Adsorption of polyhydroxy fullerene on polyethyleneimine-modified montmorillonite

Qingze Chen a,b, Runliang Zhu a,b, Yanping Zhu a,b, Jing Liu a,b, Lifang Zhu c, Lingya Ma a,b, Meng Chen a

Abstract

The environmental behaviors and potential ecotoxicity of carbon nanomaterials, such as fullerene and its derivatives, are gaining ever-increasing concerns at present. This work attempts to develop an adsorbent for the effective removal of polyhydroxy fullerene (PHF) from aqueous solution, which was synthesized by modifying montmorillonite (Mt) with a branched polymer polyethyleneimine (PEI). The adsorption results showed that the obtained adsorbent (i.e., PEI-Mt) could effectively remove PHF over a wide range of solution pH; both the electrostatic attraction and hydrogen-bond interaction between PHF and PEI-Mt contributed to the strong adsorption. Decreasing solution pH and rising PEI loading amount on Mt could both increase the adsorption amounts of PHF on PEI-Mt. The adsorption isotherms could be well fitted with the Langmuir model, and the calculated maximum adsorption value of PHF on 10%PEI-Mt reached ~213 mg/g, much higher than that on the original Mt (~16 mg/g). The adsorbents after PHF adsorption were further characterized using Fourier transform infrared spectroscopy, X-ray diffraction, and transmission electron microscopy. The results suggested that the adsorbed PHF primarily existed on the outer surfaces of PEI-Mt. This work showed that PEI-Mt can be a potentially efficient adsorbent for the removal of PHF from aqueous solution.

1. Introduction

Fullerene and its derivatives have gained immense attention so far due to their remarkable properties and potential applications in many areas, including environmental remediation (e.g., adsorbents and catalysts) (Yang et al., 2006; Meng et al., 2012), biomedicine (e.g., antibacterial and antitumor) (Markovic and Trajkovic, 2008; Kwag et al., 2013), and materials science (e.g., sensors, supercapacitors, and hydrogen storage) (Li et al., 2009; Xiong et al., 2012; Rather and De Wael, 2013). With the ever-increasing production and wide applications of fullerene-based carbon nanomaterials, they will inevitably enter into the environment (Colvin, 2003; Klaine et al., 2008). As such, significant concerns have been drawn to their influences on human health and natural environment.

Although fullerene is highly hydrophobic, water-soluble fullerenes, such as aqueous fullerene colloid aggregates (nC60) and polyhydroxy fullerene (PHF), can be conveniently synthesized by ultrasonication, solvent exchange, and chemical modification (Brant et al., 2006; Georgieva et al., 2013). The enhanced water-solubility expands their applications, but may simultaneously increase their risks to the environment and human health (Klaine et al., 2008). Indeed, the cytotoxicity of water-soluble fullerenes to both bacteria and human cell has been confirmed by many studies (Sayes et al., 2004; Wielgus et al., 2010; An and Jin, 2012). Consequently, proactive research on effective removal of water-soluble fullerenes from aqueous solution is critical to controlling the potential risks and ensuring a sustainable carbon nanomaterials industry.

At present, a number of studies have been conducted to remove nC60 from wastewater by various adsorbents, such as biomass (Kiser et al., 2010), activated sludge (Yang et al., 2013), and alum (Wang et al., 2013). In contrast, much less work about the pollution control of PHF has been reported up to now. On the other hand, Prylutskyy et al. (2014) indicated that the origin of fullerene solubility in water was probably based on the surface hydroxylation of fullerene molecules; therefore, nC60 could be considered as fullerene aggregates with a certain amount of hydroxyls on the outer surfaces. Furthermore, when exposed to the environment, fullerenes and nC60 could be transformed into PHF by many factors (e.g., long-term mixing, ultraviolet radiation, and interaction with microbes and oxidants) (Hwang and Li, 2010; Qu et al., 2010; Chae et al., 2014). Hence, being regarded as the main products of fullerenes in the environment, PHF need large concerns in terms of their pollution control.
Table 1
Meanings of the abbreviations for the samples.

