Effect of pH on binding of pyrene to hydrophobic fractions of dissolved organic matter (DOM) isolated from lake water

Yi Mei¹ · Yingchen Bai² · Liying Wang¹

Abstract In order to better understand the compositional and structural complexity of dissolved organic matter (DOM) macromolecules and provide mechanistic information on the binding of hydrophobic organic contaminants (HOCs) to DOM, we fractionated large amounts of lake water into three hydrophobic DOM-fractions. The variation of the partitioning coefficients ($K_{DOC}$) of pyrene at different pH levels was examined by florescence quenching titration. Results show that, relative to the more polar acidic DOM-fractions, the hydrophobic neutral fraction exhibits a higher sorption ability to pyrene. Generally, the sorption of pyrene to the three hydrophobic fractions is strongly pH-dependent. The $K_{DOC}$ values of pyrene generally increase with decreasing pH levels, which is especially obvious in the sorption of pyrene to the fulvic acid fractions, suggesting that the binding is controlled by hydrophobic interactions. The mechanisms underlying the binding of pyrene to the hydrophobic fractions were also discussed. Our data are beneficial to further understanding the binding of HOCs to DOM and how it has been affected, which may result in more accurate predictions of $K_{DOC}$.

1 Introduction

Dissolved organic matter (DOM) is ubiquitous in the environment and has been reported to influence the fate and behavior of hydrophobic organic contaminants (HOCs) such as polycyclic aromatic hydrocarbon (PAH) (Gauthier et al. 1997; Cox et al. 2007; Perre et al. 2014). Since DOM is a complex mixture composed of aromatic aliphatic hydrocarbon structures, it is arduous to confirm its chemical structure and reactivity. On the other hand, DOM is commonly characterized by fractionating it into secondary categories based on hydrophobic–hydrophilic characteristics with resin sorbents (Leeheer 1981). According to Leenheer's fractionation protocol, DOM can be isolated into hydrophobic fractions and hydrophilic fractions; furthermore, these two fractions can be further fractionated into so-called “acid”, “base” and “neutral” fractions. This fractionation procedure facilitates the understanding of these fractions with different physical–chemical characteristics and provides mechanistic information on the binding of HOCs to DOM (Mei et al. 2009a; Maoz and Chefetz 2010). Ilani et al. (2005) studied the interaction of hydrophobic fractions fractionated from wastewater with triazine herbicide and three PAHs (phenanthrene, fluoranthene and pyrene), and stated that the content of the hydrophobic fractions is more important to the effect of the mobility and transport process of HOCs than to the total organic carbon (TOC) concentration. In our previous studies (Mei et al. 2009a), we investigated the binding of three PAHs (anthracene, phenanthrene and perylene) to three hydrophobic and three hydrophilic fractions, determining the partitioning coefficients ($K_{DOC}$) of the three PAHs between the DOM-fractions and water by fluorescence quenching titration. Our data revealed that the hydrophobic fractions with higher aromaticity and lower
polarity possessed a higher binding ability than that of the hydrophilic fractions. Other investigators obtained similar observations when studying the interaction of HOCs with DOM fractions (Maoz and Chefetz 2010; Xi et al. 2012). All above show the significance of the hydrophobic fractions in the sorption of HOCs.

Since different DOM fractions possess different polarity and acid-base properties, solution chemistry such as pH may have varying effects on the sorption of HOCs to the DOM-fractions. Previous studies showed that the changes of pH lead to DOM conformation change and consequently affected its interaction with hydrophobic organic pollutants (Jones and Tiller 1999; Marschner et al. 2005; Pan et al. 2008; Mei et al. 2009b). However, inconsistent results were reported. With regard to the relation of pH to the $K_{DOC}$ values for HOCs, both positive and negative relationships have been observed (Schlautman and Morgan 1993; Jones and Tiller 1999; Akkanen and Kukkonen 2001; Marschner et al. 2005). In our previous work (Mei et al. 2009b), we studied the effects of pH and ion strength on DOM conformation by 3D fluorescence spectroscopy and fluorescence polarization technique; in this study, we further investigated the effects of pH on the interaction between pyrene and hydrophobic fractions. The objectives of this study are to (1) study the influence of pH on the interactive ability of hydrophobic fractions with pyrene; and (2) correlate the pH and adsorptive properties of the DOM-fractions.

