value for CBM betrays the influence of far-traveled anthropogenic pollutants (Wang et al., 2015; Wu et
15	al., 2020; Zhang et al., 2013).



Figure 2: Spatial distributions of concentrations of (a) WSOC and (b) BC in snow samples. Sampling sites are
divided into the five groups defined in Figure 1. Bubble sizes are proportional to concentrations of WSOC

19 and BC.

1	In comparison with previous studies, we observe that, with the exception of NEIM, our measured WSOC
2	concentrations are significantly higher than those reported for snow/ice from the Tibetan Plateau (TGL,
3	~0.71–1.02 μ g g ⁻¹ , Feng et al., 2016), the Alps (~0.14–0.78 μ g g ⁻¹ , Vione et al., 2021), North America
4	(~0.1–0.3 μ g g ⁻¹ , Fellman et al., 2015), and polar regions (~0.12–0.27 μ g g ⁻¹ , Antony et al., 2014), but
5	comparable to values in Laohugou glacier ice from the Tibetan Plateau (~1.02–2.6 μ g g ⁻¹ , Feng et al.,
6	2016, 2018) and seasonal snowpack in northwestern China (0.48–7.07 μ g g ⁻¹ , Zhou et al., 2021). This
7	finding implies that snowpack WSOC in northeastern China is significantly contributing to regional and
8	global climate change (Domine, 2002).
9	A similar spatial pattern is exhibited by snowpack BC (Fig. 2b). For example, of all five regions, the
10	regional mean BC concentration is highest for SNC (923 \pm 512 ng g ⁻¹), followed by SEIM (659 \pm 582
11	ng g ⁻¹), NNC (494 \pm 224 ng g ⁻¹), and the CBM (391 \pm 312 ng g ⁻¹). BC concentrations are lowest in
12	NEIM (60 ± 19 ng g ⁻¹), in agreement with the values in remote areas reported by Doherty et al. (2010).
13	3.2 Fluorescence characteristics of WSOC
14	Three fluorescent components (C1-C3) were captured by resolving the EEMs. All fluorescence
15	information is summarized in Table S2. C1 exhibits a primary peak at Ex = 240 nm, Em = 448 nm,
16	indicating the high-oxygenated HULIS found primarily in aromatic conjugated macromolecules (Chen
17	et al., 2016). The weaker secondary peak occurs at longer excitation wavelengths (Ex / Em = 308 / 448
18	nm), implying a higher aromatic content and greater molecular weight (Coble et al., 1998). Wen et al.
19	(2021) concluded that C1 was probably derived from natural terrestrial sources, such as dust and soil, as
20	proposed originally by Stedmon et al. (2003) and Osburn et al. (2016). Accordingly, we classified C1 as
21	a terrestrial, humic-like substance, hereafter referred to as HULIS-1.

1	C2 exhibits a primary (secondary) peak at $Ex = 240$ (293) nm, $Em = 398$ nm, suggestive of the
2	lower-oxygenated HULIS (Chen et al., 2016). Observed in a variety of sources, Stedmon et al. (2003)
3	reported this component in terrestrial end-member samples. Whereas, both Murphy et al. (2011) and
4	Osburn et al. (2016) have since linked C2 to anthropogenic sources, such as urban runoff and sewage.
5	Microbial activity and the degradation of phytoplankton in natural aquatic systems are also thought to
6	contribute to this component (Yamashita et al., 2008; Zhang et al., 2009). Accordingly, we classified C2
7	as humic-like substances with complex origins in terrestrial, anthropogenic, and/or microbial sources,
8	hereafter termed HULIS-2. Unlike HULIS-1 and HULIS-2, C3 is recognizable as a UVB-like protein or
9	tyrosine-like fluorescence (hereafter PRLIS) with a primary (secondary) peak at $Ex = 240$ (293) nm, Em
10	= 398 nm (Osburn et al., 2016; Stedmon and Markager, 2005). PRLIS reflects autochthonously labile
11	DOM produced by biological processes (Stedmon et al., 2003) and has been reported in previous studies
12	of seasonal snow (Zhou et al., 2019b).



