



Cite this: *RSC Adv.*, 2017, 7, 23164

Effective removal of salicylic and gallic acids from single component and impurity-containing systems using an isatin-modified adsorption resin

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In this study, an isatin-modified adsorption resin named HF-02, which effectively adsorbs and removes salicylic acid (SA) and gallic acid (GA) from aqueous phase, was prepared and characterized. Its adsorption capacity towards these two adsorbates was higher than for the commercial hyper-crosslinked resins H103 and CHA-111, and the macroporous resins HPD 500, Amberlite XAD-7 and XAD-4. This is attributed to its superior porous structure and modified groups. The adsorption capacity of HF-02 towards SA is higher than for GA and its uptake amounts for SA and GA calculated by Freundlich isotherm were 350.7 mg g⁻¹ and 160.8 mg g⁻¹, respectively. The difference is mainly because the utilization ratio of adsorption sites in resins is lower when adsorbing GA and the hydrophilicity of GA is stronger than SA. The thermodynamic parameters in single component systems indicated that the two adsorption processes are exothermic and physical in nature. Furthermore, the molecular structure, initial concentration and temperature determine the uptake amounts and adsorption rate of HF-02. It is noteworthy that 1% NaOH aqueous solution and absolute ethanol gave a more efficient desorption effect in the dynamic experiment than 363 K pure water. The salt-promoting effect of NaCl, the complex-forming effect of Ni(II) and the ionization-inhibiting effect of H⁺ would enhance the uptake amounts of HF-02 in impurity-containing systems.

Received 2nd February 2017
 Accepted 4th April 2017

DOI: 10.1039/c7ra01377a

rsc.li/rsc-advances

1. Introduction

Salicylic acid (SA) and gallic acid (GA), as valuable pure chemicals and pharmaceutical intermediates, are widely employed in drug production, and in the cosmetics and food industries.^{1–5} In many complex production processes, a large amount of raw materials will be discharged with wastewater, resulting in a great waste of raw materials. To make matters worse, as typical aromatic compounds, SA and GA have potential carcinogenic and mutagenic risks.⁶ Consequently, the efficient removal and recycling of SA and GA could play a significant role in the transformation of the traditional chemical industry and in the development of the new environmental protection industry. On the other hand, as valuable plant acids, SA and GA are widely present in organic matter.^{7,8} The high concentration of the two substances in the extract solution is of great value in the reduction of material costs and in the high-value utilization of biomass.

As a green technology, adsorption has been widely utilized in recent years because of its high separation efficiency and enrichment capacity. In addition to the natural modified material adsorbents, such as activated carbon, chitosan and biosorbents, various synthetic adsorbents have been used for the adsorption of organic matter and heavy metals. Zhang and colleagues prepared a series of three-dimensionally ordered macroporous polymers. The materials have a relatively large and uniformly distributed pore size, but the specific surface area is typically relatively low.^{9,10} The materials could adsorb SA with a high adsorption capacity, but the pore sizes are much larger for SA and the utilization ratios of pores are therefore low. Metal organic frameworks are versatile materials with high porosity, adjustable pore size and various functional groups, and can be used for catalysis and adsorption. Hasan and colleagues calcined a series of Zr-based metal organic frameworks with mean pore sizes of the materials close to microporous. This is conducive to the adsorption of SA.¹¹ However, their specific surface areas are relatively low and this would be a drawback for adsorption. Graphene has reasonable adsorption properties owing to its high specific surface area and uniform adsorption sites. Lee *et al.* prepared a series of graphene-based nanosorbents through the acid or alkali heat treatment of graphene.¹² These materials possess the ability to remove SA at low concentrations, but their adsorption capacities would generally decrease at high concentrations. In

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addition, these materials are currently difficult to produce in large-scale applications.

Resins, hyper-crosslinked resins in particular, have been widely used to remove aromatic adsorbates from aqueous solution due to their high specific surface area, tunable pore structure and their superior mechanical strength.¹³ However, generally, resins also have the shortcoming of low selectivity. In 1969, Davankov produced a hyper-crosslinked resin for the first time by a post-crosslinking reaction of polystyrene.¹⁴ Since resin adsorption has the advantages of simple operation, high adsorption capacity and outstanding regenerative performance, hyper-crosslinked resins have a wide range of applications in the recovery of phenolic compounds,¹⁵ purification of natural products¹⁶ and the treatment of wastewater containing heavy metals.¹⁷ A large number of commercial hyper-crosslinked resins including H103 and CHA-111 are used in industrial production, and they generally have a high adsorption capacity.^{13,18,19} To further improve the adsorption capacity and adsorption selectivity, modification is often conducted to introduce functional groups into chloromethyl polystyrene (CMPS),²⁰ and modified amino, hydroxyl, carboxyl and carbonyl groups are effective towards aromatic adsorbates.

According to previous studies on resin properties, physical interaction was the dominant factor during the adsorption of aromatic compounds from aqueous media. In addition, the π - π conjugating effect between aromatic compounds and the benzene ring of the resin, hydrogen-bonding effect and electrostatic interaction between the adsorbate groups and the modified functional groups of the resin also contribute to the adsorption capacity.^{21,22}

In this study, an isatin-modified hyper-crosslinked resin (HF-02) was synthesized by a one-pot reaction based on chloromethylated polystyrene resin. After the modification, its polarity was raised and the steric hindrance effect of the adsorption was weakened. In addition, the benzene ring, acylamino and carbonyl groups would form a chemical interaction with SA and GA. Compared to the series of commercial resins, HF-02 displayed greater physical adsorption of SA and GA due to its higher specific surface area and more regular pore structure. In addition, the π - π conjugating effect, hydrogen-bonding effect and electrostatic interaction between the isatin group and the adsorbates further improved its adsorption capacity. HF-02 was used to adsorb and enrich SA and GA in single component systems, and in contrast, its adsorption properties in systems containing NaCl, Ni(II) and H⁺ were also investigated. Thus, the salt-promoting effect of NaCl, the complex-forming effect of Ni(II) and the ionization-inhibiting effect of H⁺ were evaluated under different conditions. Based on this study, the structure matching relationship between HF-02 and adsorbates was preliminarily explored, and could provide a theoretical basis for resource utilization and harmless treatment of the two compounds.

2. Materials and methods

2.1 Chemicals

Chloromethylated styrene-divinylbenzene copolymer (CMPS) was obtained from Tianjin Yunneng Co., Ltd. CHA-111, H103 and

HPD 500 were supplied by Zhenzhou Qinshi Co., Ltd. The Amberlite XAD-7 and XAD-4 were supplied by Rohm & Haas Company. Isatin (C₈H₅NO₂), anhydrous aluminum chloride (AlCl₃), salicylic acid (C₇H₆O₃), gallic acid (C₇H₆O₅) and nitrobenzene (C₆H₅NO₂) were obtained from Sinopharm Chemical Reagent Co., Ltd. All reagents used in the present investigation were analytical grade and were used without further purification.