2 Materials and methods

2.1 Regents and chemicals

Pyrene was purchased from Sigma-Aldrich (99.9 % pure, St. Louis, USA) and used without further purification. Concentrated pyrene stock solution was obtained by dissolving the chemical in methanol (HPLC grade, Mallinckrodt, USA). All other chemicals and solvents used are better than analytical grade. Freeze-dried DOM-fractions were dissolved in Milli-Q water (18.2 MΩ cm, Millipore) as concentrated stock, and the solution pH was adjusted by using 0.1 mol/L HCl and 0.1 mol/L NaOH.

2.2 Sample collection and fractionation

Lake water sample was collected from the Hongfeng Lake where is located in Guizhou Province, China. A total volume of 1000 L surface water was collected in August 2010, and the water samples were filtered through a pre-combusted (450 °C for 5 h) Whatman GF/F glass fiber filters for further fractionation. DOM fractions were isolated, based on the literature (Leeheer 1981; Chefetz et al. 1998), using XAD-8/4 resin (Supelco Bellefonte, PA, USA). Detailed fractionation procedures were reported in our previous work (Mei et al. 2009a). Briefly, we adjusted the pH of the filtered original water to 2 with HCl, and it flowed through the XAD-8/XAD-4 resin columns at a rate of 4 L/h. The hydrophobic acids (HOAs), which were composed of humic acid (HA) and fulvic acid (FA), adsorbed onto XAD-8 resin were eluted with 0.1 mol/L NaOH. Then, the HOAs were further acidified to pH 1 with HCl and they settled for 24 h to precipitate HA. The suspension was recycled through the XAD-8 resin and then eluted with 0.1 mol/L NaOH. The elution is the FA fraction. Then the XAD-8 resin was air dried and Soxhlet-extracted with methanol. The fraction contained in the methanol solution is the hydrophobic neutral fraction (HON). All fractions were concentrated by rotary evaporation at 35 °C, and then were purified with 100 Da membrane dialysis and freeze-dried to a low-ash solid form.

2.3 Dissolved organic carbon and molecular weights measurements

The dissolved organic carbon (DOC) of the hydrophobic fractions were determined using a high temperature catalytic method with a TOC/N Ilanalyzer (Elementar, Germany). The relative precision of the DOC analysis was <3 %, as obtained by repeated measurement. Molecular weights were obtained using high performance size exclusion chromatography (HPSEC,Agilent 1100, PE) with a UV detector at 254 nm and an YMC-60 column (Waters, Milford, MA). The flow rate of the mobile phase composed of the phosphate buffer (0.001 mol/L Na$_2$HPO$_4$, 0.001 mol/L NaH$_2$PO$_4$ and 0.03 mol/L NaCl) was set at 0.5 ml/min. The DOM-fractions were dissolved in the phosphate buffer with the same composition as the HPSEC mobile phase. The solutions were then filtered through a per-combusted Whatman GF/F glass fiber filter. The number ($M_n$), weight-averaged ($M_w$) molecular weights, and polydispersity ($\rho$) were determined based on the method proposed by Chin et al. (1994).

2.4 Absorbance and fluorescence measurements

Absorbance was obtained using a Shimadzu UV-3000 double Beam spectrophotometer with 1 cm quartz cell to determine the molar absorptivity at 280 nm ($\varepsilon_{280}$) and to correct the inner filter effects (IFE) when conducting a fluorescence quenching experiment. Fluorescence intensities were collected on a Hitachi F-4600 spectrofluorometer (Hitachi, Japan) with a cuvette magnetic stirring system.
containing a Teflon micro stir bar. The photomultiplier voltage was set at 700 V; excitation and emission bandwidth were set at 5 and 10 nm, respectively. The fluorescence intensities of pyrene were correspondingly determined at the excitation/emission wavelengths of 271/374 nm. The $K_{DOC}$ values were determined by a fluorescence quenching titration based on the Stern–Volmer equation (Gauthier et al. 1997). The fluorescence quenching of the PAHs by the DOM have proven to be static quenching mechanisms based on diffusion, temperature studies and the fluorescence efficiency experiment (Tranina et al. 1990; Chen et al. 1994). Five dilutions of the fractions were prepared, after adjustment, to the corresponding pH with 0.1 mol/L HCl and 0.1 mol/L NaOH; then, all samples were bubbled for 5 min with pure Argon in the dark to reach anoxic conditions. The Raman peak of distilled water was collected at 348/397 nm to check the instrument stability during the experiment. A 3-mL aliquot of each fraction of dilution was pipetted into the cuvette. Then, an absorption scan from 250 to 470 nm was conducted. The cuvette was then moved to the Hitachi F-4600 to collect the background fluorescence of the fractions, which was subtracted from the spectra of next solution. An aliquot of pyrene stock solution was spiked into the cuvette to a final concentration of 2 μg, according to its solubility and fluorescence intensity; then, the cuvette was stirred for 3 min using a Teflon micro stir bar and allowed to settle for another 2 min before the fluorescence measurement. After initial spiking, the fluorescence intensity of all samples was collected for three times at 5, 7, and 9 min, and the final data was an average of the three measurements.

