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# Source identification and characteristics of dissolved organic matter and disinfection by-product formation potential using EEM-PARAFAC in the Manas River, China

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Dissolved organic matter (DOM) is ubiquitous in natural water and reacts with disinfectants to form disinfection by-products (DBPs). The analysis of DOM in raw water is helpful in evaluating the formation potential of DBPs. However, there is relatively little research on the DOM identification of raw water in northern China. In this study, the sources and characteristics of DOM were investigated in the samples collected from the Manas River. Dissolved organic carbon (DOC),  $UV_{254}$ , specific ultraviolet absorbance, and fluorescence indices (fluorescence index, humification index, and biological index) were measured to characterize the DOM, and trihalomethanes (THMs) were quantified following formation potential tests with free chlorine. The maximum amount of total trihalomethane formation potential (THMsFP) was 225.57  $\mu$ g L<sup>-1</sup>. The DOM of the Manas River consisted of microorganisms and soil resources. The excitation–emission matrix combined with parallel factor analysis (EEM-PARAFAC) identified microbial humus (C1, 54%) and tryptophan-like protein (C2, 46%). PARAFAC components were evaluated as the precursor surrogate parameters of THMsFP. Additionally, the linear THMsFP correlation was stronger with C1 + C2 (r = 0.529, p < 0.01) than with C1 (r = 0.485, p < 0.01). Thus, C1 + C2 is an accurate THMsFP precursor surrogate parameter for the Manas River, and the use of fluorescence spectroscopy may be a robust alternative for predicting DOC removal.

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# Introduction

Dissolved organic matter (DOM) is defined as the fraction of organic matter that passes through a 0.45 mm filter; it is ubiquitous in surface water bodies and poses a challenge to the effectiveness of water treatment processes. Until now, disinfection has been effective for inactivating pathogenic microorganisms in drinking water treatment. However, commonly used disinfectants can react with the organic matter in water to generate diverse disinfection by-products (DBPs), such as trihalomethanes (THMs) and haloacetic acids (HAAs), some of which have been reported to have adverse effects on human health, inducing cytotoxicity, genotoxicity, teratogenicity, and carcinogenicity. Therefore, a comprehensive understanding of DOM composition and effective removal of DBP is essential to ensure the success of a water treatment plant and to achieve the desired quality for intended water use.

DOM is the main precursor of DBP. Given its complex composition and low concentration, surrogate parameters were

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developed to predict the removal of DOM through water treatment and estimate its reactivity toward DBP formation. <sup>8,9</sup> These surrogate parameters include dissolved organic carbon (DOC), ultraviolet (UV) absorbance, and specific ultraviolet absorbance (SUVA). <sup>10,11</sup> However, they provide limited information and are time consuming.

Excitation-emission matrix (EEM) fluorescence spectroscopy is a rapid, sensitive, and informative method, which has been widely applied to trace the quality and quantity of DOM. It has been proven to have the potential to assess DBP formation potentials (FPs). 12-14 An EEM spectrum is constructed using emission wavelength, excitation wavelength, and fluorescence intensity. However, it contains a vast amount of fluorophore information, and this is difficult to unscramble and characterize because of the influence of fluorescence overlap and matrix effect. On the other hand, a fluorescence EEM coupled with parallel factor analysis (EEM-PARAFAC) can efficiently extract the information contained in EEMs and eliminate interference among components. Therefore, it can provide better insights into the DOM variation in the water treatment processes, 15 and has been applied to a variety of natural and engineered systems. The PARAFAC model can be applied to analyze large data sets of hundreds or thousands of EEMs,

making statistical analysis more reliable. It makes it possible to measure the fluorescence characteristics of various water sources and helps to identify the effective fluorescent groups in organic substances, including humic acid, proteins, fulvic acid, etc. EEM-PARAFAC method divides fluorescent organic compounds into several independent components according to their unique properties and structures, which provides a basis for further understanding the dynamic changes of fluorescent components.16-20 The intensities and the relative distributions of PARAFAC components are representative of their levels and the chemical composition of fluorescent DOM, and can be developed as surrogates for conventional water quality parameters, treatability of DOM, DBPsFPs, and performance of treatment processes. 11,21-24 For example, humic-like components identified by PARAFAC were linked to different sources of DOM (e.g., terrestrial and microbial) and were shown to have dissimilar treatability during water and waste treatment.21,25,26 Some studies have indicated a high correlation between trihalomethane formation potential (THMsFP) and some PAR-AFAC components in natural aquatic environments<sup>27,28</sup> and in laboratory-scale water treatments.29 In another study, five EEM-PARAFAC components, including four terrestrially derived humic-like substances (C1, C2, C3, and C4) and one tryptophanlike substance (C5), were identified in the sub-alpine Lake Tiancai.25 Although the application of the EEM-PARAFAC in a full-scale drinking water treatment plant of Australia has already been reported, it is noteworthy that the ambient climate (e.g., flow rate, rainfall pattern, and temperature) largely governs the relative chemical composition of DOM.30 This indicates that the specific water quality differs from source to

source, and the composition of DOM may be quite different

even though similar fluorescence levels may be present on the maps, thus resulting in varying correlations. As such, the values of EEM-PARAFAC for tracking DOM, DBPsFP, and treatment system performance require further discussion.

