Nanocomposites of Fe₂O₃@rGO for adsorptive removal of arsanilic acid **from aqueous solution**

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AbstractArsanilic acid (ASA), an organic-arsenic veterinary drug used widely, has greatly attracted attention due to its potential threats. We report the nanocomposites of the α -Fe₂O₃ nanoparticles growth on reduced graphene oxide (rGO) by a one-pot method. The α -Fe₂O₃ nanoparticles are densely covered on the surface of rGO according to the observations of transmission and scanning electron microscope. The adsorptive capacity $(357.4\pm11.2 \text{ mg g}^{-1})$ of the $\begin{array}{c} -1 \ \vdots \ \mathbf{0} \end{array}$
 $\begin{array}{c} \mathbf{1} \ \mathbf{1} \end{array}$ Fe₂O₃@rGO nanocomposites for ASA, which was more than the sum of adsorptive capacities of the pure α -Fe₂O₃ nanoparticles and rGO, revealed a remarkable enhancement due to the synergetic effect of multiple interactions and the good dispersion of α -Fe₂O₃ nanoparticles with more active binding sites in the Fe₂O₃@rGO nanocomposites. The adsorption equilibrium of ASA onto the Fe₂O₃@rGO nanocomposites was achieved for 60 min, and the adsorption of ASA was dependent of pH and temperature, and independent of the concentration of humic acid ranging from 0 to $20 \text{ mg } L^{-1}$. After five cycles of adsorption-desorption, the adsorptive amounts of ASA by the regenerative sorbent still $\begin{bmatrix} -1 \\ 8 \\ 1 \end{bmatrix}$ retained 85% of adsorptive amount by the fresh sorbents. The adsorption process of ASA can be described by the Langmuir and the pseudo-second-order equations and is exothermic and spontaneous according to thermodynamic analysis.

Keywords: Arsanilic Acid, Nanocomposites, Graphene, Iron Oxide

INTRODUCTION

Arsanilic acid (4-aminphenylarsonic acid), an organic-arsenic veterinary drug, has been widely employed to promote growth rate, improve feed efficiency and control parasitic diseases for decades in animal-feeding productions [1]. Although the toxicity of arsanilic acid (ASA) is low, the most ingested ASA can be finally excreted in their original form via animal manure and transformed into high toxic inorganic arsenic derivatives under high-moisture and hightemperature conditions during composting [2,3]. The untreated livestock and poultry breeding wastewater enter the soil environment in agriculture, resulting in potential health and environmental risks [4]. Therefore, the effective removal of ASA from the culture wastewater is a crucial requirement for controlling arsenic contamination.

The concentration of ASA in the breeding wastewater is at the contamination.

The concentration of ASA in the breeding wastewater is at the level of mg L^{-1} [5]. For the removal of the low-level ASA from the culture wastewater, adsorption is the preeminent method and especially suitable for the treatment of wastewater containing low-concentration pollutants. Some sorbents have been developed for the removal of the organic-arsenic compounds, including metal-organic framework [6], metallic oxide [7], iron (oxyhydr)oxides [8], magnetic greigite [9], molecularly imprinted polymers [10], mineral [11], chitosan [12] and carbon nanotubes [13]. Recently, nanocomposites have exhibited superior performance for the adsorption of organicarsenic compounds [14-17]. Kong and Wilson [14] reported a binary

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goethite-cellulose nanocomposite with the advantage of enhanced adsorption of organic-arsenic compound due to the good dispersion of goethite nanoparticles (NPs). Hu et al. [15] developed Femodified carbon nanotubes which have higher adsorption capacity of organic-arsenic compound than the unmodified carbon nanotubes due to synergetic effect of multiple interactions. The binary nanocomposites of $Fe₃O₄$ NPs and reduced graphene oxide (rGO) were developed for the removal of organic-arsenic compounds with higher adsorption capacity compared to the pure $Fe₃O₄$ NPs and rGO because of the multiple interactions [16,17]. The rGO as the dispersion support of NPs exhibited the highest adsorption capacity of the organic-arsenic compounds because the planar geometry of rGO is amenable modification or functionalization, providing the fabricated adsorbents with the desired properties [18]. Chen and co-workers found that Fe^{3+} , Zn^{2+} and Cu^{2+} dramatically enhanced the adsorption of organic-arsenic compounds on goethite surface [19]. Compared with other metals, $Fe³⁺$ is a common and cheap element. The α -Fe₂O₃ NPs have been confirmed to be alternative sorbents for the removal of arsenic through the surface As-Fe coordination [20]. Due to the agglomeration of α -Fe₂O₃ NPs, the surface complexation of α -Fe₂O₃ NPs with ASA was weakened. The nanocomposites of the well-dispersed α -Fe₂O₃ NPs on the rGO combine the advantages of α -Fe₂O₃ NPs and rGO, which could improve the adsorptive properties of ASA [21].