All the fluorescence intensity values were corrected for primary and secondary IFE based on the following equation (Gauthier et al. 1997):

$$F_{cor}/F_{obsd} = \frac{2.3dA_{ex}}{1 - 10^{-gA_{em}}} \frac{10^{-sA_{em}}}{1 - 10^{-sA_{em}}}$$

(1)

where $F_{cor}$ is the corrected fluorescence intensity, $F_{obsd}$ is the observed fluorescence intensity, $A_{ex}$ is the absorbance at the excitation wavelength, $A_{em}$ is the absorbance at the emission wavelength, and $d$, $g$, and $s$ are the fluorescence beam width (1 cm), distance from the edge of the sample beam to the edge of the cuvette (0.3 cm), and excitation beam thickness (0.2 cm), respectively. The binding of pyrene to the fractions was described by the Stern–Volmer equation (Gauthier et al. 1997):

$$F_0/F = 1 + K_{DOC}[DOM]$$

(2)

where $F_0$ and $F$ are the fluorescence intensity of pyrene in the absence and presence of DOM-fractions, respectively, and $K_{DOC}$ and [DOM] are the partitioning coefficients of pyrene and the concentration of the DOM-fractions, respectively.

### 3 Results and discussion

#### 3.1 DOM-fractions characterization

DOM is composed of a mixture of low-molecular weight compounds and chemically heterogeneous macromolecules (Leenheer and Croué 2003). Fractionation of the DOM is necessary to better understand the contribution of the individual fraction to its total sorption properties.

Table 1 shows the results of elemental analysis of the hydrophobic fractions. The lake-derived water in this study contained 77 % hydrophobic fractions (based on DOC analysis), which is consistent with the results obtained by other investigators (Polubesova et al. 2007). Elemental analysis exhibited that the hydrophobic fractions possess a lower C/H ratio, suggesting their distinct aliphatic property. Relative to the HA and FA fraction, the HON fraction contains a high content of carbon and lower content of oxygen. With respect to atomic ratios, HON demonstrates a lower C/H ratio and a high C/O ratio, indicating its lower polarity and higher hydrophobicity relative to the acidic fractions (i.e., HA and FA). Since the HA and FA fraction are characterized by a higher oxygen content and lower C/O ratio, and the carboxyl group may be the dominant polar functional group in the acidic fractions.

#### 3.2 Partitioning coefficients of the hydrophobic fractions

The Stern–Volmer plots for the sorption of pyrene to FA, HA and HON were curved towards the y-axis and not all plots were linear (Fig. 1). Nonlinear sorption isotherms of pyrene binding to the DOM have been reported previously (Laor and Rebhun 2002; Borisover et al. 2006; Polubesova et al. 2007). It is well known that the binding of HOC to aquatic DOM is usually ascribed to a non-specific partitioning mechanism, and the sorption isotherm is linear in this case. Laor and Rebhun (2002) argued that the non-linearity of pyrene isotherms resulted from the combination of non-specific (partitioning) and specific (adsorption) binding mechanisms. Since the three fractions contain both polar and non-polar functional groups, the polar interactions, such as the π–π electron donor–acceptor interaction (EDA) and H-bonding, may greatly contribute to the non-linearity of sorption isotherms.