2.2 Preparation of HF-02 adsorbent

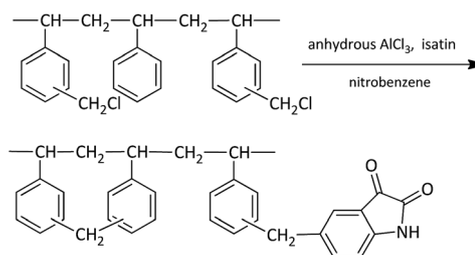
In a three-necked round-bottomed flask, 30.0 g of CMPS (17.82% chlorine equivalent weight) was swollen in 250 mL of nitrobenzene with mechanical agitation at 298 K overnight. Then 5.0 g of anhydrous AlCl₃ was added rapidly to the system and the reaction mixture was refluxed at 383 K for 4 h (8% chlorine equivalent weight remained). This post-crosslinking reaction was optimized according to an earlier study.²³ After cooling to 363 K, 11.9 g isatin (1.2 times the mole ratio to chlorine) and 2.0 g anhydrous AlCl₃ were added, and the system was refluxed for another 6 h. After that, the solid particles were filtered from the solution, and extracted by Soxhlet extractor with ultrapure water and acetone and dried at 323 K. The dried sample of resin was named HF-02.

The probable process of the reaction is shown in Scheme 1, and HF-02 was characterized by Fourier transform infrared spectroscopy (FT-IR Nicolet iN10 IR Microscope, Thermo Fisher, USA), elemental analysis (VarioELCube, GmbH, Germany) and N₂ adsorption (ASAP2020 M+C, Micromeritics, USA).

2.3 Adsorption experiments in single component systems

2.3.1 Adsorption properties of HF-02 and commercial resins. The adsorption capacity data were obtained from batch adsorption experiments; 0.200 g of HF-02 and commercial resins H103, CHA-111, HPD 500, Amberlite XAD-7 and XAD-4 were introduced into 200 mL of SA or 80 mL of GA aqueous systems (adsorbate concentrations: 100 mg L⁻¹, 250 mg L⁻¹, 500 mg L⁻¹, 750 mg L⁻¹, 1000 mg L⁻¹). The adsorption systems were placed in a 303 K constant temperature shaker and shaken at 180 rev. per min. After a 24 h static adsorption process, the adsorbents were filtered and the equilibrium concentrations of adsorbate were determined using an ultraviolet-visible spectrophotometer at specific wavelengths. Thus, the adsorption capacity of resins was calculated according to eqn (1):

$$Q_e = (C_0 - C_e) \times V/W \quad (1)$$



Scheme 1 Preparation of HF-02.



where C_0 and C_e are the concentrations of the adsorbate in the adsorption system before and after equilibrium adsorption (mg L^{-1}), V is the volume of the system (L), and W the weight of dry adsorbent (g).

The Freundlich isotherm model was employed to describe the adsorption curve according to eqn (2):²⁴

$$Q_e = K_f C_e^{1/n} \quad (2)$$

where Q_e is the equilibrium adsorbing quantity of the adsorbent (mg g^{-1}), and K_f and n are the Freundlich constants depicting the adsorption intensity.

2.3.2 Adsorption equilibrium. For the equilibrium adsorption measurements, the same adsorption systems were placed at temperatures of 288 K, 303 K and 318 K to attain equilibrium. The Langmuir and Freundlich isotherm models were employed to fit the equilibrium adsorbing quantity, and the Langmuir isotherm model was according to eqn (3):²⁵

$$\frac{C_e}{Q_e} = \frac{C_e}{Q_0} + \frac{1}{K_1 Q_0} \quad (3)$$

where Q_0 is the maximum adsorbing quantity of the adsorbent (mg g^{-1}), and K_1 is the Langmuir equilibrium constant reflecting the free energy of adsorption (L mg^{-1}).

2.3.3 Adsorption kinetics. For kinetic adsorption, about 1.0 g of HF-02 was introduced into 1000 mL of SA aqueous system (1000 mg L^{-1}) or 2.5 g of HF-02 into 1000 mL of GA aqueous system (1000 mg L^{-1}) in a conical flask, and mechanically stirred at 303 K and 120 rev. per min. During the adsorption process, the systems were sampled at preset time intervals and the residual concentration of adsorbate was monitored. The adsorption capacity at a contact time t was calculated until the adsorption equilibrium was reached. The pseudo-first-order rate equation [eqn (4)] and pseudo-second-order rate equation [eqn (5)] were employed to interpret the quantity adsorbed.^{26,27}

$$\log(Q_e - Q_t) = \log Q_e - \frac{k_1 t}{2.303} \quad (4)$$

$$\frac{t}{Q_t} = \frac{k_1 t}{k_2 Q_e^2} + \frac{t}{Q_e} \quad (5)$$

where Q_t is the adsorption capacity at time t (mg g^{-1}), and k_1 and k_2 are the adsorption rate constants of the equation.

2.3.4 Dynamic experiments with a fixed-bed column. For the dynamic adsorption-desorption experiments, 10 mL (wet volume, about 2.7 g) of HF-02 was placed in a glass column ($\Phi 7 \times 240 \text{ mm}$). SA and GA solutions with initial concentrations of 1000 mg L^{-1} were passed through the column at a rate of 6 BV per h (bed volume per hour). After the effluent concentration C_v (mg L^{-1}) became higher than $0.9C_0$, different desorption liquids, including 1% sodium hydroxide aqueous solution, absolute ethanol, 298 K pure water and 363 K pure water flowed in the reverse direction at 3 BV per h, and the adsorbate concentration in the desorption effluent was collected and monitored per volume.

The fixed-bed column was washed with ethanol and deionized water until the concentration of desorption solutions was lower than $0.02C_0$, and it was then used repeatedly in further adsorption-desorption processes and the equilibrium capacity was measured in every cycle.

2.4 Adsorption experiments in impurity-containing systems

In order to investigate the influence of inorganic salt, heavy metal ions and pH value on the adsorption behavior of SA and GA on HF-02, experiments were also carried out in systems containing various amounts of NaCl, Ni(II) and HCl or NaOH.

2.4.1 Effect of NaCl. Different mass ratios (0%, 2%, 5%, 10%, relative to weight of solution) of NaCl were added to SA and GA aqueous solutions at concentrations of 100 mg L^{-1} , 250 mg L^{-1} , 500 mg L^{-1} , 750 mg L^{-1} and 1000 mg L^{-1} . Two commercial resins, H103 and D201, were selected as references to compare their adsorption performance with HF-02.

2.4.2 Effect of Ni(II). Different concentrations of Ni(II) (0 mg L^{-1} , 20 mg L^{-1} , 50 mg L^{-1} , 100 mg L^{-1} , NiCl_2 was adopted) aqueous solutions were prepared with SA and GA at initial concentrations of 100 mg L^{-1} and 1000 mg L^{-1} , respectively, and the adsorption performance of HF-02 in the different aqueous solutions was compared.

2.4.3 Effect of pH. Different initial pH values (1, 2, 3, 4, 5, 6, 7, 8, 9 and 10, adjusted by HCl or NaOH) of SA and GA aqueous solutions with initial concentrations of 1000 mg L^{-1} were prepared, and the adsorption performance of HF-02 in the different aqueous solutions was compared. After that, the adsorption behavior was measured in different concentrations of HCl (0 mol L^{-1} , 0.5 mol L^{-1} , 1 mol L^{-1} , 2 mol L^{-1} , 3 mol L^{-1} , 4 mol L^{-1}) with initial concentrations of 1000 mg L^{-1} SA or 1000 mg L^{-1} GA.