In this work, the nanocomposites of $Fe₂O₃@rGO$ were developed and prepared via *in-situ* crystal growth of $Fe₂O₃$ NPs onto the surface of rGO with hydrothermal process. The objective of this work is to assess the adsorptive process of ASA by the $Fe₂O₃@rGO$ and their possible mechanisms, which would provide an understanding of the role of the Fe₂O₃@rGO for the treatment of ASA

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in the contaminated wastewater.

EXPERIMENTAL

1. Preparation and Characterization of the Fe₂O₃@rGO Nano**composite**

Graphene oxide (GO) was obtained as described in Supporting Information [22]. The product was obtained by centrifugation (16,000 rpm) for 5 min and washing with deionized water to no presence of sulfate ions. Finally, the as-synthesized GO powder was dried at 60 °C for constant weight.

 0.40 g of FeCl₃·9H₂O was dissolved in 50 mL of ethanol and then the FeCl₃ ethanol solution was mixed into 100 mL of the GO suspension aqueous solution (1 g L^{-1}) under magnetic stirring for 15min. NH3 solution (30mL) was mixed dropwise. After the hydrothermal reaction of the mixture at 180 °C for 48 h, the obtained products were isolated by centrifugation (16,000 rpm) for 10 min, washed with methanol, and dried in vacuum at 60 °C overnight. For comparison, the Fe₂O₃ NPs, and the rGO were also prepared using the identical procedure as above. All the reagents and apparatus used are also listed in Supporting Information. The characterization methods are described in Supporting Information.

2. Batch Experiments

The adsorption of ASA on the $Fe₂O₃@rGO$ was tested using batch 2. **Batch Experiments**

The adsorption of ASA on the Fe₂O₃@rGO was tested using batch

method. Effect of Fe₂O₃@rGO dosage (1-8 g L⁻¹) on the removal The adsorption of ASA on the Fe₂O₃@rGO was tested using bat method. Effect of Fe₂O₃@rGO dosage (1-8 g L⁻¹) on the remove efficiency was tested in ASA solution (20 mL) with 100 mg L⁻¹ efficiency was tested in ASA solution (20 mL) with 100 mg L^{-1} at pH 5 for 120 min. The ASA solution (20 mL) with varying concenefficiency was tested in
pH 5 for 120 min. The
tration $(100-700 \text{ mg L}^{-1})$ tration (100-700 mg L^{-1}) by adding a fixed dose of Fe₂O₃@rGO was stirred for a specified period at desired pH and temperature with stirring rate of 200 rpm. The dosage of the $Fe₂O₃@rGO$ was kept EVALUATE THE ASSEMBLE TO THE CORRECT THE STATE of the STATE of the starting rate of 200 rpm. The dosage of the Fe₂O₃@rGO was kept at 4 g L⁻¹. Temperature was adapted to 25, 35 and 45 °C to test the influence of temperature. To measure the influence of contact time on adsorption amounts, the contact time was varied from 10 to 120 min. Influence of pH (3-10) on the removal of ASA was tested ES That the community of the C formulation of the community at 25 °C for 60 min through modulating solution pH using HCl or NaOH solutions. The typically content of dissolved natural organic matter (DNOM) was below 20 mg NaOH solutions. The typically content of dissolved natural organic matter (DNOM) was below 20 mg L^{-1} in water [23]. The effect of DNOM as humic acid (HA) on the removal efficiency was tested matter (DNOM) was below 20 mg L^{-1} in NOM as humic acid (HA) on the remotin a solution containing ASA of 100 mg L^{-1} in a solution containing ASA of 100 mg L^{-1} and DNOM of 0, 5, 10, M a in a solution
 $20 \text{ mg } L^{-1}$ 20 mg L^{-1} . The pH of solution was controlled at pH 9 because HA is water-soluble in alkaline solution. The reuse of the $Fe₂O₃@rGO$ was tested in five cycles of adsorption-desorption using acidic ethand the pH is valued with the controlled with $\frac{1}{2}$ is water-soluble in alkaline solution. The reuse of the Fe₂O₃@rGO was tested in five cycles of adsorption-desorption using acidic ethanol acidified with 0.1 mol ing to the previous method [6]. The loaded Fe₂O₃@rGO was added in acidic ethanol with stirring for 6 h, and then the regenerative sorbents were filtrated, washed with deionized water near neutral and dried for the next cycle. All the trials were performed in triplicate and all the measurement data were expressed as the average± standard deviation (SD) at the 5% significance level. After stable situation, the concentration of the ASA in the remnant solution was estimated by a high-performance liquid chromatography (HPLC) method [24]. The adsorptive amount was obtained by Eq. (1) as follows.