1

2 Figure 3: (a-c) Three fluorescent components identified by PARAFAC analysis. (d) Relative contributions of 3 the three components to total fluorescence at each site. HULIS-1, HULIS-2, and PRLIS are represented by 4 the specific colors shown in the legend (top left corner). The distributions of BC emissions and desert areas in 5 our study area are indicated by gray and light yellow, respectively, with darker gray colors indicating higher 6 black carbon concentrations. (e-i) Average contributions of the three components in different groups of 7 samples. BC emission data are derived from the research group at Peking University 8 (http://inventory.pku.edu.cn/ home.html, Wang et al., 2014a); the Chinese desert (sand) distribution dataset 9 is provided by the National Tibetan Plateau Data Center (http://poles.tpdc.ac.cn/zh-hans/data/122c9ac2-53ee-10 4b9a-ae87-1a980b131c9b/; Wang et al., 2013a).

Figure 3d depicts the spatial distribution of the relative contribution of three components to fluorescence, with the regional averages given in Figure 3e–g. In SEIM, the greatest contribution is that of HULIS-1 (47 %), followed by HULIS-2 (38 %) and PRLIS (15 %), indicating that the signal is dominated by local soil/dust sources, consistent with the local environment (Figs. 2 and 3d). HULIS-2 plays a greater role in SNC, where it accounts for half of the total fluorescence signals, with HULIS-1 is being next most important. This difference in key components between SEIM and SNC illustrates the change in the primary source of fluorescence intensity. Indeed, although HULIS-2 might be derived from any

1	combination of terrestrial, anthropogenic, and microbial sources (Osburn et al., 2016; Stedmon et al.,
2	2003; Yamashita et al., 2008; Zhang et al., 2009), human activity (e.g., agriculture, industrial emissions)
3	is most intensive in SNC (Figs. 1a and 3d; Guo and Hu, 2022). Therefore, our combined analysis suggests
4	that anthropogenic source is the main contributor to seasonal snow in SNC. The conclusion is also in
5	good agreement with the previous study (Zhou et al., 2019b). As in SNC, HULIS-2 also represents
6	approximately half of the fluorescence signal in both NNC and CBM. The background environment of
7	NNC is similar to that of SNC, with dense urban cities and populations. In the CBM, which is heavily
8	forested (Fig. 1a; Guo and Hu, 2022), the long-range transport of anthropogenic pollutants is responsible
9	for the dominance of HULIS-2, as discussed in Sect. 3.1. HULIS-1 accounts for less than PRLIS in both
10	NNC and the CBM, which we posit reflects the concealment of bare soil surfaces by deep snow and the
11	importance of biological processes due to the heavy vegetation cover. PRLIS accounts for >50 % of the
12	total fluorescence signal in NEIM, followed by HULIS-2; HULIS-1 contributes relatively little in this
13	region. We attribute this pattern to both the extensive grassland and forest cover, which obscures bare
14	soil surfaces and the distance from anthropogenic pollution. These together serve to amplify the
15	importance of biological processes (Zhou et al., 2019a). Taken as a whole, the respective contributions
16	of HULIS-1, HULIS-2, and PRLIS to the fluorescence signals in our study area are ~30 %, ~50 %, and
17	\sim 20 %, respectively. We note that these findings correspond well with the background environmental
18	conditions.