3. Results and discussion

3.1 Characterization of HF-02

3.1.1 FT-IR spectra. Compared to CMPS (Fig. 1a), the adsorption bands at 673 cm^{-1} and 1266 cm^{-1} which were assigned to the C-Cl bond (stretching band of $-\text{CH}_2\text{Cl}$) and C-H bond (bending vibration of $-\text{CH}_2\text{Cl}$) clearly decreased in HF-02 (Fig. 1b), suggesting that the chlorine was consumed in the

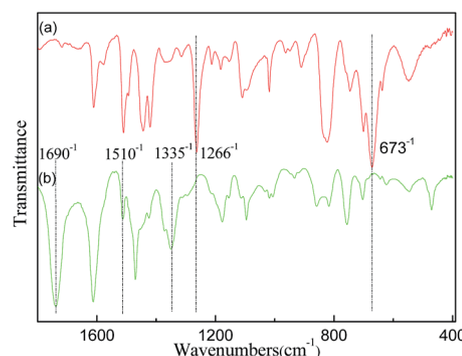


Fig. 1 IR spectra of CMPS (a) and HF-02 (b).



Friedel–Crafts reaction.²⁸ In addition, three new peaks appeared at 1690 cm^{-1} , 1510 cm^{-1} and 1355 cm^{-1} which were ascribed to the stretching vibrations of the C=O group, N–H group and C–N group, respectively (Fig. 1b), indicating that a carbonyl group and secondary amides were successfully introduced into the framework.²⁹

3.1.2 Elemental analysis. The elemental analysis results for C, O, N and Cl in the resins are shown in Table 1. The content of nitrogen and oxygen increased to 3.72% and 9.58%, respectively in HF-02, while the chlorine content decreased from 17.82% to 0.85%. The content of nitrogen indicated that the isatin group capacity of HF-02 was 2.66 mmol g^{-1} .

3.1.3 Porous structure. The adsorption and desorption isotherms of N_2 on HF-02 during pore size distribution (PSD) analysis are shown in Fig. 2, which displays the IV profiles according to the IUPAC classification with hysteric loops at a relative pressure higher than 0.9, indicating that uniform pore radii had been formed.³⁰ PSD analysis showed that the specific surface area and pore volume of HF-02 had greatly decreased (Table 2). By the post-crosslinking reaction, the larger pores are subdivided into a large number of smaller pores, and, accordingly, the specific area reaches 59.2 times that of CMPS. These transformations would provide more combining sites for pore adsorption and surface adsorption, and contribute to the higher adsorption capacity overall.³¹

3.2 Adsorption in single component systems

3.2.1 Comparison of adsorption capacity of HF-02 and commercial resins. As shown in Fig. 3, the uptake amounts of the adsorbents decrease in the following order: HF-02 > H103 > CHA-111 > HPD 500 > XAD-7 > XAD-4. The isothermal

Table 1 Elemental analysis results of the resins

Resin	Element content (%)			
	C	O	N	Cl
CMPS	75.82	—	—	17.82
HF-02	79.66	9.58	3.72	0.85

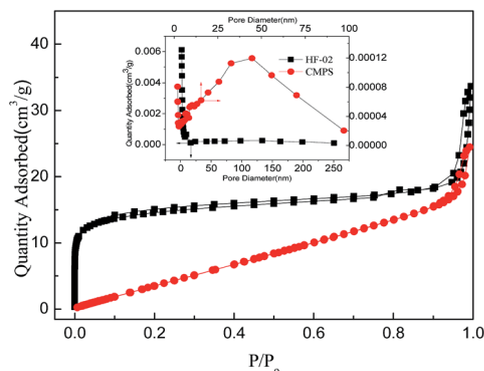


Fig. 2 N_2 adsorption–desorption isotherms and PSD curves for CMPS and HF-02.

Table 2 PSD analysis of the resins

Resin	S_{BET} ($\text{m}^2\text{ g}^{-1}$)	V_t ($\text{cm}^3\text{ g}^{-1}$)	Pore size (nm)
CMPS	20.627	0.85	26.0
HF-02	1222.9	1.17	3.8

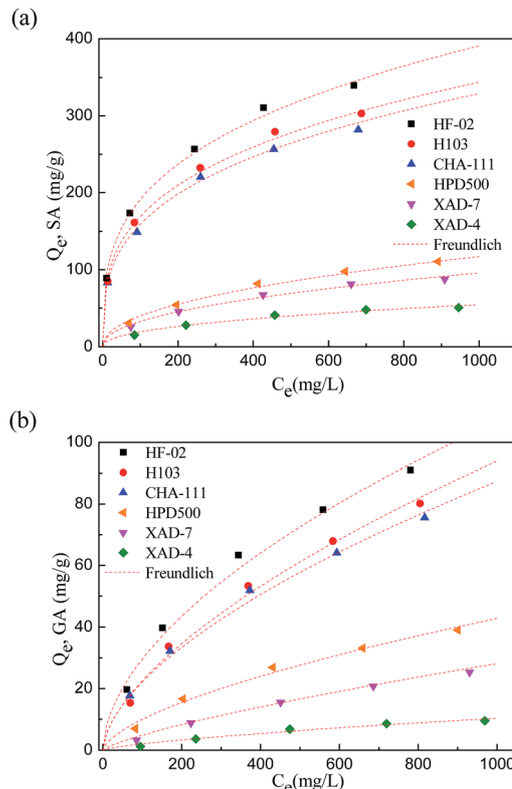


Fig. 3 Comparison of adsorption of resins towards SA (a) and GA (b) and the fitting of the Freundlich equation.

equilibrium adsorption data for SA and GA on different adsorbents are fitted by the Freundlich isotherm model, and the constants evaluated by the model are included in Table 3. HF-02 exhibited the largest K_f value among the resins, implying that it has the strongest affinity for the two adsorbates.

The different adsorption properties of the series of resins are closely related to their structural properties, and both physical and chemical interactions would be involved in the adsorption process. From the point of view of physical structure, H103 and CHA-111 are typical hyper-crosslinked resins, and XAD-7, XAD-4 and HPD 500 are representative macroporous resins. The related physical and structural parameters are presented in Table 4.

The hyper-crosslinked resin exhibited better adsorption capacities towards SA and GA due to their higher Brunauer–Emmett–Teller (BET) surface areas, and this would provide more combining sites for the adsorption of adsorbate molecules than macroporous resins.



Table 3 Fitting results of the Freundlich equation constants for the adsorbents towards its adsorption system

Adsorbent	Adsorbate	Freundlich equation	K_f	n	R^2
HF-02	SA	$\ln Q_e = 0.3109 \times \ln C_e + 3.8201$	45.610	3.2162	0.9900
	GA	$\ln Q_e = 1.2608 \times \ln C_e + 0.8182$	2.2664	0.7931	0.9690
H103	SA	$\ln Q_e = 0.3079 \times \ln C_e + 3.7134$	40.994	3.2482	0.9882
	GA	$\ln Q_e = 0.6199 \times \ln C_e + 0.2608$	1.2980	1.6133	0.9766
CHA-111	SA	$\ln Q_e = 0.3151 \times \ln C_e + 3.6193$	37.313	3.1737	0.9883
	GA	$\ln Q_e = 0.5969 \times \ln C_e + 0.3458$	1.4131	1.6751	0.9775
HPD 500	SA	$\ln Q_e = 0.4691 \times \ln C_e + 1.5217$	4.5849	2.1317	0.9859
	GA	$\ln Q_e = 0.6339 \times \ln C_e - 0.6210$	0.5374	1.5775	0.9755
XAD-7	SA	$\ln Q_e = 0.4696 \times \ln C_e + 1.3249$	3.7618	2.1341	0.9819
	GA	$\ln Q_e = 0.7613 \times \ln C_e - 1.9921$	0.1463	1.3135	0.9917
XAD-4	SA	$\ln Q_e = 0.4572 \times \ln C_e + 0.8401$	2.3166	2.1873	0.9651
	GA	$\ln Q_e = 0.6958 \times \ln C_e - 2.4747$	0.0842	1.4372	0.9582