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Meaning</th>
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<tr>
<td>PHF</td>
<td>Polyhydroxy fullerene</td>
</tr>
<tr>
<td>PEI</td>
<td>Polyethylenimine</td>
</tr>
<tr>
<td>Mt</td>
<td>Montmorillonite</td>
</tr>
<tr>
<td>2%PEI-Mt</td>
<td>2%PEI-modified Mt</td>
</tr>
<tr>
<td>10%PEI-Mt</td>
<td>10%PEI-modified Mt</td>
</tr>
<tr>
<td>Mt-PHF</td>
<td>Mt after the adsorption of PHF (C_0 = 400 mg/L)</td>
</tr>
<tr>
<td>2%PEI-Mt-PHF</td>
<td>2%PEI-Mt after the adsorption of PHF (C_0 = 400 mg/L)</td>
</tr>
<tr>
<td>10%PEI-Mt-PHF</td>
<td>10%PEI-Mt after the adsorption of PHF (C_0 = 400 mg/L)</td>
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</table>

With 2D nano-sized structure, high cation exchange capacity, and large specific surface area, montmorillonite (Mt) as a low-cost and environmentally-friendly adsorbent for cationic contaminants (e.g., cationic dyes and heavy metal cations) has been widely applied in environmental remediation (Bergaya and Lagaly, 2013; Yuan et al., 2013; Zhu et al., 2016). However, owing to the negatively charged layers, raw Mt exhibits a poor affinity toward anionic contaminants (Zhu et al., 2016). A number of studies indicated that Mt could be easily modified by cationic modifiers to synthesize various adsorbents, such as the cationic surfactant modified Mt for organic contaminants (Roberts et al., 2006; Zhu et al., 2011), and the hydroxymetal polycations modified Mt for oxyanions (Zhou et al., 2010; Ma et al., 2015). As an important type of modifiers for Mt, the cationic polymers have received broad attentions because of their large charge/mass ratio (Yue et al., 2007; Zhu et al., 2010). In this regard, cationic polymers could readily saturate the CEC of Mt and change the surface charge of Mt from negative into positive. As a result, the modified Mt could provide adsorption sites for anionic contaminants (Kang et al., 2009; Li et al., 2011). Given that PHF generally exhibit negative surface charges in water (Brant et al., 2007), one could propose that cationic polymers-modified Mt may be an efficient adsorbent for PHF.

Herein, polyethylenimine (PEI), a water-soluble cationic polymer with a large number of amino groups and high charge density (Fig. S1) (Zhang et al., 2014), was selected to modify Mt. The adsorption capacities of Mt and the PEI-modified Mt (PEI-Mt) toward PHF were investigated, and the potential adsorption mechanisms were further elucidated through X-ray diffraction (XRD), transmission electron microscopy (TEM), and Fourier transform infrared spectroscopy (FTIR). Our results indicated that PEI-Mt exhibited high affinity toward PHF, with the maximum adsorption amount of ~213 mg/g. As such, PEI-Mt could be used as a promising adsorbent toward PHF.

2. Materials and methods

2.1. Materials

The sodium-exchanged Mt used in this work was prepared by dispersing the calcium-rich Mt (CEC was 110.5 mmol/100 g, purity >95%, Inner Mongolia, China) in a 0.5 mol/L solution of NaCl, followed by stirring for 24 h at 80 °C (repeated 3 times). The obtained suspension was centrifuged and washed repeatedly with distilled water until AgNO3 test confirmed no chloride in the supernatant. After being dried at 60 °C, the sodium-exchanged Mt was collected for the follow-up experiments. For brevity, Mt was used to denote the sodium-exchanged Mt in the following text. The cationic polymer, branched PEI (molecular mass = 10,000, 99%), was supplied by Shanghai Aladdin Bio-Chem Technology Co., Ltd. PHF was purchased from Suzhou Dade Carbon Nanotechnology Co., Ltd.

2.2. Synthesis of adsorbents

PEI-Mt was prepared as follows: a desired amount of PEI was dissolved into distilled water under stirring for 10 min, followed by adding 10 g of Mt. The mixture was vigorously stirred for 12 h at room temperature, and then the resulting OMts were collected by centrifugation, washed with distilled water (3 times), and freeze-dried for 24 h. The added amounts of PEI were 2% and 10% of the weight of Mt, and the corresponding OMts were denoted as 2%PEI-Mt and 10%PEI-Mt, respectively. For clarity, the meanings of all the abbreviations for the samples are listed in Table 1.

2.3. Adsorption experiments

In order to determine the adsorption equilibration time, the adsorption kinetics experiments were conducted with the adsorption time ranged from 10 min to 24 h. In detail, 0.1 g of the adsorbents (i.e., Mt and 2%PEI-Mt) were added into 20 mL of the PHF solution with the concentration of 200 mg/L. The suspension were steadily agitated in an incubator shaker at 200 rpm and 25 °C. Subsequently, the influences of pH on the adsorption of PHF on PEI-Mt were investigated. In this case, the pH was initially adjusted using 0.1 M NaOH or 0.1 M HCl, and the initial concentration of PHF for 2%PEI-Mt and 10%PEI-Mt were 200 and 800 mg/L, respectively. Besides, batch adsorption experiments were conducted to evaluate the adsorption capacities of the raw Mt and PEI-Mt toward PHF from aqueous solutions. Specifically, 0.1 g of the adsorbents were mixed with 20 mL of PHF solutions containing different

Fig. 1. Effects of pH on the adsorption capacities of 2%PEI-Mt and 10%PEI-Mt toward PHF (a). Zeta potential of Mt, 2%PEI-Mt, and 10%PEI-Mt as a function of solution pH (b).
3. Results and discussion

The adsorption kinetics of PHF on Mt and 2%PEI-Mt were first investigated (Fig. S2). Both Mt and 2%PEI-Mt exhibited rapid adsorption rates toward PHF within the first 2 h, which could be ascribed to the abundant adsorption sites at the beginning. According to the adsorption kinetics results, the adsorption time was set at 24 h in the follow-up batch adsorption experiments.