The main objects of this study were (1) to investigate the variation in DOM characteristics and DBP precursors in the water source of drinking water plants by fluorescence PARAFAC and (2) to determine the relationships between DOM spectroscopic properties and DBPFPs to assess the application potential of fluorescence spectroscopy for monitoring DBPs in raw water, providing valuable information for realtime monitoring of DBP, and optimization of the drinking water treatment process.

# 2. Methods

### 2.1. Study area and sample collection

The Manas River (43°24′-45°12′ N, 85°41′-86°32′ E) is located in the northern part of Xinjiang, China. It originates in the Tian Shan Mountains and is an inland river. Its ecological status plays a key role in the development of the economic belt of the northern slope of Xinjiang and the industrial, agricultural, and trade development of the Urumqi, Changji, and Shihezi urban agglomerations. This study used a section of the river that flows through Shihezi City, which is approximately 30.4 km long. As shown in Fig. 1, 12 sampling sites (M01–M12) were established in this section of the river, which were divided into upstream (M01–M04), midstream (M05–M08), and downstream (M09–M12). Samples were collected from October to December in 2020 and once per month. The samples were collected from surface water (0–0.5 m) with a sampling device, each sampling cite close to the edge of the river. Three replicates were collected

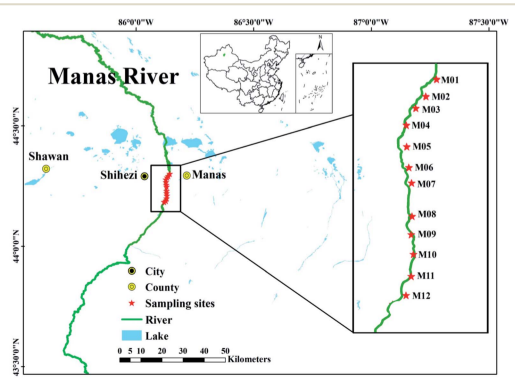


Fig. 1 Sampling map showing the Manas River Basin, China.

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in 1.25 L clean plastic bottles from each sampling site, and immediately transferred to the laboratory. All the samples were stored at 4 °C before further treatment and analysis. Pretreatment of the samples included filtration through a pre-washed 0.45 µm cellulose acetate membrane filter. All analytical characterizations were performed within 24 h of sample collection.

### 2.2. Analytical characterization

Electrical conductivity (EC), temperature, and pH of the samples were measured on site using portable meters (2100Q and HQ40d, HACH, Japan). The NH<sub>3</sub>-N and total dissolved solids (TDS) concentrations were determined using the standard examination methods for drinking water parameters, China (GB/T 5750.5-2006). The DOC concentrations were measured using a TOC-VWP total organic carbon analyzer (Shimadzu, Japan). Samples were acidified before measurement to purge inorganic carbon. Therefore, the amount of nonpurgeable organic carbon (NPOC) was equal to the DOC. Furthermore, UV absorbance at 254 nm (UV<sub>254</sub>) was measured using a Varian Cary 50 UV-Vis spectrophotometer and by consulting Standard Method 5910. SUVA<sub>254</sub> was calculated as the 100-fold ratio of UV<sub>254</sub> to DOC, which correlated positively with the aromaticity of DOM.

### 2.3. EEM and EEM-PARAFAC analysis

EEM fluorescence spectroscopy was conducted using a Hitachi F-4700 luminescence spectrometer equipped with a xenon excitation source. The scan rate was set at 1200 nm min<sup>-1</sup>, and the excitation/emission slits were 5 nm band-pass. The scanning field was set at emission (Em) spectra ranging from 280 to 550 nm and excitation (Ex) spectra from 200 to 400 nm, with an interval of 5 nm. Milli-Q water was used as a control and was subtracted to eliminate the effect of Raman scattering.

Fluorescence data were modeled in MATLAB software using the DOMFluor toolbox.31 Outlier samples, which were identified using leverage comparison, were removed, and PARAFAC models with two to eight components were generated. Split-half analysis was used to determine the number of components. The fluorescence intensity of each component (i.e., F<sub>max</sub> of C1, C2, or C3) was used to

represent the relative concentration. Additionally, three common DOM quality indices were obtained from the EEMs: (1) biological index (BIX), which is indicative of the relative significance of microbial or biological DOM, (2) humification index (HIX), which is an indicator of humification degree or humic character, and (3) fluorescence index (FI) to distinguish algal and microbial sources from terrestrial DOM sources. The BIX was determined as the ratio of Em ranging between 380 nm and 430 nm with Ex at 310 nm; HIX was obtained by dividing the integrated intensity from Em between 435 and 480 nm by that of Em ranging between 300 and 345 nm at Ex 254 nm; FI is the ratio of the fluorescence intensity at Em 450 nm to that at Em 500 nm, given that  $Ex = 370 \text{ nm.}^{13,32,33}$ Correlation analyses were conducted using SPSS 22.0. Correlation coefficients (r), rather than  $R^2$ , were used to differentiate negative and positive correlations.