Table 4 PSD analysis of the resins

Resin	S_{BET} ($m^2 g^{-1}$)	Average pore size (nm)	Polarity	Framework
HF-02	1222.9	3.8	Polar	Styrene (with isatin-group)
H103	1207.8	4.0	Nonpolar	Styrene
CHA-111	934.0	1.2	Weak-polar	Styrene (with carbonyl-group)
HPD 500	520	11.0	Weak-polar	Styrene (with cyano-group)
XAD-7	539	6.2	Polar	Acrylate
XAD-4	747.1	9.9	Nonpolar	Styrene

In addition, pore size is also very important for the combination of adsorbate and adsorbent. The molecular sizes of SA and GA are $0.60 \times 0.48 \times 0.11$ (nm) and $0.79 \times 0.58 \times 0.11$ (nm), respectively. According to a previous study, when the pore size of the adsorbent is 2–6 times as large as the molecular size of the adsorbate, the optimum adsorption interaction is formed.³² The narrower porous structure of CHA-111 would definitely reduce its adsorption capacity.

Due to the partial combination of Cl-sites with isatin, the post-8 crosslinking reaction of HF-02 is incomplete, but HF-02 possesses a similar porous structure to complete post-crosslinked resin H103 owing to the optimized post-crosslinking reaction researched previously.²³ Beyond that, the introduced isatin group possesses both carbonyl and secondary amine functional groups. These are able to form hydrogen bonds with the active hydrogen atoms, and the secondary amine could also act as a hydrogen donor to interact with the oxygen in the acids. All of these chemical interactions will no doubt enhance the uptake amount of adsorbate. Moreover, after the modification, part of the adsorption sites will transfer onto the introduced group, which can reduce the adsorptive steric hindrance and increase the polarity of the resin. These changes provide HF-02 with a better compatibility with the two polar adsorbate molecules.

For macroporous resins HPD 500, Amberlite XAD-7 and XAD-4, the BET surface area is generally lower and the average pore size is larger than in the hyper-crosslinked resin, and these differences provide a weaker surface adsorption capacity and a greater space wastage. For the three adsorbents, the BET

surface area, the polarity and the framework have a mutual effect on the adsorption capacity. Compared with XAD-4, the weakly polar HPD 500 is better able to contact the polar adsorbate and, in addition, its cyano group can form hydrogen bonding with the adsorbate. Similarly, the polarity and acrylate structure of XAD-7 contribute to its adsorption capacity. The adsorption capacity of HPD 500 is higher than that of XAD-7, but they possess similar BET surface areas. Besides the pore factor, the π - π conjugation adsorption probably makes a higher contribution to the adsorption capacity.

The values of K_f , a relative indicator of adsorption capacity in Freundlich theory, indicate that the adsorption capacity of the resin for SA is higher than that for GA. There are two main reasons for the difference.

On the one hand, the physical structure of the adsorbent has a selective adsorption effect towards the two adsorbates. As a sort of synthetic adsorbent, resin generally has a uniform pore distribution. The pore size distribution of resins can range from 1.1 to 25 nm, and the micropores in the applied resins account for more than 30% of the pore volume. For the two adsorbates with different molecular volumes, the interception ability of the surface layer of resin towards them would be different. Most micropores are sensitive to the molecular size of the adsorbate. The larger adsorbate molecule is more likely to be intercepted on the surface layer of the resin, and less likely to enter the inner layer of the resin and be adsorbed by the inner adsorption sites. The utilization ratio of adsorption sites is lower for larger adsorbates, and the macroscopic result of this phenomenon is that the adsorption capacity towards larger adsorbates is lower.



Accordingly, since the molecular size of GA is larger than that of SA, the adsorption of the applied resins towards GA is less efficient than towards SA.

On the other hand, the hydrophilic nature of GA is stronger as a result of its three hydroxyl groups. For SA, there are fewer hydrophilic groups on its benzene ring, and its neighboring hydroxyl and carboxyl groups are more inclined to form an intramolecular hydrogen bond, thus its hydrophilic nature is weaker than for GA. For the more hydrophilic GA, the adsorption site must overcome a stronger resistance to form an adsorption effect with it, and as a result, the uptake amounts of the adsorbent towards GA are lower.

It is the interaction of these two factors that results in the different adsorption performances of the two adsorbates. What is more, it is interesting to note that hyper-crosslinked resin exhibited a larger advantage over macroporous resin when subjected to SA. This is probably because the higher specific surface area of hyper-crosslinked resin is favorable for the adsorption of smaller-sized adsorbate molecules.

3.2.2 Adsorption equilibrium. The equilibrium adsorption isotherms of HF-02 towards SA and GA from aqueous phase at 288, 303 and 318 K are presented in Fig. 4, and it was observed that the adsorption capacities increased as the equilibrium concentration increased and the temperature decreased.

The constants evaluated by the Langmuir and Freundlich isotherm equations are included in Table 5. The monolayer adsorption capacities of SA and GA calculated by the Langmuir

Table 5 Fitting results for adsorption isotherms of SA and GA on HF-02

Adsorbate	Temperature (K)	Langmuir isotherm			Freundlich isotherm		
		Q_m	b	R^2	K_f	n	R^2
SA	288	365.2	0.019	0.908	54.21	3.394	0.993
	303	350.7	0.016	0.928	45.67	3.218	0.990
	318	326.5	0.012	0.924	35.84	3.021	0.994
GA	288	171.3	0.002	0.992	1.472	1.575	0.991
	303	160.1	0.002	0.990	1.452	1.593	0.992
	318	141.6	0.002	0.983	1.377	1.616	0.986

equation are 350.7 mg g^{-1} and 160.1 mg g^{-1} at an equilibrium concentration of 1000 mg L^{-1} at 303 K. The decrease in temperature benefits the adsorption process, which indicates that the process is exothermic and physical in nature. Freundlich equations are more suitable for correlating the equilibrium isotherm data, which shows the heterogeneity of the adsorbent surface, and the adsorption behavior was a preferential process with $n > 1$.²⁵

The adsorption equilibrium constants $K_d (=Q_e/C_e)$ at different temperatures and different initial concentrations are given in Table 6. The constants could reflect the binding capacity between the adsorbent and adsorbate.³³ K_d decreases with increasing initial concentration, and this is mainly because the adsorbate has priority to form an adsorption effect with the surface adsorption sites of the adsorbent at high concentrations, and the adsorption effect of the inner sites would be hindered, which leads to a decrease in overall binding ability. Furthermore, K_d increases with decreasing adsorption temperature, which shows that the adsorbate and adsorbent will combine more readily at lower temperatures. In addition, $K_d(\text{SA})$ is larger than $K_d(\text{GA})$ under the same adsorption conditions, which is the same sequence as for equilibrium adsorption capacity.