$$
q = (C_0 - C_t) V/m \tag{1}
$$

value of $q=(C_0-C_1)V/m$ (1)
where q (mg g⁻¹) is the adsorption capacity, C_0 (mg L⁻¹) is the ini-

Fig. 1. XRD patterns of the Fe₂O₃@rGO.

Fig. 1. XRD patterns of the Fe₂O₃@rGO.
tial concentration of ASA, C_t (mg L^{-1}) is the remnant concentration of ASA, V (mL) is the volume of solution and m (g) is the mass of Fe₂O₃@rGO nanocomposite.

RESULTS AND DISCUSSION

1. Characterization

From Fig. 1, the strong and sharp XRD peaks for the crystal phase of well-crystallized Fe₂O₃ NPs are consistent with the peaks of α -Fe₂O₃ particles (JCPDS No. 33-0664) [25], illustrating that the α -Fe₂O₃ particles have been introduced into the hybrid sorbent. Lack of XRD peaks of rGO at 26° is detected due to the overlap of the large amounts of α -Fe₂O₃ particles onto the rGO surface [26].

The 1-D rod-like $Fe₂O₃$ NPs are formed in the width range of 20-50 nm (Fig. S1(a)) and the well-packed layers of RGO sheets are observed in Fig. S1(b). Large amount coverage of rod-like $Fe₂O₃$ NPs anchored uniformly on both sides of the rGO sheets can be observed in Fig. 2(a), suggesting the rGO sheets as the stable support for anchoring 1-D Fe₂O₃ NPs. Fe₂O₃ NPs are randomly and well dispersed on the rGO sheets, indicating an efficient assembly between the $Fe₂O₃$ NPs and rGO sheets during the hydrothermal process. From Fig. 2(b), the SEM image of the $Fe₂O₃@rGO$ agrees with its TEM image and the rod-like $Fe₂O₃$ NPs are anchored fully on the rGO surface. The presence of a large amount of Fe element in the Fe₂O₃@rGO from the EDS results can confirm the growth of Fe₂O₃ NPs on the rGO surface. The Fe₂O₃@rGO displays the in the Fe₂O₃@rGO from the EDS results can confirm the growth
of Fe₂O₃ NPs on the rGO surface. The Fe₂O₃@rGO displays the
BET surface area of 163 m² g⁻¹ and pore volume of 0.306 cm³ g⁻¹. The average pore diameter is estimated to be 0.548 nm. From Fig. 3, the characteristics of Fourier transmission infrared spectrum (such 22.1 strated at a b 105 m g and pote voltaire of 0.500 cm g .
The average pore diameter is estimated to be 0.548 nm. From Fig.
3, the characteristics of Fourier transmission infrared spectrum (such
as a band of Fe-O bon The average pote unanteer is estimated to be 0.5-to finite. From Fig. 3, the characteristics of Fourier transmission infrared spectrum (such as a band of Fe-O bonds at 571 cm⁻¹, the aromatic C=C stretching vibration at s, the characteristics of Fourier that
as a band of Fe-O bonds at 571 c
vibration at $1,550 \text{ cm}^{-1}$, a should
nated carboxylates at $1,625 \text{ cm}^{-1}$ nated carboxylates at $1,625 \text{ cm}^{-1}$ and a band of carboxyl group at wibration
nated carb
 $1,742 \text{ cm}^{-1}$ $1,742 \text{ cm}^{-1}$) reveal the presence of Fe₂O₃ and rGO in the Fe₂O₃@rGO [27]. The surface element distribution of the Fe₂O₃@rGO was analyzed by XPS spectrum (Fig. S1). The two peaks at 710.8 and 724.4 eV are assigned to the transitions of Fe 2p3/2 and Fe 2p1/2, which is consistent with hematite phase [28].