### 2.4. DBP analysis

Chlorine disinfection experiments were performed in dark brown bottles at 25 °C for three days. Phosphate solution (KH<sub>2</sub>PO<sub>4</sub>/NaOH) was used to maintain the pH of the sample at 7.0  $\pm$  0.1. To ensure the presence of measurable residual chlorine in each sample, chlorine was added at a dose of 5 mg Cl<sub>2</sub> per mg C.14 Sodium hypochlorite (NaOCl, Sinopharm Chemical Reagent Co., China) was used as a disinfectant. The residual chlorine was quenched using ascorbic acid, and the samples were collected for DBP analysis.

Typical DBPs include chloroform (TCM), dichlorobromomethane (DCBM), dibromochloromethane (DBCM), and tribromomethane (TBM). These were extracted and measured according to the U.S. EPA Method 551.1.34 All DBPs were determined using a gas chromatography/electron capture detector (GC-ECD; 7890A, Agilent, USA).

### 3. Results and discussion

### DOM quantity and quality

Raw water absorbance (UV<sub>254</sub>), DOC, pH, chloride, conductivity, TDS, NH3-N, and TN (total nitrogen) are summarized in Table 1. Between October and December, Manas River had a pH

Table 1 Summary of water characteristics of surface water samples collected from Manas River

Sample/ index	Absorbance $(UV_{254})$	$\frac{\mathrm{DOC}}{\mathrm{(mg\ L^{-1})}}$	рН	Chloride (mg L <sup>-1</sup> )	Conductivity (ms m <sup>-1</sup> )	${ m TDS} \atop { m (mg~L^{-1})}$	NH <sub>3</sub> -N (mg L <sup>-1</sup> )	TN
M01	$0.030 \pm 0.05$	$4.698 \pm 2.29$	$7.40 \pm 0.12$	$5.33 \pm 0.54$	$38.6 \pm 1.03$	$302.5 \pm 2.11$	$0.66 \pm 0.07$	$0.077 \pm 0.07$
M02	$0.027\pm0.11$	$6.069 \pm 3.35$	$7.45\pm0.08$	$\textbf{5.38} \pm \textbf{1.13}$	$37.5 \pm 0.98$	$277.5 \pm 1.09$	$0.62\pm0.13$	$0.129\pm0.11$
M03	$0.031 \pm 0.03$	$6.707 \pm 2.14$	$7.39 \pm 0.18$	$4.85\pm0.66$	$37.6 \pm 1.24$	$276.2 \pm 1.26$	$0.75\pm0.06$	$0.031\pm0.14$
M04	$0.028\pm0.04$	$6.229 \pm 1.37$	$7.50\pm0.21$	$\textbf{4.75} \pm \textbf{1.47}$	$36.0\pm2.07$	$268.7 \pm 2.57$	$\textbf{0.68} \pm \textbf{0.22}$	$\textbf{0.147} \pm \textbf{0.12}$
M05	$0.027\pm0.05$	$6.263\pm0.16$	$7.39 \pm 0.16$	$5.08 \pm 1.15$	$37.6\pm1.63$	$251.2\pm1.94$	$0.64 \pm 0.14$	$\textbf{0.048} \pm \textbf{0.15}$
M06	$0.025\pm0.13$	$5.314 \pm 2.85$	$7.43 \pm 0.17$	$\textbf{5.17} \pm \textbf{0.84}$	$37.2\pm1.09$	$321.3\pm1.57$	$0.63\pm0.06$	$0.041\pm0.23$
M07	$0.026\pm0.06$	$6.000 \pm 2.73$	$7.54\pm0.25$	$5.25\pm0.61$	$37.4 \pm 0.77$	$321.3\pm3.02$	$0.64\pm0.17$	$0.082\pm0.17$
M08	$0.031\pm0.16$	$\textbf{4.489} \pm \textbf{2.66}$	$\textbf{7.56} \pm \textbf{0.09}$	$\textbf{5.48} \pm \textbf{0.57}$	$37.1\pm0.85$	$253.8 \pm 4.18$	$\textbf{0.62} \pm \textbf{0.09}$	$0.030\pm0.05$
M09	$0.027\pm0.13$	$5.247 \pm 1.57$	$7.56\pm0.23$	$5.50\pm1.29$	$46.0\pm1.25$	$240.0\pm1.04$	$\textbf{0.67} \pm \textbf{0.27}$	$0.036\pm0.04$
M10	$0.026\pm0.04$	$5.558 \pm 1.44$	$7.60\pm0.19$	$5.30 \pm 0.79$	$34.4 \pm 2.44$	$228.7 \pm 1.19$	$0.67 \pm 0.19$	$0.151\pm0.19$
M11	$0.025\pm0.23$	$\textbf{5.287} \pm \textbf{3.08}$	$\textbf{7.51} \pm \textbf{0.13}$	$5.02\pm1.22$	$34.7\pm1.72$	$202.5\pm0.99$	$\textbf{0.66} \pm \textbf{0.04}$	$0.075\pm0.03$
M12	$0.027\pm0.19$	$5.294 \pm 0.79$	$7.56\pm0.26$	$\textbf{5.38} \pm \textbf{1.01}$	$38.3\pm1.88$	$238.7 \pm 1.28$	$0.64\pm0.05$	$0.175\pm0.21$
Mean	$0.030\pm0.14$	$5.596 \pm 1.08$	$\textbf{7.49} \pm \textbf{0.77}$	$\textbf{5.34} \pm \textbf{1.86}$	$\textbf{38.6} \pm \textbf{2.03}$	$310.0\pm3.01$	$0.52\pm0.21$	$\textbf{0.085} \pm \textbf{0.37}$