2 Figure 4: (a) Variations in fluorescence-derived indices among the five groups. Boxes denote the 25th and 75th 3 quantiles, and horizontal lines represent median values. Averages are shown as small boxes, the whiskers 4 denoting maximum and minimum data. (b) Comparison plots of HIX versus FI for the seasonal snow surface 5 samples (colored dots) from northeastern China, together with the average and standard deviation of different 6 types of WSOC (grey markers) adapted from Arctic aerosols (^a Fu et al., 2015), Spanish urban aerosols (^b 7 Mladenov et al., 2011), seasonal snowpack in northwestern China (^c Zhou et al., 2019b), Chinese lakes and 8 rivers (^{d, e} Zhou et al., 2017), rainwater from Rameswaram, India (^f Salve et al., 2012), fog water from Tai 9 Mountain, China (^g Birdwell and Valsaraj, 2010), cave water from Frasassi Caves, Italy (^h Birdwell and Engel, 10 2010), and spring water in the USA (ⁱ Birdwell and Engel, 2010). Shaded areas represent mixed WSOC 11 signatures.

12 The FI, BIX, and HIX indices reveal spatial variability in fluorescence characteristics and thus permit

13 the tracing of potential sources. Regionally averaged FI, BIX, and HIX values are depicted in Figure 4a.

14 Our results show that, the FI values are 1.34-1.94 (mean = 1.49), BIX are 0.46-1.17 (mean = 0.74), and

- 15 HIX are 0.30–0.84 (mean = 0.64). By comparison, reported mean FI, BIX, and HIX values for seasonal
- snow in Xinjiang (northwestern China) are 1.42, 0.76, and 0.55, respectively (Zhou et al., 2019b),
- 17 suggesting that the impacts of WSOC humification and aromaticity are slightly greater in the study area

1	than in Xinjiang. This outcome implies a relatively strong terrigenous signal and correspondingly weaker
2	biogenic signal in the seasonal snowpack of northeastern China. Regionally, SEIM exhibits the lowest
3	FI (mean = 1.40) and BIX (mean = 0.49) values but the largest HIX value (mean = 0.80), further
4	confirming the strong influence of highly aromatic, terrestrially derived WSOC in this region relative to
5	the others. In contrast, NEIM returns the highest FI (mean = 1.74) and BIX (mean = 0.89) values, but the
6	lowest HIX value (mean = 0.38), indicating the dominance of low-aromatic WSOC of microbial origin.
7	Intriguingly, our results reveal that FI and BIX rise generally with decreasing (increasing) fractional
8	contributions of HULIS-1 (PRLIS), whereas HIX exhibits a similar but contrasting pattern. Together, the
9	comprehensive dataset described above verifies the regional variability in the terrestrial contributions to
10	WSOC, with SEIM > SNC > NNC > CBM > NEIM. This pattern is reversed for microbially sourced
11	WSOC.
12	Figure 4b illustrates HIX versus FI as a scatterplot, compared with published data for different sample
13	types; the shaded area depicts the region in which the FI value is >1.4 but <1.9. In general, FI exhibits a
14	rising trend with declining HIX across northeastern China. For both SEIM and SNC, FI occupies a
15	narrow range centered on 1.4, indicating either a predominantly terrestrial or mixed origin. We note that
16	these results are comparable to those of fog water from the Tai Mountain, Arctic atmospheric aerosols,
17	and seasonal snow in northwestern China (Birdwell and Valsaraj, 2010; Fu et al., 2015; Zhou et al.,
18	2019b). Further, we highlight that HIX values are marginally higher in SEIM than elsewhere, suggesting
19	a stronger influence from highly humified WSOC that probably reflects the extensive deserts and
20	exposed earth in this region. FI values for NNC and CBM fall within the range of 1.4–1.7 and thus reflect
21	a mixed origin, in line with previous data from urban aerosols in Spain and Chinese river water samples
22	(Mladenov et al., 2011; Zhou et al., 2017). When combined, the FI and HIX values for NNC and CBM

snowpack indicate that WSOC in these regions bears a stronger terrestrial signature than do water
 samples from Chinese lakes and Italian cave (Birdwell and Engel, 2010; Zhou et al., 2017). Finally, FI
 values for NEIM fall within a range of 1.6–2.0, comparable to values for spring water in the USA
 (Birdwell and Engel, 2010), thus implying a predominantly microbial or mixed origin.