3.2.3 Thermodynamic investigations. The adsorption thermodynamic parameters adsorption enthalpy ΔH (kJ mol^{-1}), adsorption entropy ΔS ($\text{kJ mol}^{-1} \text{ K}^{-1}$), and adsorption free energy ΔG (kJ mol^{-1}) for the adsorption process are calculated using the Van't Hoff equation [eqn (6) and (7)].^{33,34}

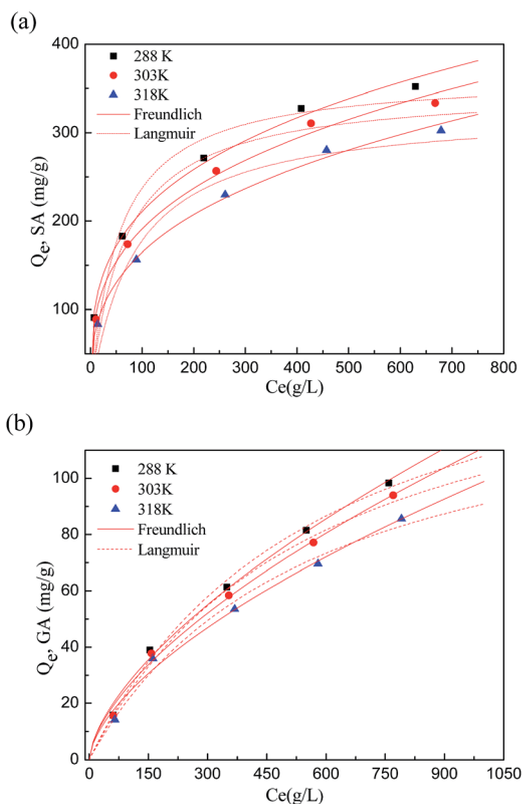


Fig. 4 Adsorption isotherms of SA (a) and GA (b) on HF-02 (solid line for the Freundlich equation, dotted line for the Langmuir equation).

Table 6 Adsorption equilibrium constants for SA and GA on HF-02

Adsorbate	C_0 (mg L^{-1})	K_d		
		288 K	303 K	318 K
SA	100	12 839.87	8015.81	5999.77
	250	3017.00	2460.29	1802.02
	500	1278.40	1050.15	916.81
	750	837.10	755.45	640.27
	1000	589.19	498.29	473.13
GA	100	256.46	251.98	214.27
	250	249.67	233.59	219.52
	500	172.95	164.90	142.39
	750	145.38	127.76	117.43
	1000	126.86	119.38	105.70



$$\Delta G^0 = -RT \ln K_d \quad (6)$$

$$\ln K_d = \frac{\Delta S^0}{R} - \frac{\Delta H^0}{RT} \quad (7)$$

where K_d is the adsorption equilibrium constant (mL g^{-1}), R is the universal gas constant ($8.314 \text{ J mol}^{-1} \text{ K}^{-1}$) and T the adsorption temperature (K).

The thermodynamic parameters for SA and GA are summarized in Table 7. The values of ΔH^0 for SA and GA are all negative and their absolute values are all less than 40 kJ mol^{-1} , which illustrates that the adsorption process is exothermic and physical in nature. Similarly, the negative values of ΔS^0 imply the decreasing degree of system chaos, and this is because the thermal motion of adsorbate molecules is impeded because of the adsorption effect. The two parameters illustrate that the adsorption process is exothermic and spontaneous reaction.

The negative values of ΔG^0 imply that the adsorption process is an exothermic reaction, and its absolute value is in the range of $10\text{--}20 \text{ kJ mol}^{-1}$ indicating that the adsorption is basically a physical process. Furthermore, it is noted that ΔG^0 increases with temperature indicating that elevated temperature can promote the spontaneous process of adsorption. In addition, the absolute values for SA are larger than those for GA, coinciding with the uptake amounts of HF-02 towards them.

3.2.4 Adsorption kinetics. The kinetic curves for SA and GA adsorbed by HF-02 under a series of conditions are shown in Fig. 5. The adsorption capacity increases over time and reaches its adsorption equilibrium in 1440 minutes.

HF-02 has a higher adsorption rate in the initial stages, as can be seen in the inset graph in Fig. 5; the adsorption capacities towards SA and GA after 60 minutes can reach 40.4% and 75.6% of the equilibrium adsorption capacities (Q_e), respectively. This can be explained by the mass transfer process. The interception effect of the surface layer of HF-02 towards GA is higher than for SA, and the utilization of inner adsorption sites is correspondingly lower. After the adsorbate is adsorbed on the surface, the inner driving force of the concentration difference is reduced accordingly. Thus, a higher utilization of inner adsorption sites needs a longer time to reach the adsorption equilibrium, and the ability of SA to reach equilibrium in unit time is poorer.

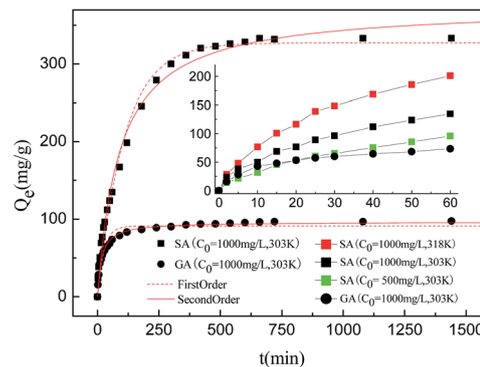


Fig. 5 Kinetic adsorption curves of HF-02 towards SA and GA.

By comparison of the adsorption behavior of SA at different temperatures and initial concentrations, it was found that the adsorption capacities towards SA at 318 K , 1000 mg L^{-1} and 303 K , 500 mg L^{-1} can reach 66.5% and 37.3% of the Q_e in 60 minutes, respectively. This could directly reflect the driving process of adsorption. A higher temperature results in a shorter time required to reach equilibrium due to the higher diffusion rate of the SA molecules into the pores of the resin, and a lower initial concentration of SA requires more time to reach equilibrium owing to the weaker driving force of the concentration gradient.

The pseudo-first-order and pseudo-second-order equations were used to fit the kinetic adsorption data, and all the constants evaluated by the application of equations are included in Table 8.

Table 8 Kinetic parameters of SA and GA on HF-02

Adsorbate	Pseudo-first-order equations			Pseudo-second-order equations		
	Q_e	k_1	R^2	Q_e	k_2	R^2
SA	327.22	0.0090	0.9858	374.67	0.0000	0.9906
GA	91.045	0.0397	0.9403	96.237	0.0006	0.9897

Table 7 Thermodynamic parameters of SA and GA on HF-02

Adsorbate	C_0 (mg L^{-1})	ΔG^0 (kJ mol^{-1})			ΔH^0 (kJ mol^{-1})	ΔS^0 ($\text{J K}^{-1} \text{ mol}^{-1}$)
		288 K	303 K	318 K		
SA	100	-22.65	-22.64	-20.83	-40.44	-60.52
	250	-19.18	-19.67	-17.95	-31.39	-41.45
	500	-17.13	-17.52	-16.33	-25.03	-26.14
	750	-16.11	-16.70	-15.47	-22.57	-21.41
	1000	-15.27	-15.65	-14.75	-20.52	-17.04
GA	100	-13.28	-13.93	-12.85	-17.71	-14.42
	250	-13.22	-13.74	-12.91	-16.43	-10.41
	500	-12.34	-12.86	-11.87	-17.06	-15.24
	750	-11.92	-12.22	-11.41	-17.01	-17.25
	1000	-11.60	-12.05	-11.16	-16.01	-14.82



Both equations fit well for SA adsorbed by HF-02 owing to the high correlation factors ($R^2 > 0.98$), which indicates that both physical diffusion and chemical adsorption are the rate-limiting factors.³⁵ The utilization of inner sites is relatively high when HF-02 adsorbs SA, and a weak driving force of the inner adsorption effect limits the adsorption rate. Furthermore, the chemical adsorption involving adsorption force through sharing or exchange of electrons between HF-02 and SA is another rate-limiting factor; this is because the secondary amine could have an electrostatic interaction with dissociated carboxyl or hydroxyl groups.²² But for GA, the kinetic adsorption data are better fitted by the pseudo-second-order equation compared to the pseudo-first-order equation, which suggests that the rate-limiting factor is primarily the chemical adsorption.