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(c) (a) (b) Upstream Midstream Downstream Upstream Midstream Upstream (d) (f) 0.8 5 0.029 0.027 Midstream Upstream Downstream Midstream Downstream Midstream Downstream (g) 0.031 0.030 0.028 Midstream Midstream Downstream

Fig. 2 Box plot of  $UV_{254}$ , DOC, and SUVA in the Manas river (a-c) October; (b-f) November; (g-i) December. Upstream: sample site 1-4; midstream: sample site 5-8; downstream: sample site 8-12.

ranging from 7.39 to 7.60, indicating a weakly alkaline environment. It had an aromatic content of 0.030 as measured by UV<sub>254</sub>, a mean DOC of 5.596 mg  $\rm L^{-1}$ , a chloride value of 5.34 mg  $\rm L^{-1}$ , and a conductivity of 38.6 ms m<sup>-1</sup>.

Fig. 2 and 3 shows the DOC content and  $UV_{254}$  values of surface water samples collected from Manas River. In October, the average DOC levels in the upstream, midstream, and downstream of Manas River were 8.212, 8.315, and

 $5.820~{
m mg}~{
m L}^{-1}$ , respectively, and the average values of UV<sub>254</sub> were 0.032, 0.026, and 0.021 AU cm<sup>-1</sup>, respectively. The highest SUVA content was upstream (0.411), while the SUVA content midstream and downstream were 0.318 and 0.370, respectively.

In November, the DOC content downstream of the Manas River was 5.11, which was similar to the value estimated in October, while the DOC levels of upstream and midstream decreased to 4.783 and 4.118, respectively. The average UV<sub>254</sub>

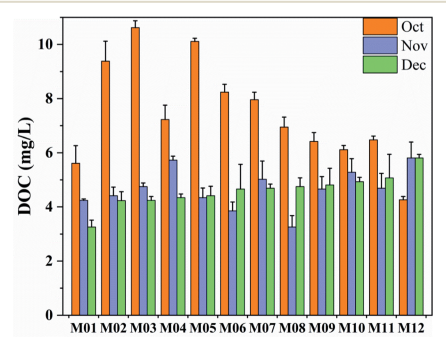


Fig. 3 Seasonal changes of DOC content in the Manas River.

values upstream, midstream, and downstream were 0.028, 0.028, and 0.029 AU cm $^{-1}$ , respectively. In December, the average DOC levels upstream, midstream, and downstream were 4.02, 4.63, and 5.16 mg L $^{-1}$ , respectively, and the average UV $_{254}$  values were 0.027, 0.028, and 0.030 AU cm $^{-1}$ , respectively. Furthermore, in December, the maximum SUVA upstream was 0.68, followed by midstream (0.61) and downstream (0.57).

The average DOC content in Manas River  $(5.596 \text{ mg L}^{-1})$  was higher than that of Mississippi River (3.55 mg  $L^{-1}$ ), Yellow River (2.35 mg  $L^{-1}$ ), and Poyang Lake (3.5 mg  $L^{-1}$ ). 35-37 In terms of spatial trend, the DOC content displayed an expected decreasing trend from upstream to downstream because the downstream regions of the Manas River are mainly rural and agricultural areas, whereas the urban and industrial areas are located in the upstream regions.<sup>38</sup> The values upstream may also be higher owing to the contribution of domestic sewage and industrial wastewater to the DOC content in the upstream, as well as the water flow from the Jiahezi Reservoir upstream. Owing to the low drift and long retention time, the water quality of the reservoir undergoes complex physical, chemical, and biological changes; it usually contains a high DOC content.39 There are a few villages and towns near the middle and lower reaches. Although they contribute to the organic matter content in the Manas River, they have little impact. In terms of temporal changes, the DOC was significantly low in November and December, as compared to October (Fig. 3). This could be because the impact of precipitation on the DOC in the water body is weakened in peak winter, that is, as the river reaches a withered water period, the input of external DOC is greatly reduced.40,41 Meanwhile, as water temperature ranges from 0.9 °C to 6.7 °C in winter, this low temperature further reduces the DOC produced on site. All these results indicate that the Manas River contains a higher amount of DOC, which is the main component of naturally DOM. 42,43 Therefore, the material structure and content of DOM exhibit spatial and temporal differences. In other words, higher DOC levels were observed in October and upstream of the river, and domestic activities frequently affected the DOC content in Manas River.

The SUVA index is an indicator of molecular weight and is used to characterize the aromaticity of DOM.44 It has been reported that an SUVA value less than 3 generally indicates that the majority of DOM consists of more aliphatic compounds, while SUVA greater than 3.5 suggests more complex aromaticity. 41 As shown in Fig. 2(c), (f) and (i), all the measured SUVA values between October and December were less than 3, that is between 0.239 and 0.890. This suggests that the major components of DOM in Manas River were low molecular weight aliphatic compounds.45 It can be seen that as the SUVA decreases from upstream to downstream, the aromaticity decreases. This may be due to the reduction of aromatic components by sediment adsorption or microbial decomposition in the water flow process. It is also evident that the higher frequency of human activities leads to stronger aromaticity of DOM in upstream. The overall SUVA demonstrated a gradual increase from October to December, indicating that the organic matter was more aromatic in winter and could contain more aromatic compounds with unsaturated carbon-carbon bonds. In general, the Manas River contained aliphatic compounds that did not absorb at 254 nm. Furthermore, the aromaticity gradually decreased with the direction of the river (from upstream to downstream), and was stronger in winter.