5

3.3 Comparisons of fluorescence and absorption characteristics

6 Figure 5a describes the spatial distribution of $a_{WSOC}(280)$ as WSOC absorption in the snowpack of 7 northeastern China; Figure 5b depicts TFV as a measure of the spatial distribution of absolute WSOC 8 fluorescence intensity for comparison. In general, TFV and a_{WSOC} (280) both exhibit large spatial 9 variability in the range of 690-18600 RU nm² and 0.42-16.98 m⁻¹, respectively. Regional mean values are 7700 \pm 2800 RU·nm² (TFV) and 6.90 \pm 2.39 m^{-1} ($a_{WSOC}(280))$ for SEIM, 12400 \pm 4300 RU·nm² 10 11 (TFV) and 11.48 \pm 3.96 m⁻¹ (a_{WSOC}(280)) for SNC, 7800 \pm 500 RU·nm² (TFV) and 7.02 \pm 0.88 m⁻¹ $(a_{WSOC}(280))$ for NNC, 3900 ± 2500 RU·nm² (TFV) and 3.97 ± 2.46 m⁻¹ ($a_{WSOC}(280)$) for the CBM, 12 13 and 1000 \pm 300 RU·nm² (TFV) and 0.83 \pm 0.23 m^{-1} ($a_{WSOC}(280)$) for NEIM. We note that both 14 distributions are consistent in space (Fig. 5e), with the highest concentrations in SNC and the lowest in 15 NEIM. Moreover, the $a_{WSOC}(280)$ value for SNC is an order of magnitude larger than that for NEIM, 16 implying that the impact of WSOC on snow albedo at UV wavelengths is significant in SNC but less 17 notable in NEIM in general (see Sect. 3.5). Previous work has reported a similarly broad range of 18 snowpack $a_{WSOC}(280)$ (0.15–10.57 m⁻¹) in northwestern China (Zhou et al., 2019b).



Figure 5: Spatial distribution of (a) a_{WSOC}(280) (m⁻¹), (b) TFV (RU nm²), (c) MAC₂₈₀ (m² g⁻¹) and (d) NFV
(RU nm² (mg L⁻¹)⁻¹). Regional averages for (e) a_{WSOC}(280), TFV, and (f) MAC₂₈₀, NFV for the five groups.
Error bars in (e) and (f) represent the standard deviations of a_{WSOC}(280), MAC₂₈₀, TFV, and NFV for the five
groups, respectively.



HULIS-2 fraction measured in the NNC samples (Fig. 3g), as the lower-oxygenated BrC (e.g., HULIS-

2 2) has a higher absorption capacity (Browne et al., 2019).

3 Scatterplots of a_{WSOC}(280) versus TFV, F_{max}(HULIS-1), F_{max}(HULIS-2), and F_{max}(PRLIS) are provided 4 in Figure S4 to further demonstrate the close relationship between fluorescence and the absorption 5 characteristics of WSOC in our snow samples. As samples Q480, Q484, and Q477 deviate considerably 6 from the respective confidence intervals, we did not include them in our analyses. Surprisingly, we found 7 that TFV is closely correlated to $a_{WSOC}(280)$, with P < 0.001 and all datapoints located close to the line 8 of best fit, indicating that the three components (HULIS-1, HULIS-2, PRLIS) contributing to the total 9 fluorescence are also responsible for the majority of absorption. For each component, our data show that 10 F_{max}(HULIS-1) is most closely correlated with a_{WSOC}(280), followed by F_{max}(HULIS-2). The correlation 11 between $F_{max}(PRLIS)$ and $a_{WSOC}(280)$ is the weakest, yet it is still significant (P < 0.001). Together, our 12 results imply that HULIS-1 is probably the greatest contributor to light absorption, with PRLIS being the 13 least important.