 (b) (a) Rod-like Fe2O3 NPs Elements wt ^o 34.3
28.1 $\tilde{\circ}$ 37.6 100.0 Total RGO 200 nm

Fig. 2. (a) TEM and (b) SEM images of the Fe₂O₃@rGO.

Fig. 3. Fourier transmission infrared spectrum of the Fe₂O₃@rGO.

2. Effect of Fe₂O₃@rGO Dosage

The removal efficiency of ASA in solution increased from 42.4% to 97.2% with the increase in the $Fe₂O₃@rGO$ dosage in the range The remeter of
to 97.2% with the 1-4 g L⁻¹ of 1-4 g L^{-1} (Fig. S2). There was no obvious difference of removal to 97.2% with the increase in the Fe₂O₃@rGO dosage in the range
of 1-4 g L⁻¹ (Fig. S2). There was no obvious difference of removal
efficiency in the Fe₂O₃@rGO dosage from 5 to 8 g L⁻¹. These results for 37.276 with the fincted of the Fe₂O₃@rGO dosage from 5 to indicate that the Fe₂O₃@rGO dosage from 5 to indicate that the Fe₂O₃@rGO dosage of 4 g L⁻¹ indicate that the Fe₂O₃@rGO dosage of 4 g L^{-1} was sufficient.

3. Effect of Contact Time

The adsorption of ASA by the $Fe₂O₃@rGO$ took place very quickly within the first 30 min (Fig. 4) due to a large amount of available binding sites in the Fe₂O₃@rGO and a large concentration gradient on the interface of solid-liquid at the beginning, and hence the transfer of ASA onto the surface of the $Fe₂O₃@rGO$ was faster. Within the second 30 min, the uptake was slowed because ASA took more time to transport from the surface to internal binding sites. After 60 min, the uptake was settled off and changed insignificantly, indicating that a state of equilibrium was reached. As time increased, the binding sites on the surface of the Fe O_3 @rGO became exhausted. Our findings were also reported by other researchers, such as a quick uptake of roxarsone in the initial stage by the carbon nanotubes [13] and by the metal-organic frameworks [29].

Fig. 4. Effect of contact time.

Therefore, 60 min was suitable as the contact time of ASA adsorption for the subsequent trials.

4. Effect of pH

ASA contains benzene ring, amino group and one arsenic ion with the aqueous dissociation constants ($pK_{a1}=1.9$ $pK_{a2}=4.1$ and $pK_{a3}=9.2$) [7] and there exists the various chemical speciation in the variation of solution pH. The point of zero charge of pH (pH_{PZC}) for the Fe₂O₃@rGO was obtained as in the previous method [30]. The pH_{pzC} of the Fe₂O₃@rGO was 6.7±0.4 according to the pH drift method [30], suggesting that the surface of the $Fe₂O₃@rGO$ was cationic below pH_{PZC} and anionic above pH_{PZC} . The pH -dependence of ASA adsorption is observed as illustrated in Fig. 5. It was found that there was no notable change in the adsorptive amount of ASA by the $Fe₂O₃@rGO$ in the range of pH 3-6 because there was an electrostatic interaction between the negatively charged ASA and the cationic surface of the $Fe₂O₃@rGO$ as the dominant interaction forces. In pH ranging from 6 to 10, the adsorptive amount of ASA by the Fe $O₃@rGO$ obviously reduced because of the electrostatic repulsion of the same charged ASA and Fe₂O₃@rGO