### 3.2. PARAFAC analysis

PARAFAC analysis was performed on all samples by selecting two to seven components. Half split analysis was also performed to find a suitable number of components. Only the models containing four and five components could be split-half validated. Tucker's congruence coefficients ( $r_c$ ) were applied to identify the similarity between two pairs of Ex and Em loadings. Some of the fluorescent components extracted by PARAFAC indicated the presence of specific organic substances, which could also represent organic compounds with similar fluorescent properties. The EEM measurements were performed at a constant temperature ( $25 \pm 1$  °C) and pH to minimize variation. The maximum fluorescence intensity of each component was obtained to illustrate the quantitative and qualitative differences between the samples.

Two fluorescent components (C1 and C2) were identified from the samples using PARAFAC (Fig. 4, Table 2). Component C1 had Ex/Em maxima at (235, 335)/390 nm, which resembled the traditionally defined peak A (at 260/380-460 nm) and peak M (290-310/370-430 nm), 21,46 assigned to a humic-like component associated with fulvic acids. Component C2 showed Ex/Em maxima at (≤200, 275)/300 nm, which was similar to the tryptophan-like peak T47 observed in farmland environments. Accordingly, C1 represented terrestrial humus, which is primarily observed in surface water. However, it is also found in sewage and agricultural environments; it mainly originates from land-derived and microbial organic matter. 48 C2 is an amino acid that is free or bound to proteins. It is similar to free tryptophan, which is widely present in aquatic environments and is an indication of recent biological activity.24 Thus, a protein-like component may be associated with refractory

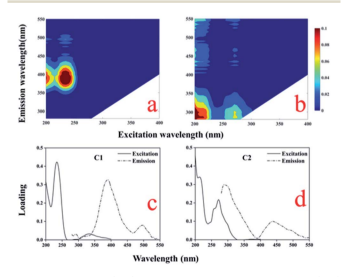


Fig. 4 EEM contours (a, b) and excitation and emission loadings (c, d) of the two fluorescent components identified using PARAFAC.

Table 2 Description of the PARAFAC components and their comparison with previous studies<sup>6</sup>

Component	Ex/Em (nm)	DOM groups and descriptions by previous studies	This study
C1	(235, 335)/390	C2: <250 (320)/390, humic-like materials <sup>26</sup> C1: 240(320)/404, humic-like materials produced from organic matter as a result of microbial activities <sup>50</sup>	Humic-like components
C2	(≤200, 275)/300	C7: 275/340, tryptophan-like components <sup>14</sup> C3: 230(280)/319, tryptophan-like components <sup>38</sup> C4: 260–280/250–340, protein-like components <sup>47</sup>	Protein-like components

<sup>&</sup>lt;sup>a</sup> Ex/Em represents the excitation and emission maxima.

DOM, which is mainly affected by autogenous sources produced by phytoplankton and microbial metabolism in water bodies. Humus components are produced by terrestrial and microbial sources, and are mainly affected by domestic sewage, agricultural irrigation drainage, and microbial degradation. Studies have shown that DOM in the water body primarily contains humic acid, hydrophilic organic acids, proteins, carbohydrates, and lipids.49 Its main source is the sewage from domestic activities, industrial wastewater, and the natural decomposition of dead organisms or the by-products of bacterial decomposition. The city of Shihezi, which lies the northwest region of the Manas River, has airports, factories, and small villages and towns near the middle and lower reaches. Due to factory sewage, domestic sewage, and farmland irrigation discharged into Manas River, as well as the degradation of microorganisms and the self-growth of phytoplankton, terrestrial humus (C1), and protein-like substances (C2) were detected in the river.

The mean fluorescence intensity of the two components and their respective contribution to total DOM fluorescence intensity (the percentage of fluorescence maximum score for each component to the total fluorescence maximum scores of all components) using the PARAFAC model differed by location (Fig. 5). In the entire Manas River, the proportion of humus substances (C1, 54%) was higher than that of protein substances (C2, 46%), which indicated that the humus components had a greater impact on the organic matter content. The data revealed that from upstream to downstream, the humus and protein-like components showed a downward and upward trend, respectively, indicating that the land source gradually decreased along the river direction, while the selfgenerating source gradually increased. However, sampling site M11 had a higher proportion of the humus substances component, which was different from the other downstream sites. This may be due to the influence of man-made inputs or the influence of a small iceberg meltwater runoff at this site. This means that DOM contained more contributions from terrestrial humus upstream of the river, while aquatic systems and microorganisms contributed more downstream. This was consistent with the results of the above.