14 3.4 Fractional contributions of different WSOC components to light absorption

15 In this study, we applied a multiple linear regression method comparable to that of Chen et al. (2019a) 16 to derive the fractional contribution of each WSOC component to light absorption. Table S3 lists the 17 statistical results of the fitted light absorption coefficient, based on the F_{max} data for three fluorescent 18 components of EEM analysis. As the fitted results can explain ~94 %-99 % of the variance in measured 19 light absorption within the 280-400 nm range, we conclude that the fusion of multiple fluorescent 20 components is an effective means of describing most of the spatial features of WSOC light absorption 21 throughout northeastern China. Accordingly, the wavelength-dependent fractional contributions of each 22 component of light absorption in this band (280-400 nm) are reported in Figure 6. For the entire study area, light absorption is dominated by high-oxygenated HULIS-1, which accounts for ~56 %-65 % of
the contribution across UV wavelengths. Furthermore, the contribution of HULIS-1 first increases with
increasing wavelength from 280 to ~340 nm, and then decreases. In contrast, HULIS-2 exhibits a
valley-type pattern in terms of its fractional contribution at wavelengths between 280 and 400 nm and is
responsible for ~19 %-30 % of all light absorption.





Figure 6: Relative contributions of the three components to the total absorption of samples in (a) the whole
study area and (b-f) each of the five groups.

9 PRLIS contributes the least (~12 %-17 %) to light absorption and exhibits a similar wavelength-10 dependent pattern to HULIS-1. These results are consistent with the qualitatively comparative analysis 11 described in Sect. 3.3. Previous studies have also highlighted this dominance of high-oxygenated 12 compounds in WSOC light absorption, based on samples impacted by naturally and anthropogenically 13 derived soils (Zhou et al., 2022). Conversely, the total absorption coefficient of WSOC decreases with 14 the increasing wavelength between 280 and 400 nm, in accord with previous studies (Andreae and 15 Gelencser, 2006; Chakrabarty et al., 2010; Gustafsson et al., 2009; Wu et al., 2019). The AAE lies 16 primarily between 5.0 and 8.0 (mean = 6.6) in the range of 280–400 nm, which is in agreement with

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results from snow collected from the Arctic, the northern Tibetan Plateau, and northwestern China (Voisin et al., 2012; Yan et al., 2016; Zhou et al., 2021).

3 For each component, the wavelength-dependent variability in light absorption is similar among all five 4 regions, although the magnitude of each contribution varies from region to region. Moreover, compared 5 with the spectral results, that the solar-radiation-weighted broadband light absorption is a more 6 meaningful parameter for researchers studying climate change and atmospheric radiation. Therefore, the 7 broadband results in Figure 7a for 280-400 nm absorption contributions-HULIS-1 (62 %), HULIS-2 8 (21%), and PRLIS (17%)—are average values for the whole study area. On a regional scale, the HULIS-9 1 contribution to light absorption (280–400 nm) follows the spatial pattern SEIM > SNC > NNC > CBM >10 NEIM. We note that HULIS-1 dominates light absorption in SEIM, SNC, and NNC but has a minor 11 impact in NEIM compared with the other two components. In contrast, the impact of HULIS-2 varies 12 only slightly among the five regions, with the greatest contributions in NNC and CMB, and the lowest 13 in NEIM. The contribution of PRLIS is essentially opposite to that of HULIS-1, being dominant in NEIM 14 but of relatively minor importance elsewhere. As shown in Figure 7b, light absorption contributions at 15 280 nm are consistent with the broadband results (Fig. 7a) in terms of the regional pattern, although 16 specific values differ because of the different wavelength-dependent properties of light absorption for 17 the three WSOC components.



Figure 7: Regional averages for the relative contributions of the three fluorescent components to light
absorption at wavelengths of (a) 280–400 nm and (b) 280 nm.