Table 2 shows the log$K_{DOC}$ values of pyrene binding to the three hydrophobic fractions at pH 7. Our data are well consistent with the reported results (Wijnja et al. 2004; Marschner et al. 2005; Polubesova et al. 2007). Increasingly, the sorption ability of HON for pyrene was higher than that of HA and FA, with log$K_{DOC}$ values of 4.61, 4.49, and 4.25 for HON, HA and FA, respectively. It is well
known that the binding of neutral PAHs molecules to DOM is governed by hydrophobic interactions, which is the combination of dipole-induced dipole interactions (i.e., dispersion forces) and the thermodynamic gradient driving force. The magnitude of inducing a dipole moment is related to the polarizability of the molecules, which can be estimated by the sum of their bond polarizability. Therefore, an increase in the number of conjugated double bonds (–HC=CH–) will increase the molecular polarizability and thereby increase the magnitude of the dispersion forces. As a result, an increase in the aromaticity of DOM will increase the polarizability of the molecules, and accordingly increase its binding ability to the PAHs (Gauthier et al. 1997; Peuravuori 2001). The atomic C/H ratios, molecular weights of DOM and UV absorbance at 280 nm ($\varepsilon_{280}$), were thought to be surrogate for the aromaticity of DOM (Gauthier et al. 1997; Tanaka et al. 1997; Peuravuori 2001). However, our data was somewhat inconsistent with the aforementioned conclusions. Although the HON fraction possesses a lower atomic C/H ratio, its molecular weight and $\varepsilon_{280}$ are lower than those of the HA fraction. Besides higher aromaticity (lower C/H ratio), the higher sorption ability of HON to pyrene may result from its lower polarity. Xing et al. (1994) proposed a polarity index (PI) to represent the magnitude of DOM polarity, and defined FI as the atomic (O + N)/C ratio. Although the HA fraction possesses a higher molecular weight and $\varepsilon_{280}$ value, a higher content of polar compounds in HA impede the sorption more pyrene. Our data was well corroborated by the reported observations (Salloum et al. 2002; Simpson et al. 2003; Gunasekara et al. 2003). Salloum et al. (2002) reported an increase in $K_{DOC}$ values after removing the polysaccharides of soil organic matter, and speculated that aromatic structures might be physically constrained by polar structures. Likewise, when carbohydrates were selectively removed from humic acids, Simpson et al. (2003) discovered an increase in phenanthrene binding ability. After the similar observations were obtained, Gunasekara et al. (2003) proposed that the mobile sorption domains were free and would become more accessible to HOCs with the removal of rigid structures. Another possible contribution to the higher binding ability of HON may be the EDA interaction. The HON fraction was reported to be rich in methyl and carbonyl groups, which could accept electrons from electron-donor pyrene.

### 3.3 Effect of pH on the binding characteristics

Table 3 shows the $K_{DOC}$ values of pyrene binding to HA, FA and HON at pH 4, 7, and 10. The $K_{DOC}$ values of pyrene were higher at pH 4 and lower at pH 10, suggesting the strong pH-dependent sorption property. Since the DOM-fractions contain large amounts of polar and acidic function groups, any changes in pH may affect its

### Table 1 Elemental analysis and atomic ratios of the hydrophobic fractions (HA, FA and HON)

<table>
<thead>
<tr>
<th>Sample</th>
<th>Elemental analysis (%)</th>
<th>Atomic ratios</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C</td>
<td>H</td>
</tr>
<tr>
<td>HA</td>
<td>43.38</td>
<td>4.45</td>
</tr>
<tr>
<td>FA</td>
<td>35.36</td>
<td>3.68</td>
</tr>
<tr>
<td>HON</td>
<td>52.22</td>
<td>7.64</td>
</tr>
</tbody>
</table>

### Table 2 Log $K_{DOC}$ values of pyrene with hydrophobic fractions, molecular weight and UV at 280 nm ($\varepsilon_{280}$)

<table>
<thead>
<tr>
<th>Sample</th>
<th>$M_w$</th>
<th>$M_n$</th>
<th>Polydispersity</th>
<th>$\varepsilon_{280}$</th>
<th>Log $K_{DOC}$a</th>
</tr>
</thead>
<tbody>
<tr>
<td>HA</td>
<td>2355</td>
<td>1928</td>
<td>1.22</td>
<td>190</td>
<td>4.49 ± 0.03</td>
</tr>
<tr>
<td>FA</td>
<td>2045</td>
<td>1471</td>
<td>1.39</td>
<td>143</td>
<td>4.25 ± 0.04</td>
</tr>
<tr>
<td>HON</td>
<td>2100</td>
<td>1338</td>
<td>1.57</td>
<td>175</td>
<td>4.61 ± 0.01</td>
</tr>
</tbody>
</table>

* a data are expressed as mean value and a standard deviation.
Table 3 Partitioning coefficients ($K_{DOC}$) of pyrene binding to HA, FA and HON at pH 4, 7 and 10

<table>
<thead>
<tr>
<th>Sample</th>
<th>$K_{DOC}$ ($10^3$ L/kg C)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>pH 4</td>
</tr>
<tr>
<td>HA</td>
<td>48.5 ± 4.45</td>
</tr>
<tr>
<td>FA</td>
<td>32.05 ± 4.30</td>
</tr>
<tr>
<td>HON</td>
<td>52.30 ± 3.87</td>
</tr>
</tbody>
</table>

* Mean ± standard deviation

conformation and sorption domains, depending on its physicochemical properties and composition.