[31]. Another factor might be that the dissociated species of ASA at high pH inhibited the formation of hydrogen bonds with rGO [15]. These made an unfavorable adsorption of ASA by the Fe₂O₃@rGO at pH>6. However, about 65% of adsorptive amount of ASA at $pH \leq 6$ was observed in the pH range of 6-10, illustrating that the in the adsorption of ASA existed other interactions such as (As-Fe coordination) other than electrostatic interactions and/or hydrogen bonds [8,32]. Adamescu and co-workers calculated the binding energy of the As-Fe coordination up to 20 kcal and/o
and/o
lated
mol⁻¹ $mol⁻¹$ [33]. Tian and co-workers found that the contribution of the interactions to the adsorption of the organic-arsenic compounds was in the order: As-Fe coordination>hydrogen bonding> π - π interaction [16]. Therefore, the Fe₂O₃@rGO still kept a relatively high adsorptive amount at pH>6 due to the stable interaction of As-Fe coordination. Similar results had been exhibited by other research [6,34]. Therefore, the pH of 5 was used as the desired pH for the ASA adsorption in the subsequent trials.

5. Adsorbed Amounts

As the initial concentration enhanced, the adsorptive amount of ASA by the Fe₂O₃ NPs, the pure rGO and the Fe₂O₃@rGO increased (Fig. 6). The Fe₂O₃@rGO was used as an example, and the adsorptive amount of ASA was dependent on the ASA concentrations Fig. 6). The Fe₂O₃ errors was used as an example, and the adsorptive amount of ASA was dependent on the ASA concentrations from 100 to 500 mg L^{-1} in feed solution. After 500 mg L^{-1} , a small

Fig. 6. Effect of the initial ASA concentration.

variation in the adsorptive amount of ASA by the $Fe₂O₃@rGO$ was exhibited. The experimental values of the adsorptive amount of ASA by the Fe₂O₃ NPs, the rGO and the Fe₂O₃@rGO were was exhibited. The experimental values of the adsorptive amount
of ASA by the Fe₂O₃ NPs, the rGO and the Fe₂O₃@rGO were
found to be 127.5±6.7, 150.5±12.3 and 357.4±11.2 mg g^{-1} , respectively. The adsorptive amount of ASA by the $Fe₂O₃@rGO$ was more than the sum of adsorptive amounts of the $Fe₂O₃$ NPs and the rGO. This phenomenon is attributed to (1) the synergetic effect of multiple interactions of $Fe₂O₃@rGO$ with ASA (such as electrostatic force, hydrogen bonding, As-Fe coordination and π - π interaction) [16], (2) the good dispersion of $Fe₂O₃$ NPs on the rGO in the hybrid process which is a benefit for the formation of As-Fe surface coordination [35,36], and (3) the formation of more active binding sites of the Fe $_2O_3$ @rGO. These factors led to a notable increase in the adsorption amount of ASA by the $Fe₂O₃@rGO$ compared with the $Fe₂O₃$ NPs and the rGO. Similar phenomena were also described for other nanocomposites [16,17]. There was a comparison of the $Fe₂O₃@rGO$ nanocomposites in the adsorptive amount of ASA with previously reported sorbents as listed in Table 1 $[6,8,17,32,36-38]$. The Fe₂O₃@rGO nanocomposites has relatively high adsorptive amounts and relatively rapid adsorptive rate compared with most of previously reported sorbents. Fig. S3 displays that the presence of HA has no significant matrix effect. Inhibition influence on the removal efficiency of ASA was not ob-

Fig. 7. Effect of temperature.