4 We find it noteworthy that, for each component, the overall regional pattern of its contribution to light 5 absorption aligns with its impact on fluorescence signals, thereby confirming the viability of the 6 attribution analysis employed in this study. Nonetheless, we observed that the magnitude of each 7 component's contribution varies relative to its respective fluorescence signal. For instance, HULIS-1 8 returns a greater contribution to light absorption than its fluorescence signal, in contrast to HULIS-2. 9 One plausible explanation for this discrepancy is that the apparent fluorescence quantum yields (AQY), 10 which are essentially the ratio of fluorescence intensity versus absorption intensity, are different for each 11 component. Indeed, in their comprehensive field-based study of BrC fluorescence and absorption

properties in northern China, Wen et al. (2021) reported that the AQY of WSOC decreases with
 increasing HIX, meaning that components with higher HIX values, such as HULIS-1, have lower AQY
 than HULIS-2. Thus, the contribution of HULIS-1 to the fluorescence signals will be smaller than its
 contribution to light absorption, and vice versa for HULIS-2.

5 3.5 Albedo reduction and radiative forcing attributed to snowpack WSOC

6 The strong light absorption of WSOC in UV bands has important ramifications for snow albedo and 7 radiative forcing throughout northeastern China. However, owing to the chemical and optical complexity 8 of WSOC components, quantitative estimates for snowpack light absorption remain poorly understood. 9 For example, although prior work in northeastern China has focused on BC (Wang et al., 2013b) and 10 other water-insoluble light-absorbing particles (Wang et al., 2017; Zhao et al., 2014) via field 11 measurements, model simulations, and satellite remote sensing (Pu et al., 2019), the specific impacts of 12 WSOC have not been studied. Consequently, ours is the first study to report on the impact of WSOC on 13 snow albedo and radiative forcing in northeastern China and to compare these data with BC results to 14 highlight the non-negligible role of WSOC.





Figure 8: (a) and (b): Simulated snow spectral albedo and broadband albedo reductions—under various
contamination scenarios and for different regions—for old snow (radius = 1000 μm). (c), (d) Simulated snow
spectral albedo and broadband albedo reductions—under various contamination scenarios and for different
regions—for fresh snow (radius = 100 μm). Colors represent the different types of snow (pure snow, BC- or
WSOC-contaminated snow, and snow polluted by both WSOC and BC).

Figure 8 shows the regional-mean spectral snow albedo as well as the reduction in albedo due to WSOC,

8 BC, and WSOC + BC. We assume a snow radius of 100 μ m for fresh snow and 1000 μ m for old snow. 9 Our findings reveal that WSOC induces a marked decline in albedo within the UV and short-wave VIS 10 bands, with the magnitude of albedo reduction growing rapidly as wavelength shrinks owing to the large 11 AAE value of WSOC. In comparison, BC induces a widespread albedo reduction spanning the UV to 12 NIR bands, and wavelength-dependent variations are significantly smaller than those of WSOC. For VIS 13 and NIR, the reduction in albedo is dominated by BC, whereas the impacts of WSOC and BC are 14 comparable at UV wavelengths, a pattern that is consistent with the results of studies of atmospheric 15 aerosols (Shamjad et al., 2016). We note that these characteristics persist throughout northeastern China 16 despite regional variability in environmental conditions and snowpack types (old vs. fresh snow).