The carboxyl groups of DOM were reported to be increasingly deprotonated above pH 4, resulting in an increase of its polarity and solubility (Carter and Suffet 1982; Marschner et al. 2005). Furthermore, the conformation of DOM could be affected by the change of solution pH (Jones and Tiller 1999; Ferreira et al. 2002). Humic substances are in an elongated conformation in alkaline solutions due to electrostatic repulsion, but the protonation of carboxyl groups at low pHs facilitates intramolecular interactions and leads to the formation of a more condensed hydrophobic structure. Pan et al. (2008) observed an aggregation of DOM molecules via atomic force microscopy and an increase in zeta potential, with decreasing pH. They suggested that, in this case, electrostatic repulsive force decrease, and the aggregation of DOM molecules formed large hydrophobic domains in which more hydrophobic compounds were adsorbed, as if individual molecules possessed lower apparent molecular sizes due to self-curling. In our previous study (Mei et al. 2009b), we studied the effects of pH on the florescence properties and molecular conformation of two humic acids by the three-dimensional excitation–emission matrix florescence (3DEEM) and steady-state florescence polarization (FP). As pH decreased, we observed a decline in the florescence intensity and a blue shift in the florescence maxima. In the case of the blue shift in florescence maxima, the polarity of humic substances declined and the hydrophobic domains of the DOM functional groups were exposed in the solution (Mobed et al. 1996). At the same time, decreasing florescence intensities suppressed the ionization of the functional groups. The conformational change of the humic acids was corroborated by florescence polarization: the FP decreased with decreasing pH. This indicated the decreasing of the apparent molecular size, because of the self-curling of individual molecule. In addition to enhancing the hydrophobic interaction resulted from molecular aggregation, the lower pH is beneficial to polar interactions such as H-bonding and EDA interactions. Gu et al. (2007) observed the H-bonding ability of carboxyl groups of DOM declining due to deprotonation with increasing pH. Zhu et al. (2004) stated that lower pH would induce a stronger π–π EDA interaction. Furthermore, hydrogen ions could also act as the promoters for EDA interactions via aromatic moieties or heterocyclic compounds within the DOM molecules.

It is clear that a greater decrease in the $K_{DOC}$ values of pyrene binding to FA occurs with increasing pH (Table 3). With regard to the HON fraction, only a slight effect was observed with changing pH from 4 to 10, while the influence of changing pH on the pyrene sorption to HA was lower than that to FA and higher that to HON. The different extent of pH-dependent binding observed among the three DOM-fractions might be ascribed to their respective compositional and structural properties. Relative to the HON fraction, the FA fraction has a much higher polarity, as shown by the (O + N/C) ratio (Table 1), and is more enriched in polar groups. Accordingly, the distinct components of FA become tightly bound via intramolecular H-bonding at lower pHs, leading to more sensitive pH-dependent binding. While HON is less polar with a PI = 0.6 and is composed of highly cross-linked, large-sized hydrophobic domains, its conformation and variation of hydrophobicity would be less affected by the protonation–deprotonation transition of carboxylic groups.

4 Conclusions

Our data reveals that the sorption of pyrene to lake-derived hydrophobic fractions is strongly correlated to the aromatic domains in DOM macromolecules. Relative to the more polar acidic DOM-fractions, the hydrophobic neutral fraction (HON) exhibits a more efficient sorption ability to pyrene. Generally, the sorption of pyrene to the three hydrophobic fractions is strongly pH-dependent. The partitioning coefficients ($K_{DOC}$) of pyrene generally increase with decreasing pH, which is especially obvious with the sorption of pyrene to the FA fraction, suggesting the binding is controlled by hydrophobic interactions. Among the three hydrophobic fractions, HON exhibited the highest sorption ability to pyrene, which may result from its lower polarity and stronger EDA ability. The effect of pH on the sorption of pyrene to HON is relatively limited due to its less polar groups; furthermore, HON possesses larger-sized hydrophobic domains whose conformation and hydrophobicity would not be much affected by the protonation–deprotonation transition of the carboxylic groups. As the carboxyl groups of DOM are protonated with decreasing pH (i.e., increasing H$^+$), the electrostatic repulsion between the DOM molecules decrease and the molecular aggregation happen; furthermore, the aggregation results in the formation of larger hydrophobic domains in which more
HOC are adsorbed, especially those with large hydrophobicity. Our data are beneficial to further understanding the binding of HOCs to DOM and how it has been affected, which may lead to a more accurate prediction of $K_{DOC}$.

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References