3.2.5 Dynamic adsorption-desorption of HF-02 and its regeneration. The dynamic adsorption curves for SA and GA on HF-02 from aqueous solution with an initial concentration of 1000 mg L^{-1} and a flow rate of 6 BV per h are presented in Fig. 6. The breakthrough points (at $C/C_0 = 0.05$) for SA and GA adsorption on an HF-02 column were 174 and 35 BV, respectively. The column lost its adsorption capacity ($C/C_0 > 0.9$) after 375 and 97 BV for SA and GA aqueous solutions, respectively, and the dynamic adsorption capacity towards SA and GA is similar to its behavior under static adsorption.

The Thomas model supposes that the flowing behavior in the fixed-bed adsorption column is the plug flow process.³⁶ The Thomas model is employed to correlate the adsorption curve and it is given by:

$$\ln\left(\frac{C_t}{C_0} - 1\right) = \frac{mk_T Q_T}{q} - C_0 k_T t \quad (8)$$

where m is the mass of HF-02 in the column (0.68 g), k_T is the Thomas rate constant [$\text{mL} (\text{min}^{-1} \text{mg}^{-1})$], Q_T is the adsorption capacity of the adsorbate on the adsorbent (mg g^{-1}), q is the volumetric flow rate (0.25 mL min^{-1}), and t is the time (min).

The Thomas model could fit well to the dynamic curves of the two adsorbates [$R^2(\text{SA}) = 0.9972$, $R^2(\text{GA}) = 0.9976$]. This model is based on the assumption that external and internal diffusion in the adsorbent are not the primary rate-determining steps.

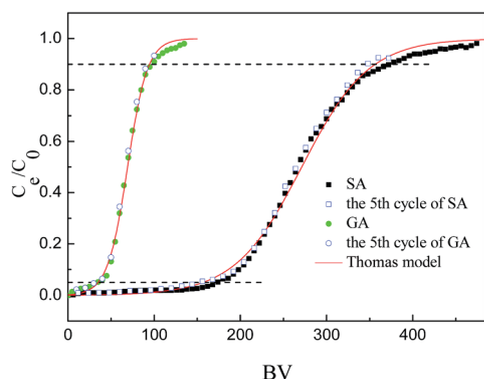


Fig. 6 Dynamic adsorption curves SA and GA on HF-02.

It is generally believed that the adsorption process consists of three stages, including external diffusion, internal diffusion and the combination process. The combination process on the adsorption sites probably limits the adsorption rate. This process between the adsorbate molecules in the solvent and the sites on the solid surface is complex, and it can be assumed that an aqueous membrane covers the solid surface, and that the molecules transferring from the aqueous membrane to the surface of the adsorbent can be reduced to a free diffusion process. In order to combine with the adsorption sites, it is necessary for the adsorbate molecules to overcome the resistance of the liquid membrane, and this is the most likely rate-determining step.³⁷

It was found that the time to reach adsorption equilibrium for SA was longer than for GA. This is because the inner adsorption process could not achieve its maximum mass transfer driving force until the external adsorption sites had achieved adsorption equilibrium, and a weak impetus driving force results in a longer time to reach adsorption equilibrium.

After HF-02 had achieved its adsorptive saturation, different desorption liquids including 1% NaOH aqueous solution, absolute ethanol, 363 K pure water and 298 K pure water, were employed to concentrate the adsorbates. The desorption curves of SA and GA with the different desorption liquids are shown in Fig. 7.

Absolute ethanol is an effective low-cost organic solvent and it could terminate the sorption effect and release the adsorbate

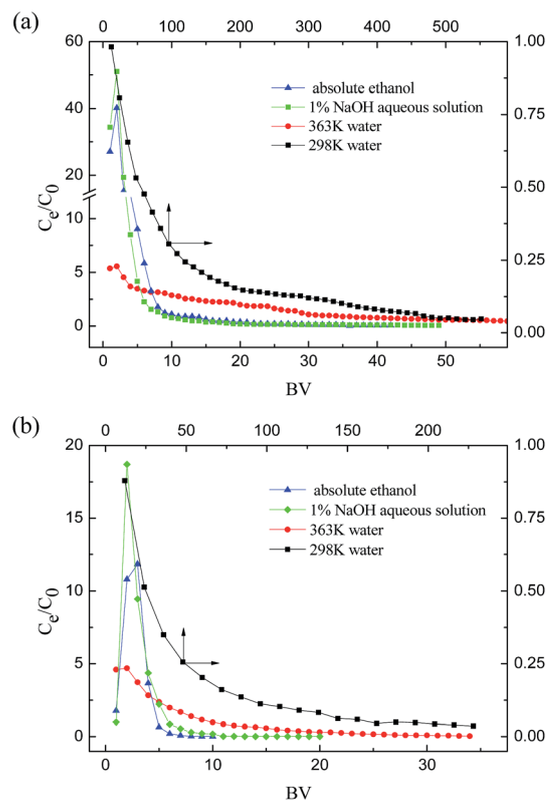


Fig. 7 Desorption curves for SA (a) and GA (b) with different desorption liquids.



into the organic phase. In addition, the hydroxyl structure of the adsorbate could be destroyed by salification in alkaline solution, and the adsorbate would be dissolved in the aqueous phase after the combined effect between the acids and adsorbent was disrupted.³⁸ For hot water, because the adsorption behavior is a kind of exothermic process, hot solvent could reduce the adsorption capacity of the adsorbent and achieve an enrichment effect.

As can be seen in Fig. 7, absolute ethanol and 1% NaOH aqueous solution achieved a high enrichment of SA and GA, and the concentration of SA and GA in the desorption liquid could reach 30 and 10 times the feed concentration, respectively. At the same time, the two desorption agents also have an effective desorption capacity, because they could collect 90% of SA and GA within 4 BV. Compared with room temperature water, 363 K pure water has an enrichment effect, and the desorption liquid can reach four times the feed concentration in the first 2 BV. In addition, it could also washup the column ($C/C_0 < 0.02$) using less solvent volume than room temperature water. Compared with the other two desorption agents, a greater volume of 363 K pure water is required to washup the column, and the concentration in solution is lower. These different desorption properties show that the destructive effect of hot water towards the combination between the adsorbates and HF-02 is weaker compared with absolute ethanol and 1% NaOH.

According to a comparative study of the three desorption methods, it is concluded that all three methods possess their respective features. Absolute ethanol has the advantages of adsorbate purification and solvent recovery because the organic solvent can be easily recycled by evaporation; however, this solvent is flammable in commercial production. Alkaline solution could treat the acid-containing wastewater more safely, but has the possibility to introduce impurities and produce secondary contamination. Hot water could recycle the adsorbate purely and safely, but its desorption technology requires more energy consumption.