1	For broadband wavelengths, our results indicate that the WSOC-induced (mean = $3.6 \ \mu g \ g^{-1}$) albedo
2	reduction for 280-400 nm wavelength in old (fresh) snow is 0.16 (0.05) across the whole study area,
3	which corresponds to approximately 50 % (46 %) the impact of BC (mean = 0.6 μ g g ⁻¹). Regionally, the
4	greatest decline in albedo occurred in SNC, where a mean WSOC of 5.7 μ g g ⁻¹ resulted in a reduction
5	of 0.20 (0.06) in the 280-400 nm range for old (fresh) snow. In contrast, the smallest decline in albedo
6	was observed in NEIM, with a reduction of 0.06 (0.02) resulting from an average WSOC concentration
7	of 0.5 μ g g ⁻¹ . Compared with the UV bands, a WSOC-induced albedo reduction of 0.03 (0.009) over the
8	UV–NIR range (280–1500 nm) accounts for only ~19 % (17 %) of that due to BC in the study area. The
9	regional mean for old (fresh) snow falls in the range of 0.01–0.04 (0.003–0.012), with the highest (lowest)
10	values occurring in SNC (NEIM). However, we observed the highest ratio of WSOC- to BC-induced
11	albedo reduction in NEIM. Together, these results indicate that WSOC plays a potentially important role
12	in altering UV snow albedo in NEIM, despite its relatively low concentrations in the regional snowpack.





Figure 9: Radiative forcing due to different pollutants in (a) old or (b) fresh snow. Gray, blue, and red indicate
the radiative forcing of BC, WSOC, and BC + WSOC, respectively.

Radiative forcing is an important index that directly reflects the impact of snowpack WSOC on the
regional radiation balance and climate (Beres et al., 2020). Previous studies have tended to focus on
calculating instantaneous radiative forcing values; however, in reality, time-averaged results are more
valuable for climate research. Here, we present data on the daily mean radiative forcing due to WSOC,
BC, and WSOC + BC (Fig. 9). For northeastern China, the mean radiative forcing of WSOC in old (fresh)
snow is 3.8 (0.8) W m⁻², with regional mean values varying from 1.2 (0.2) to 4.9 (1.0) W m⁻². Zhou et
al. (2021) reported daily mean radiative forcing by regional WSOC (0.6–7.1 µg g⁻¹) was between ~0.04

1 and ~ 0.6 W m⁻² for northwestern China, which is comparable to our values in fresh snow. Furthermore, 2 the ratio of WSOC-driven to BC-driven radiative forcing varies within the range of 10 %-32 % (10 %-3 31 %) for old (fresh) snow, which is consistent with the results of our calculated albedo reductions. These 4 results confirm that the role of WSOC must not be ignored in discussions about radiative balance in 5 northeastern China. Similarly, the sizeable impact of WSOC on the absorption of UV radiation has the 6 potential to influence biogeochemistry (Helms et al., 2013; Seekell et al., 2015), and snow photochemical 7 processes (e.g., photolysis of nitrate (NO₃⁻) and nitrite (NO₂⁻) in snow, in addition to the release of NO_x 8 $(NO + NO_2 \text{ and HONO})$. Snow photochemistry is beyond the scope of this study, however, the high 9 concentrations of WSOC and nitrate (not shown) pollution in northeastern China make this a logical next 10 step for research in this field.

11 4 Conclusions and atmospheric implications

12 During 2020 and 2021, we collected 34 surface samples of seasonal snow from sites throughout 13 northeastern China to investigate the fluorescence characteristics, optical properties, and radiative effects 14 of snowpack WSOC. With an average concentration of WSOC of $3.6 \pm 3.2 \ \mu g \ g^{-1}$, our results returned regional mean values of $3.4 \pm 1.5 \ \mu g \ g^{-1}$ (SEIM), $5.7 \pm 3.7 \ \mu g \ g^{-1}$ (SNC), $2.7 \pm 0.8 \ \mu g \ g^{-1}$ (NNC), $2.0 \pm 1.5 \ \mu g \ g^{-1}$ (NNC), $2.0 \pm 1.5 \ \mu g \ g^{-1}$ (SIC), $2.7 \pm 0.8 \ \mu g \ g^{-1}$ (NNC), $2.0 \pm 1.5 \ \mu g \ g^{-1}$ (SIC), $2.7 \pm 0.8 \ \mu g \ g^{-1}$ (SIC), $2.8 \ \mu g \ g^{-1$ 15 1.3 μ g g⁻¹ (CBM), and 0.5 \pm 0.2 μ g g⁻¹ (NEIM), indicating a considerable degree of regional variability 16 17 of WSOC mass loadings. Measured values of WSOC fluorescence intensity (690-18600 RU nm²) and light absorption $(0.4-17 \text{ m}^{-1})$ are also highly variable. 18 19 Moreover, we also used EEMs and PARAFAC to identify three fluorescence WSOC components 20 prevalent in northeastern China and analyzed their regional differences. In SEIM, which is characterized 21 by desert and bare soil surfaces, the signal of high-oxygenated and terrigenous HULIS-1 is dominant