### 3.3. Fluorescence indices

The ratios among fluorescence peak intensities can also be useful fluorescence indicators, including the fluorescence index (FI), biological source index (BIX), and humification index (HIX). The FI was used to indicate the source of humus components in DOM, where FI < 1.4, indicated land and soil origins, FI > 1.9 reflected a microbial source, and FI between 1.3 and 1.9 demonstrated that the water body was a mixture of terrestrial and self-generating source.50 As shown in Fig. 6, the average FI value (3.26) upstream in October was greater than 1.9, indicating that the DOM mainly consisted of microbes or aquatic resources upstream. However, this may be because there were more plants near the upper reaches, and the decaying of their leaves in autumn probably increased the microbial activities. In the midstream and downstream regions, the FI values were lower, that is 1.62 and 1.41, respectively. Furthermore, the DOM may be affected by microorganisms and terrestrial sources. In contrast, in November, the upstream and midstream FI values were 1.52 and 1.06, respectively, while downstream had a higher FI of 4.91, especially M10. This showed that there was a high external source input, which was

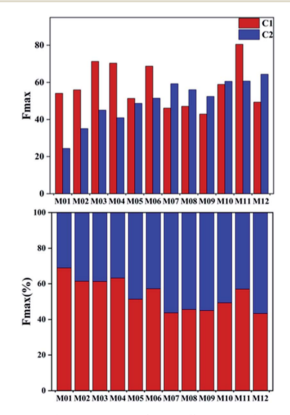


Fig. 5 Parallel factor component (PARAFAC) maximum fluorescence intensity ( $F_{max}$ ) values organized by sampling location.

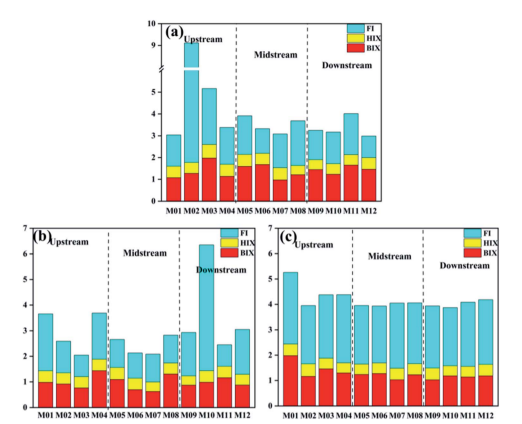


Fig. 6 FI, HIX, and BIX values of Manas River: (a) October; (b) November; (c) December.

consistent with the PARAFAC result. The overall FI value of Manas River in December was relatively high (2.37–3.56) due to the greatly reduced land-source input in winter and the slow flow of the river, which was mainly related to the metabolism of microorganisms in the water body. This indicated that the DOM may not be fully microbiologically derived but could also be the result of terrestrial matter input, as assumed by the ground-water LNE (sample location).<sup>51</sup> Thus, there may be significant differences in the DOM characteristics in different areas of the Manas River. The main sources of DOM in the upper reaches were terrestrial humus and inputs from human activities, followed by the metabolism of microorganisms. This was in stark contrast to the downstream regions. In winter, the river contained higher amounts of DOM.

HIX and BIX were used to infer the degree of humification and the sources of organic matter, respectively. HIX indicates the degree of humification; a high HIX (10–16) suggests a terrestrial origin of DOM and a low HIX (<4) suggests the *in situ* formation of DOM due to biological activities (*i.e.*, a lower humification degree).<sup>52</sup> BIX was used as an indicator of DOM traceability. The BIX values of 0.6–0.7, 0.7–0.8, and 0.8–1.0 indicate weak, intermediate, and strong autochthonous component of DOM, respectively; BIX > 1.0 indicates a biological or aquatic bacterial origin of DOM. The study by Osburn *et al.*<sup>53</sup> found that the trend of BIX values followed that of the FI values. For most groundwater samples, BIX was 0.8–1 and HIX was 4–6, which indicated a weak humic character and strong autochthonous component.<sup>54</sup> In contrast, Manas River had

a higher BIX (1.02–1.47) and lower HIX (<4) values, demonstrating that surface water had a more autochthonous component or aquatic biological DOM. Compared to December, the variations in BIX in October and November were more obvious. This showed that in autumn, the river was affected by inputs from different sources, resulting in obvious changes. The HIX in the Manas River showed a downward trend from upstream to downstream, indicating that even under the influence of external sources, the characteristics of the river's self-generating source increased with the migration and transformation of the river.

Based on the above results, organic matter in Manas River primarily originated from the decomposition of microorganisms in the water and changes in the water body. At the same time, the input of humus and human activities also affected the DOM. External inputs affected upstream, and there was an increase the degradation of microorganisms and bacteria in winter.

### 3.4. Relationship between DBPsFP and DOM characteristics

Natural organic matter reacts with chlorine disinfectants to produce DBPs, which are commonly found in surface water and groundwater. Among them, THMs including chloroform, bromodichloromethane, dibromomonochloromethane, and tribromomethane are the most abundant forms in the chlorination process and pose a threat to human health. According to the regulations of the United States Environmental

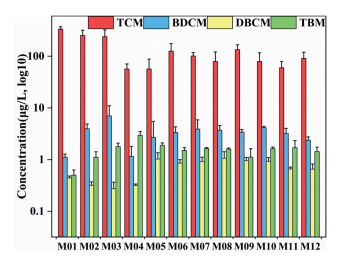


Fig. 7 The formation potential of trihalomethanes in the Manas River. (TCM: trichloromethane; BDCM: bromoxylene; DBCM: dibromomonochloromethane; TBM: tribromomethane).