Owing to a large number of primary and secondary amine groups on the macromolecular chains, the surface properties of PEI are highly dependent on solution pH (Amara and Kerdjoudj, 2003; Gao et al., 2009). As such, the effects of pH on the adsorption capacities of PEI-Mt toward PHF were investigated (Fig. 1a). The adsorption of PHF on both 2%PEI-Mt and 10%PEI-Mt decreased as the solution pH increased from 4 to 12. Given that about 70% of the primary and secondary amine groups in PEI molecular were protonated in acidic solution (Amara and Kerdjoudj, 2003; Gao et al., 2009), the increase in PHF adsorption could be mainly attributed to the enhanced protonation of amine groups, which could facilitate the electrostatic interaction between negatively charged PHF and positively charged amine groups in PEI molecular. Moreover, 10%PEI-Mt showed much better adsorption capacity than 2%PEI-Mt toward PHF, suggesting that increasing PEI loading amount on Mt could provide PEI-Mt more adsorption sites for PHF. This hypothesis could be further confirmed by the change in zeta potentials of the adsorbents (as shown below).

The zeta potential of Mt was affected evidently by the modification of PEI (Fig. 1b). Specifically, the isoelectric point (pHzpc) of Mt could not be observed in the pH range of 3–12. The modification by PEI caused the appearance of the pHzpc of 10%PEI-Mt at pH 9.6, which could be attributed to the protonation of amine groups on the surface. Similar phenomena were reported in previous studies using PEI to modify other substances, such as silicon carbide (Zhu et al., 2003), nano-zirconia powders (Tang et al., 2000), and hematite (Chibowski et al., 2009). When the solution pH was below pHzpc, the positively charged 10%PEI-Mt may effectively adsorb PHF through electrostatic interaction. Noticeably, the surface of 2%PEI-Mt was negatively-changed in the whole pH range, and its zeta potentials only positively shifted as compared with that of Mt, demonstrating that a small amount of PEI could not saturate the CEC of Mt and alter its surface charge. On the other hand, the adsorption amount of PHF was still relatively large when the solution pH was above the pHzpc of 10%PEI-Mt, although the electrostatic interaction between PHF and PEI became weak because of the formation of the neutral PEI-Mt composites. Several previous studies disclosed that a great number of amino groups on PEI could generate strong attraction with other organic compounds through hydrogen bonding (Chibowski et al., 2009; Wang et al., 2015). As such, one might propose that the hydrogen bonding between PEI and PHF could contribute to the high adsorption capacity at high pH values.

<table>
<thead>
<tr>
<th>Adsorbent</th>
<th>Q0 (mg/g)</th>
<th>b (L/mg)</th>
<th>R²</th>
<th>Kf</th>
<th>n</th>
<th>R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mt</td>
<td>16.42</td>
<td>0.037</td>
<td>0.9962</td>
<td>2.10</td>
<td>0.828</td>
<td>0.8928</td>
</tr>
<tr>
<td>2%PEI-Mt</td>
<td>86.21</td>
<td>0.013</td>
<td>0.9941</td>
<td>1.41</td>
<td>1.157</td>
<td>0.9937</td>
</tr>
<tr>
<td>10%PEI-Mt</td>
<td>212.77</td>
<td>0.026</td>
<td>0.9986</td>
<td>2.39</td>
<td>5.528</td>
<td>0.9609</td>
</tr>
</tbody>
</table>

Table 2: Langmuir and Freundlich equation parameters for the adsorption isotherms of PHF on various adsorbents.
The adsorption isotherms were obtained to assess the adsorption capacities of the adsorbents toward PHF (Fig. 2a). The adsorption capacities of PHF on PEI-Mt were much higher than that on Mt. The electrostatic repulsion between the negatively charged Mt and PHF should be the reason for the poor adsorption of PHF on Mt (~16 mg/g). The adsorption amounts may be ascribed to some special interactions, such as hydrogen bonding between PHF and terminal hydroxyl groups of Mt or 'cation-bridging' through the exchangeable cations (Theng, 1982; Fortner et al., 2012). By contrast, PEI-Mt showed strong interaction with PHF, and the maximum adsorption amount of PHF on 10%PEI-Mt could reach ~210 mg/g (Fig. 2b). Combined with the results of zeta potentials above, the protonation of amine groups endowed the surface of 10%PEI-Mt with a number of positive charges under the adsorption experiment condition (pH = 6). Thereby, the electrostatic interaction between the adsorbents and the adsorbates were enhanced.