22 (47 %). The high degree of humification and minimal bioavailability of WSOC indicate that snowpack

1	WSOC originates primarily from soil source. In contrast, the autochthonous PRLIS signal (58 %)
2	dominates in remote and clean NEIM. Low-oxygenated and anthropogenic HULIS-2 dominates the
3	densely populated and intensively farmed SNC (51 %) and NNC (57 %) regions, leading us to conclude
4	that the snowpack WSOC in SNC and NNC are influenced more by anthropogenic source. In CBM of
5	the forest environment, the impact of long-distance transport of pollutants is greater than that of the
6	background environment. The above conclusions are also verified by fluorescence-derived indices.
7	We employed multiple regression analysis to estimate the fractional contributions of different WSOC
8	components to snowpack light absorption. Throughout our study area, HULIS-1 tends to be the greatest
9	contributor (~56 %–65 %) over the 280–400 nm range, followed by HULIS-2 (~19 %–30 %) and PRLIS
10	(~12 %-17 %). In contrast to its primary role in fluorescence, the contribution of HULIS-2 to light
11	absorption is relatively low across all regions, potentially reflecting the variable molecular structure of
12	different components. Finally, we highlight that the average RF due to WSOC in old (fresh) snow in
13	northeastern China is 3.8 (0.8) W m ⁻² , which is equal to 19 % (17 %) of the BC-induced radiative forcing.
14	Therefore, we demonstrated the important impacts of WSOC on the snow energy budget and potentially
15	on triggering snow photochemistry. We indicate that our study could contribute to the understanding
16	of carbon cycling processes, regional air quality, hydrological processes, and climate change in the earth
17	systems. For example, the abundant WSOC concentrations measured in this study imply the
18	significant carbon input from the atmosphere to the snowpack through wet or dry depositions in
19	northeastern China. While the complex chemical compositions of snowpack WSOC could further
20	influence the carbon balance of the snow environment by affecting microbial activities (Stedmon et al.,
21	2007). The strong absorption properties of WSOC in the UV-Vis band also imply its important role in

22 initiating snow photochemistry (McNeill et al., 2012). It will change the composition of organic

1	compounds in the snow in turn (Grannas et al., 2007), and affect the surrounding air quality by releasing
2	oxidizing gas like NO _x into the atmosphere (Zatko et al., 2013). Moreover, the non-negligible influence
3	of WSOC on the snow albedo and radiative effect indicate that it could not only accelerate snow melting,
4	but also change the periods and mass of water and carbon exchange between snowpack and underlying
5	soils or vegetation (Meyer and Wania, 2008), and potentially affect regional climate through changing
6	the surface radiative balance (Beres et al., 2020).

2	following data repository: https://doi.org/10.5281/zenodo.6541956.
3	Supplement. The supplement related to this article is available online at:
4	Author contributions. XN and WP designed the study and wrote the first draft with contributions
5	from all coauthors. XN designed and conducted the lab experiments with the assistance of YZ and
6	HW. XN processed the data with the assistance of DW and TS. XN, WP, YC, YX, TS designed
7	and conducted the field campaign. XW supervised this study. All co-authors commented on the
8	paper and improved it.
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16	

Data availability. Data presented and used throughout this study can be accessed through the

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