The HF-02 fixed-bed resin column was then used repeatedly for five cycles in the continuous adsorption–desorption process, and its dynamic adsorption curves for the fifth cycles are presented in Fig. 6. The uptake amounts of HF-02 towards SA and GA decreased to 96.3% and 93.7%, respectively, indicating that the resin exhibited excellent reusability and possessed a reliable regeneration performance.

3.3 Adsorption experiments in impurity-containing systems

3.3.1 Effect of NaCl on adsorption of SA and GA. NaCl, widely existing in nature and in organisms, was chosen for the study of the effects of an inorganic salt on the adsorption behavior of resin towards SA and GA. As shown in Fig. 8, with mass concentrations of aqueous solution rising from 0% to 10% NaCl, the adsorption capacity of HF-02 and H103 towards SA increased by 12.88% and 9.83%, respectively. But for the ion-exchange resin D201, the adsorption capacity decreased by 86.12%. The trend of adsorption towards GA was similar to SA.

This can be interpreted by the salting-out effect.³⁹ On the one hand, the dielectric constant of the water molecule is much larger

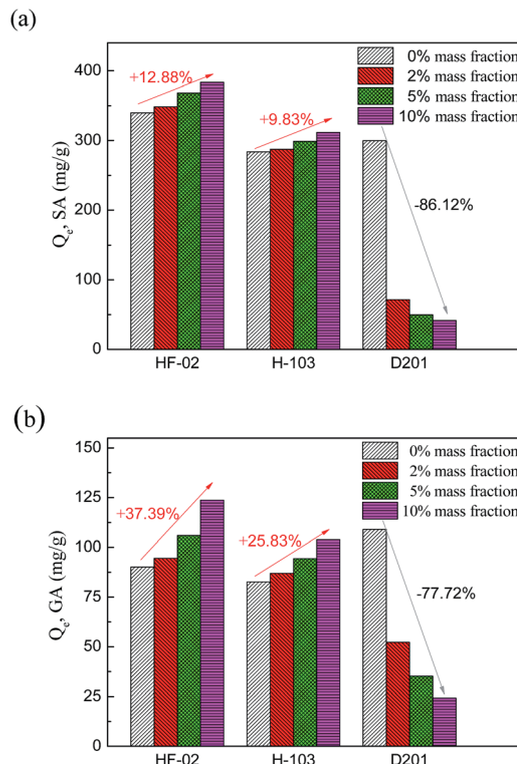


Fig. 8 Variation of adsorption capacity of HF-02, H103 and D201 towards SA (a) and GA (b) influenced by inorganic salt content.

than that of aromatic molecules and the inorganic ions in the system will attract water molecules and reject acid molecules. After the water molecules are collected by the inorganic ions, the excluded adsorbate molecules are more inclined to be adsorbed by the adsorbent. Thus, a higher concentration of inorganic salt results in a lower solubility of organic solutes and this kind of hydrophobic interaction will increase the uptake amount of the adsorbent. Furthermore, according to the theory of compression of the electric double layer,⁴⁰ more ions will thin the layer of adsorption sites and increase its conductivity, which will enhance the hydrogen bond interaction and lead to a stronger electrostatic interaction between carboxyl or hydroxyl groups and amide group. For this reason, the increased adsorptive quantity of HF-02 is much higher than that of H103. The two effects jointly promote the influence of NaCl impurities on HF-02.

For D201, as a kind of ion-exchange resin, the electrostatic interaction between hydroxyl, carboxyl and functional groups in D201 dominates its adsorption capacity. Therefore, the strong competition for adsorption sites between salt anions and adsorbate molecules could reduce the equilibrium capacity during the adsorption process,⁴¹ and the adsorption capacity of D201 decreases sharply.

The influence of NaCl at different initial concentrations of GA and SA is shown in Fig. 9. For GA, a higher initial concentration results in a larger increased proportion of adsorption capacity owing to the stronger salt-promoting effect. But for SA, the increased proportion goes against this trend at an initial concentration of 1000 mg L⁻¹ SA. Since the salting-out effect



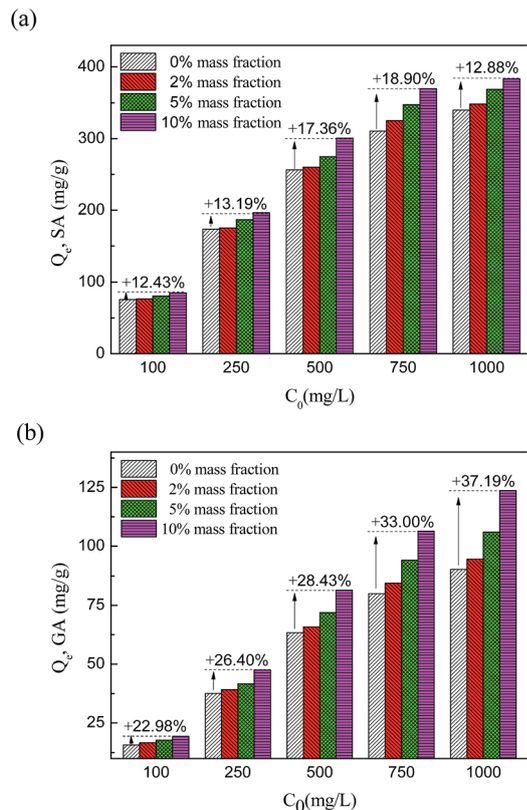


Fig. 9 Increase in adsorption capacity of HF-02 towards different concentrations of SA (a) and GA (b) influenced by inorganic salt.

could be regarded as a partial hydrophobic effect, this effect would not be obvious in a high concentration system of a hydrophobic substance.

3.3.2 Effect of Ni(II) on the adsorption of SA and GA. To study the influence of heavy metal ions, nickel ion was selected because it is widely distributed in soil and wastewater. As presented in Fig. 10, a higher concentration of Ni(II), at initial acid concentrations of 1000 mg L⁻¹, leads to higher adsorption capacities of HF-02 towards SA and GA. This is mainly because the phenolic hydroxyl groups participated in forming complex compounds with Ni(II)^{22,42} and more adsorbate molecules would be adsorbed after a single group of complex compounds formed an adsorption effect with HF-02. In the case of a limited number of adsorption sites, a higher combination capacity of the single site leads to a higher adsorption capacity of HF-02.

In contrast to the increase proportions of the two uptake amounts, the proportions of SA are much higher than those of GA. This can be explained by the structure of the complex. A Ni(II) ion can form a complex compound with four hydroxyl groups, which requires four SA molecules to form the complex with it. But for GA, which has three hydroxyl groups, fewer gallic acid molecules are required to form a stable complex compound with Ni(II), and it is considered that two molecules of GA could form a stable complex with Ni(II).⁴² The structural differences of the complex compounds lead to different uptake amounts of HF-02 towards the two molecules.

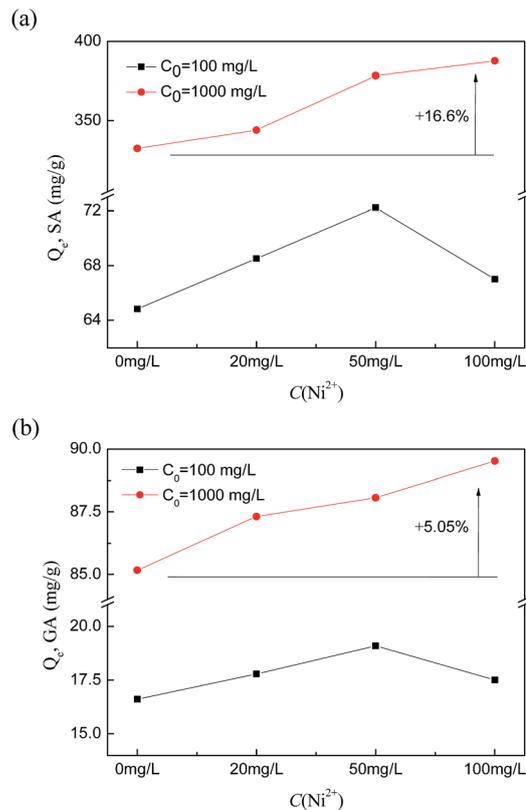


Fig. 10 Effect of Ni(II) on the adsorption of SA (a) and GA (b).