Protection Agency (US EPA), the highest pollution level of trihalomethane in drinking water is 100 µg L<sup>-1</sup>.54 The formation potential of THMs in the Manas River is shown in Fig. 7. The maximum amount of THMsFP in Manas River reached 225.57 μg  $L^{-1}$ , which included a chloroform FP (TCMFP) of 220.34  $\mu$ g  $L^{-1}$ . At the same time, upstream contained higher content of THMsFP than the middle and lower reaches. Compared to other studies, <sup>29</sup> the measured DBPsFP in raw drinking water was approximately 150  $\mu$ g L<sup>-1</sup>. Ruecker et al. 55 collected water samples from the Waccamaw River (a typical blackwater river in the southeastern United States) and measured the THMsFP to be 71-448  $\mu g L^{-1}$ . Thus, Manas River had a higher THMsFP value, which far exceeded the USEPA pollution standards. This could lead to potential health risks caused by DBPs. However, the method for determining and evaluating DBP precursors through laboratory formation potential is time consuming (chlorination and subsequent extraction and detection takes several days). Therefore, alternative parameters for continuous monitoring of DBP precursors in drinking water treatment plants have been developed, which are essential for optimizing the water treatment process, and effectively reducing and controlling DBPs.

The PARAFAC method is increasingly used in the research on DBPs. The THMsFP, especially TCMFP, which is the most abundant species of THMs, was estimated in most DBP-PARAFAC studies. Some studied reported that the total THMsFP and TCMFP were generally correlated with humic-like components and protein-like components. Further studies on the feasibility of PARAFAC components as surrogates for many emerging DBPs are required to unveil the DOM constituents related to DBP formation and to improve the ability to assess the quantity of DBP precursors in drinking water treatment and distribution systems.

Linear correlations were sought between DBPsFP and DOM properties, such as DOC, SUVA<sub>254</sub>, and  $F_{\text{max}}$  for individual PARAFAC components. There were positive correlations among DOC and PARAFAC humic-like component C1 (r = 0.535, p < 0.5350.01), HIX (r = 0.417, p < 0.01) and C1 + C2 (r = 0.377, p < 0.05), as shown in Table 3. Thus, DOM contained more humus components, which affected the DOC content. There was a significant negative correlation (r = -0.825, p < 0.05) between the SUVA<sub>254</sub> and DOC. Scholz et al.<sup>39</sup> studied a drinking water reservoir in North Wales and found that the SUVA254 had a significant negative correlation (r = -0.581, p < 0.01) with the removal of DOC (the difference between the water inflow and outflow of the reservoir). Yates et al.58 investigated the catchments of two large rivers in Conwy and Nadder and found that DOC had a significant positive correlation with the SUVA (r =0.756, p < 0.01). Although SUVA<sub>254</sub> and DOC have a good correlation, it differs for different water bodies. Therefore, the SUVA<sub>254</sub> index cannot be used as a substitute parameter for DOM. However, in this study, both the SUVA<sub>254</sub> and C1 correctly predicted the DOC content in the Manas River. In addition, the correlation between the fluorescence component was high, which showed the mutual transformation between fluorescence components, meaning that these DOM were from the same organic source. (r = 0.754, p < 0.01). The protein-like components were easily affected by microbial activities, while the humus and protein-like components in the Manas River had a strong correlation. This indicated that microbial activities was the common source of the two components. Moreover, the correlation between THMsFP, HIX, and C2 was weak (r = 0.344, r = 0.316, p < 0.01), but THMsFP had a strong linear correlation

Table 3 Correlation plot of DOC characters and THMsFP

	DOC	SUVA <sub>254</sub>	FI	BIX	HIX	C1	C2	C1 + C2	THMsFP
DOC	1								
SUVA <sub>254</sub>	$-0.825^{a}$	1							
FI	0.079	-0.057	1						
BIX	0.163	$-0.336^{a}$	0.281	1					
HIX	$0.417^{b}$	$-0.385^{b}$	-0.046	0.301	1				
C1	$0.535^{b}$	$-0.516^{b}$	$-0.419^{a}$	0.141	$0.654^{a}$	1			
C2	0.200	-0.201	$-0.636^{b}$	-0.224	0.288	$0.754^{b}$	1		
C1 + C2	$0.377^{a}$	$-0.449^{a}$	$-0.556^{b}$	-0.079	$0.513^{b}$	$0.915^{b}$	$0.911^{b}$	1	
THMsFP	0.203	$0.491^{a}$	-0.009	0.171	$0.344^{a}$	$0.485^{a}$	$0.316^{a}$	$0.529^{a}$	1

<sup>&</sup>lt;sup>a</sup> p < 0.05. <sup>b</sup> p < 0.01.

with other indicators. For example, stronger linear correlations were found between THMsFP and SUVA254 (r = 0.491, p < 0.05), C1 (r = 0.485, p < 0.01), and C1 + C2 (r = 0.529, p < 0.01). THMsFP were correlated strongly with humic-like components than with protein-like components. This result revealed that both humus and protein substances contributed to the formation of trihalomethanes, while humus components contributed more to the formation of THMs. The EEM-PARAFAC approach used in this study was an improvement over DOC and SUVA254. Magbool et al.11 found that the C1/C2 ratio established a direct relationship between seasonal changes and the composition of soil organic matter. They used the ratio of C1/C2 to explore the water quality changes in the treatment units in drinking water plants. In this study, the ratio of the two components of PAR-AFAC was not correlated with THMsFP, while C1 + C2 had a strong correlation with THMsFP (r = 0.529, p < 0.01). This suggested that humic-like fluorophores and protein-like fluorophores were important THMs precursors. Moreover, drinking water plants' exposure to trihalomethanes could be evaluated by changing the main components of organic matter in the water treatment process. The removal effect of the precursors indicates that this index may be useful for optimizing the DBP reduction and control processes.