Table 2. Isotherms parameters for the adsorption of ASA at 25 °C

Langmuir	Freundlich	$D-R$
q_{max} =423.7 mg g ⁻¹	$K_F = 19.81$ L g ⁻¹	$k_{ad} = 0.0049$ mol ² kJ ⁻²
$b = 0.01145$ L mg ⁻¹	$1/n = 0.471$	$q_s = 1,517.8$ mg g ⁻¹ $E=10.1$ mol kJ^{-1}
R^2 =0.9991	$R^2 = 0.9866$	R^2 =0.9921

served as the HA concentration ranged from 0 to 20 mg L⁻¹. Due to the formation of firm bond via surface coordination of ASA with the Fe₂O₃@rGO, ASA was not suffering the competitive adsorption with HA on the surface of $Fe₂O₃@rGO$.

6. Effect of Temperature

The adsorptive amount of ASA by the $Fe₂O₃@rGO$ decreased slightly with the increase of temperature 25-45 °C (Fig. 7), indicating that the adsorption of ASA was exothermic and a higher temperature was unfavorable for the adsorption of ASA. The exothermic characteristics of the Fe₂O₃@rGO for the ASA adsorption could be attributed to the fact that the complexation of As-Fe and the π - π interaction between rGO and ASA were exothermic reactions [13,33]. Similar trends were found by the previous reports [15,16]. **7. Adsorption Isotherm**

The equilibrium data were further fitted with three adsorption isotherms: Langmuir, Freundlich and Dubinin-Radushkevich (D-R) [39,40]. The fitting equations of these isotherms are described in Supporting Information. From Table 2 and Fig. S5, the R^2 values from Langmuir (0.9991) and D-R (0.9921) models are higher than that from Freundlich (0.9866) model, illustrating that the ASA

Table 4. Thermodynamic parameters for adsorption of ASA

Thermodynamic	Temperature (°C)		
constants	25	35	45
lnb	9.35	9.19	9.14
ΔG° (kJ mol ⁻¹)	-23.16	-23.54	-24.15
ΔH° (kJ mol ⁻¹)		-8.38	
ΔS° (J mol ⁻¹ K ⁻¹)		49.47	

adsorption maybe follows the Langmuir and D-R models. Moreover, the theoretical maximum adsorption amount of ASA from adsorption maybe follows the Langmuir and D over, the theoretical maximum adsorption amo
Langmuir model was found to be 423.7 mg g^{-1} Langmuir model was found to be 423.7 mg g^{-1} and closer to the experimental value than the theoretical value from D-R model of 1,517.8 mg g^{-1} , illustrating that Langmuir model is more suitable experimental value than the theoretical value from D-R model of 1,517.8 mg g^{-1} , illustrating that Langmuir model is more suitable to reflect the feature of the ASA adsorption on the $Fe₂O₃@rGO$ with a higher reliability. The E value from D-R model was calculated to reflect the feature of the ASA adsorption on the Fe₂O₃@rGO with a higher reliability. The E value from D-R model was calculated to be 10.1 mol kJ^{-1} , which follows the energetic range of chemical adsorption [41], illustrating that the chemical interaction between ASA and the $Fe₂O₃@rGO$ may occur during the adsorptive process [42]. In brief, the Langmuir model provides a more representative description for the ASA adsorption by the $Fe₂O₃@rGO$.

8. Kinetic Study

The data were also fitted with three kinetic models (such as pseudo-first-order (PFO), pseudo-second-order (PSO) and Elovich models) [43]. The fitting kinetic equations are described in Supporting Information. According to Table 3 and Fig. S6, the R^2 values from PFO and Elovich equations were found to be 0.9626 and 0.9176, respectively, which were lower than that obtained from the PSO equation (0.9948), demonstrating that the PFO and Elovich plots were not satisfactory for fitting the kinetic data. In addition, the q_{eq} value calculated of ASA from the PSO equation is nearly consistent with the experimental value, while the q_{eq} value calculated of ASA from the PFO equation is significantly lower than its experimental value. Therefore, the PSO model can well represent the kinetic process of ASA adsorption by the $Fe₂O₃@rGO$.