Furthermore, at lower initial concentration of acids and higher initial concentration of Ni(II), the increase proportion goes against the rising trend. This is mainly because the acylamino group of HF-02 can coordinate with the heavy metal,³⁹ and the competitive effect towards adsorption sites between Ni(II) and complex compounds would reduce the adsorption capacity towards acid molecules.

3.3.3 Effect of pH on adsorption of SA and GA. As types of weak acid, SA and GA are prone to bring about an ionization effect. Their molecular state would be transformed in different acidic and basic environments, which leads to a fluctuation of the adsorption capacities of the adsorbent towards them. The uptake amounts of HF-02 towards SA and GA influenced by the solution pH are shown in Fig. 11. In addition, the dissociation curves for SA and GA calculated from pK_a are also presented in the figure insets, as they can accurately predict the proportion of the molecular state.⁴³

It was found that the adsorption capacity of HF-02 is extremely sensitive to the pH of the aqueous solutions containing SA or GA. The tendency for adsorption is similar to the dissociation curves for SA and GA, which implies that the molecular form of the adsorbate is favorable for adsorption. In addition, the isatin group and acids all demonstrate negative charge in the alkaline solution and the charge repulsion between the two molecules will reduce the amount of adsorption of SA and GA.

What is more, it was found that HF-02 has a better adsorption capacity in strongly acidic solution, as shown in the inset



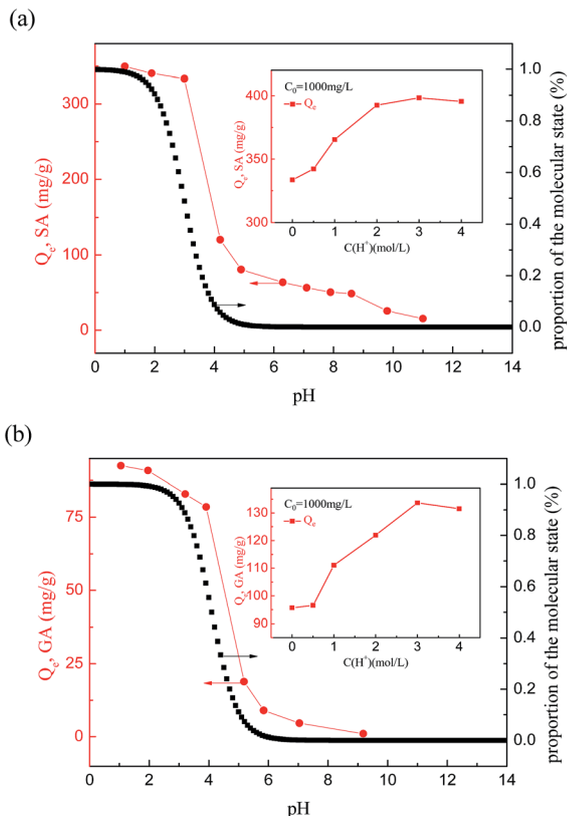


Fig. 11 Effect of pH and H⁺ concentration on the adsorption of SA (a) and GA (b).

graph in Fig. 11, and whose uptake amounts towards SA and GA increase by 19.5% and 38.9%, respectively, in the system with 4 mol L⁻¹ of H⁺. This is because the higher H⁺ concentration results in a higher degree of molecular structure due to inhibition of the ionization effect,⁴⁴ which is more favorable for the formation of hydrogen bonds between acids and isatin group. At the same time, a large amount of existing H⁺ would promote the hydrophobic interaction of adsorbate molecules owing to the formation of hydronium ions, and this extra interaction would further add to the uptake amounts of HF-02.

3.4 Structure–activity relationship between SA, GA and HF-02

After isatin had been introduced into the framework, the polarity of HF-02 increased owing to the introduction of a benzene ring and its accessory structure. The steric hindrance of the adsorption behavior became weakened owing to protrusion of the adsorption site. The joint effects of the superior specific surface area, the regular pore structure and the functional adsorption of isatin, contribute to the outstanding adsorption capacity of HF-02.

As shown in the Fig. 12, π - π conjugation of the two benzene rings between the framework and adsorbate molecules is the typical chemical adsorption effect. The acylamino group of isatin could form a stable hexatomic-ring structure by dual

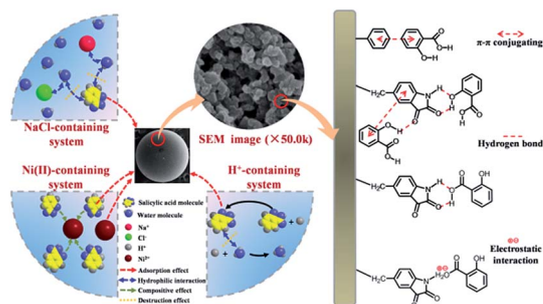


Fig. 12 Diagrammatic sketch of the structure–activity matching relationship (taking SA as an example).

hydrogen bonding with the hydroxyl groups of SA and GA, which would enhance its adsorption capacity and enthalpy. In addition, the carbonyl group and benzene ring of isatin could form additional hydrogen bonds and π - π conjugation with more adsorbate molecules. Furthermore, from the perspective of ionization, the secondary amine could form an electrostatic interaction with a dissociated carboxyl or hydroxyl group. Correspondingly, the carbonyl group and benzene ring, as kinds of nonionic functional groups, would be more stable in an alkaline environment.

4. Conclusions

An isatin-modified hyper-crosslinked resin, HF-02, which effectively adsorbs and removes SA and GA from aqueous phase, was prepared and characterized by FT-IR and elemental analysis for its functional groups, and by N₂ adsorption for its porous structure. The adsorption capacities of HF-02 towards SA and GA were higher than those of commercial hyper-crosslinked resins H103 and CHA-111, and macroporous resins HPD 500, Amberlite XAD-7 and XAD-4. Its adsorption capacity towards SA and GA at 303 K was calculated to be 350.7 mg g⁻¹ and 160.8 mg g⁻¹, respectively, using the Freundlich isotherm. In addition, the uptake amounts of SA by HF-02 were higher than those of GA; the different molecular volumes and hydrophilicities of the two adsorbates determine the adsorption capacity of HF-02. Furthermore, 1% NaOH aqueous solution and absolute ethanol had a more efficient desorption effect on HF-02 than 363 K pure water. Moreover, the adsorption capacity of HF-02 could be enhanced in systems containing NaCl, Ni(II) and H⁺. Thus, resin HF-02 is potentially useful in separating and concentrating GA and SA from single, inorganic salt-containing, heavy metal-containing and acidic systems.

Acknowledgements

The authors gratefully acknowledge the generous support provided by the Natural Science Foundation of Shandong Province (ZR2016BQ39) and Taishan Scholars Climbing Program of Shandong (No. tspd20150210).



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