SUVA, C1 + C2, and THMsFP have a high correlation; however, it varies with season. Thus, it is important to understand which index is more representative. Fig. 8 shows that the correlation between THMsFP and C1 + C2 (r = 0.9828, 0.9205, 0.8265) was higher than that with SUVA (r = 0.6105, 0.7608, 0.6386) from October to December. Throughout the three

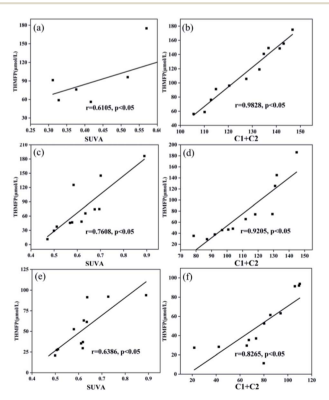


Fig. 8 Correlations between THMsFP and selected DOM properties (a, c, e) THMsFP and SUVA; (b, d, f) THMsFP and C1 + C2 (a, b) October, (c, d) November, and (e, f) December.

months of observation, it was found that the correlation between C1 + C2 and THMsFP was more stable, and the prediction of THMsFP was more accurate than that of SUVA.

As described in Section 3.2 and 3.3, the components of PARAFAC are closely related to the water quality parameters of the Manas River. Therefore, they can predict the origin and composition of DOM and they have the potential to monitor organic matter removal. In this study, the use of PARAFAC analysis proved that the main components of DOM in the Manas River were humus and protein substances. The organic matter in the river was mainly from microbial metabolism and human activities, such as land use, sewage, and wastewater discharge. Thus, the Manas River, a source of drinking water, has a high DOC level, which poses a threat to people's drinking water health. Second, it was proved that the sum of PARAFAC components, C1 + C2, had a strong correlation with the generation potential of DBP (r = 0.529, p < 0.01), thereby predicting the effect of the actual drinking water purification process on the precursors of DBP removal. Finally, the disinfection technology of Xinjiang drinking water plants is relatively traditional and does not involve advanced monitoring equipment; moreover, the determination of DBPs using the present disinfection technology is troublesome and time consuming. The PARAFAC method avoids these shortcomings and it can conveniently and accurately predict disinfection. Changes in the potential of byproduct formation in drinking water plants play an important role in the safety of drinking water for the surrounding residents.

In general, the PARAFAC component was a good substitute for the organic content in the Manas River; the high sensitivity of fluorescence monitoring can detect low-level of organic matter in the water. In addition, the PARAFAC component has high potential to monitor the sub-components of DOC, such as humus and protein.

# 4. Conclusions

DOC, UV254, SUVA, FI, and  $F_{\rm max}$  values of fluorophore components for a diverse group of raw samples from the Manas River provided the following insights:

Due to the surrounding vegetation coverage, the uneven distribution of precipitation throughout the year, the seasonal difference, and the unique hydrological and environmental characteristics of the reservoir during the past three months, the content of organic matter in water changed obviously with the seasons, which was more in October and upstream of the Manas River and less in December and downstream. Meanwhile, the organic matter in the Manas River was dominated by aliphatic compounds. Furthermore, aromaticity gradually decreased along the river and it was higher in winter. Authigenic or aquatic organisms were determined to have a higher DOM and a lower degree of humification. The main sources of DOM were microorganisms and aquatic systems, followed by terrestrial humus and human input. Owing to microbial activity, there are more self-generating sources of DOM downstream, and more external sources upstream, such as domestic

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sewage, industrial wastewater, and surrounding agricultural irrigation discharge.

Moreover, EEM and PARAFAC analysis revealed that the main components of organic matter in the Manas River were terrestrial humus (C1) and tryptophan-like protein (C2); the proportion of humus-like components was higher than that of protein-like substances. The upper reaches of the Manas River contained more humus components. The maximum amount of THMsFP in the Manas River was higher than the standard prescribed by the US EPA. Additionally, the upstream contained higher THMsFP than the midstream and downstream. Meanwhile, the THMsFP and PARAFAC components C1 and C1 + C2 had a significant linear correlation. The results of this study suggest that fluorescence spectroscopy has the potential to assess DBPsFP in situ, which can aid in the management of drinking water quality by providing a theoretical reference for the removal of DBP precursors in the water purification process of water treatment plants.

# Author contributions

Xinlin Wang: validation, conceptualization, investigation, visualization, writing - original draft, writing - review & editing. Yanbin Tong: validation, writing - editing. Jianjiang Lu: validation, resources, ronceptualization, supervision, writing review & editing. Qigang Chang: validation, supervision. Teng Ma: validation, investigation. Fangdong Zhou: validation, investigation. Jiaqi Li: validation, investigation.

# Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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