9. Thermodynamic Study

The thermodynamic parameters were calculated using the equations as listed in Supplementary Material. The standard enthalpy change (ΔH°), the standard entropy change (ΔS°) and the change in standard free energy of Gibbs (ΔG°) were calculated as described
previously (as listed in Table 4) [44]. The value of ΔG° at 25, 35
and 45 °C was found to be -19.4, -19.6 and -20.1 kJ mol⁻¹, respecpreviously (as listed in Table 4) [44]. The value of ΔG° at 25, 35 and 45 °C was found to be -19.4 , -19.6 and -20.1 kJ mol⁻¹, respectively, confirming the feasibility and spontaneity of the ASA adsorption by the Fe₂O₃@rGO [45]. The negative value of ΔH° (-8.4) tively, co
sorption
kJ mol⁻¹ kJ mol⁻¹) revealed that the uptake of ASA was exothermic, suggesting that a low temperature in the tested range was more favorable for the adsorption of ASA, which was in accordance with the results obtained from Fig. 3(d). The positive value of ΔS° (36.8 J

Fig. 8. Reusability of the regenerated Fe₂O₃@RGO.

Fig. 8. Reusability of the regenerated Fe₂O₃@RGO.
mol⁻¹ K⁻¹) suggested the increase of randomness, which might be attributed to the desorption of several molecules of solvent as each molecule of ASA adsorbed [46]. Similar results were also observed in previous literature [6,47].

10. Reusability

Five cycles of the adsorption-desorption were carried out to test the reusability of the Fe O_3 @rGO (Fig. 8). The regenerative efficiency (RE) can be calculated by Eq. (2) as follows:

$$
RE \% = (Q_{Regenerated}/Q_{Fred}) \times 100\%
$$
\n
$$
(2)
$$

RE %=($Q_{Regenerated}$ $(Q_{Fresh} \times 100\%$ (2)
where $Q_{Regenerated}$ (mg g^{-1}) is an adsorptive amount of ASA by the $\begin{split} \operatorname{RE}\% =& (\operatorname{Q}_{\textit{Regenerated}}/\operatorname{Q}_{\textit{First}}) \times 100\% \ \text{where}\ \operatorname{Q}_{\textit{Regenerated}}\left(\operatorname{mg}g^{-1}\right) \text{ is an adsorption} \ \text{regenerative} \ \text{Fe}_2\text{O}_3\varnothing\text{rGO, and}\ \operatorname{Q}_{\textit{First}}\left(\operatorname{mg}g^{-1}\right) \end{split}$ regenerative Fe₂O₃@rGO, and Q_{first} (mg g^{-1}) is an adsorptive amount of ASA by the fresh Fe₂O₃@rGO. The values of ER% decreased slightly with the increase of in cycle numbers. The adsorptive amounts of ASA by the regenerative $Fe₂O₃@rGO$ were close to 85% of the fresh Fe₂O₃@rGO, indicating that the Fe₂O₃@rGO was a reusable sorbent and could be recycled. The concentration of Fe ions in the eluent was not detected by flame atomic absorption spectrometry, indicating that there was no notable Fe leach from the $Fe₂O₃@rGO$ during the process of elution.

CONCLUSION

A nanocomposite of the Fe $O₃@rGO$ with the good performance of the ASA adsorption was prepared. The $Fe₂O₃@rGO$ nanocomposites had better-adsorbed amounts, faster-adsorbed rate and good reusability. The higher adsorptive amount of ASA by the $Fe₂O₃@rGO$ was observed in the range of pH 3-6. The adsorption amount of ASA by the Fe O_3 @rGO was improved significantly due to the synergetic effect of multiple interactions, the good dispersion of Fe $_2$ O₃ NPs on the surface of rGO and the formation of more active binding sites. There was no significant inhibition influence of HA on the removal efficiency of ASA in the range of HA from active binding sites. There was no significant inhibition influence
of HA on the removal efficiency of ASA in the range of HA from
0 to 20 mg L^{-1} . A decrease in the adsorptive amount of ASA by the Fe₂O₃@rGO was found with the increase in temperature during 25-45 °C due to its exothermicity. The ASA adsorption followed well the Langmuir and PSO models and was a spontaneous process.

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SUPPORTING INFORMATION

Additional information as noted in the text. This information is available via the Internet at http://www.springer.com/chemistry/ journal/11814.

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