Langmuir and Freundlich isotherm models were used to quantitatively analyze the adsorption isotherms (Table 2). As shown by the $R^2$ values, the adsorption isotherms of PHF on three adsorbents were better fitted with the Langmuir model, suggesting the monolayer adsorption on the adsorbents with identical adsorption sites. In light of the calculated $Q_m$ values (Table 2), the adsorption capacities of PHF on the adsorbents increased in the order Mt < 2%PEI-Mt < 10%PEI-Mt.

As a control experiment, the PHF removal efficiency from aqueous solution by PEI (i.e., in the absence of Mt) was also studied under the same adsorption conditions. As shown by the photographs of the PHF solution and the supernatants after the adsorption experiments, the PHF solution was dark brown (Fig. 3a). The treatment of PHF by PEI hardly changed the color of the supernatants (Fig. 3b and c). In effect, no visible precipitates were obtained after the centrifugal separation.
of the mixture of PHF and PEI. These results showed that the PEI alone
was not effective in removing PHF from water. In contrast, the superna-
tants became light brown and even completely transparent after the ad-
dition of 2%PEI-Mt and 10%PEI-Mt, respectively (Fig. 3d and e). The
distinctly different results from the visual examination further mani-
fested that PEI-Mt exhibited outstanding adsorption capacity toward
PHF, and can be desirable for the treatment of the PHF wastewater.

The FTIR spectra of the samples were recorded to study the structur-
al changes of the functional groups during the adsorption processes (Fig.
4a). The absorption bands at 3623 and 3428 cm$^{-1}$ on the spectrum of
Mt could be assigned to the structural OH and water molecule
stretching vibrations, respectively (He et al., 2004, 2006b; Chen et al.,
2014). The spectrum hardly altered for Mt after the adsorption of PHF
(i.e., Mt-PHF), which may be due to a very small adsorption amount of
PHF on Mt. As for PEI-Mt, the intensity of these absorption bands de-
creased, implying the reduction of their concentration resulting from
the diluting effect by the loaded PEI. Moreover, the two absorption
bands at ~2925 and 2854 cm$^{-1}$ (arising from the asymmetric and sym-
metric stretching vibrations of CH$_2$ groups, respectively) on PEI-Mt fur-
ther con

Fig. 6. TEM images of Mt(a), Mt/PHF(b), 10%PEI-Mt(c), and 10%PEI-Mt/PHF(d).
PHF significantly clustered on the surfaces of Mt. The reason might be that the special interactions (e.g., the limited terminal hydroxyl groups and ‘cation-bridging’) could not provide enough adsorption sites for PHF, so that PHF tended to cluster by itself on the surfaces. Compared to that of Mt/PHF, a uniform morphology was obtained for 10%PEI-Mt/PHF (Fig. 6d), which could be ascribed to the even distribution of the adsorbed PHF on the adsorption sites of the surfaces, i.e., without forming PHF aggregates.

The above results clearly showed that PEI-Mt had favorable adsorption capacities toward PHF and can be used as a low-cost and high-efficient adsorbent in the removal of PHF from aqueous solution. Meanwhile, given that Mt is ubiquitous in soils and sediments and may form various composites with natural organic macromolecules (Bergaya and Lagaly, 2013), the present work may provide novel information for understanding the migration and sequestration of carbon nanomaterials in the environment. In other words, PHF may be well immobilized by natural clay-macromolecules composites.

4. Conclusion

In summary, the modification of Mt with a branched polymer PEI developed a novel adsorbent, PEI-Mt, for the remediation of the wastewater containing PHF. The adsorption experiments demonstrated that PEI-Mt showed superior adsorption capacity for PHF over a wide variety of solution pH range, which could be attributed to the combined contributions of the electrostatic attraction and the hydrogen-bond interaction between PHF and PEI-Mt. The adsorption amounts of PHF on PEI-Mt increased with rising PHF loading amount on Mt and with decreasing solution pH. The adsorption equilibrium data of PHF on all the absorbents increased with rising PEI loading amount on Mt and with decreasing solution pH range, which could be attributed to the combined contributions of the electrostatic attraction and the hydrogen-bond interaction between PHF and PEI-Mt. The adsorption amounts of PHF on PEI-Mt increased with rising PHF loading amount on Mt and with decreasing solution pH. 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