Metal-organic frameworks derived magnetic carbon-αFe/Fe₃C composites as a highly effective adsorbent for tetracycline removal from aqueous solution

Weiping Xiong^{a,1}, Zhuotong Zeng^{b,1}, Guangming Zeng^{a,*}, Zhaohui Yang^{a,*}, Rong Xiao^{b,*}, Xin Li^a, Jiao Cao^a, Chengyun Zhou^a, Hongbo Chen^c, Meiying Jia^a, Yang Yang^a, Wenjun Wang^a, Xiang Tang^a

^a College of Environmental Science and Engineering, Hunan University and Key Laboratory of

Environmental Biology and Pollution Control, Ministry of Education, Hunar Heiversity,

Changsha 410082, China;

^b Department of Dermatology, Second Xiangya Hospital, Consal South University, Changsha

410011, Hunan, P R China;

^c College of Environment and Resources, Xiangtan University, Xiangtan 411105, China.



^{*}Corresponding authors at: Hunan University, College of Environmental Science and Engineering, Changsha 410082, PR China. E-mail: <u>zgming@hnu.edu.cn</u> (G.M. Zeng); <u>yzh@hnu.edu.cn</u> (Z.H. Yang) ; <u>xiaorong65@csu.edu.cn</u> (R. Xiao)

¹These authors contribute equally to this article.

Abstract

Efficient removal of tetracycline (TCN) from aqueous solution is important for water environment security. Herein, magnetic carbon-aFe/Fe₃C derived from metal-organic frameworks (MOFs) were synthesized by a carbonization method and used for TCN adsorption. The magnetic carbon-aFe/Fe₃C adsorbent was characterized by scanning electron microscope (SEM), energy dispersive X-ray spectroscopy (EDS), X-ray diffraction (XRD), Brunauer-Emmett-Teller (BET), thermal gravimetric analyzer (TGA), X-ray photoelectron spectrum (XPS) and brating sample magnetometer (VSM). The resultant magnetic carbon- α Fe/ hibited outstanding TCN adsorption capacity (511.06 mg g⁻¹) from acceous solution due to its large specific surface area (171.72 m² g⁻¹) and point y une $(0.31 \text{ cm}^3 \text{ g}^{-1})$, and abundant active adsorption sites. The main adsorption mechanisms were physisorption and ct and electrostatic adsorption. Moreover, chemisorption, including pore could remove TCN at low concentrations ($<5 \text{ mg L}^{-1}$) in magnetic carbon-αFe/Fe₃C ely low level, which was very significant for the actual lativ aqueous solution аł application. The high absorbability and good reusability in real water samples insured the magnetic carbon- α Fe/Fe₃C with great potential in practical application.

Keywords: Tetracycline removal; magnetic carbon-αFe/Fe₃C; electrostatic adsorption; practical application

1. Introduction

Antibiotics as the antibacterial agents are widely used for the treatment and prevention of diseases for human and animal, as they could effectively inhibit and kill bacteria [1, 2]. However, only less than 30% of antibiotics could be adsorbed by the human or animal and most of antibiotics are discharged into water resources via urine and could cause serious water pollution [3-5]. As a typical antibiotic, the production and usage of tetracycline (TCN) was the world's second, and its concentration in wastewater from aquaculture farms reached 20 mg L^{-1} [6, 7]. To cannot be easily biodegraded or removed by conventional wastewater plant (WWTPs) because of its high stability, causing residual tetraccline to be detected in surface water and drinking water [8, 9]. Residual tetracy ai will accumulate in the aquatic ecosystem, which may lead to the significant increase of antibiotic resistance of TCN may eventually enter the human body microorganisms [10, 11]. In the sing significant long-term threats and potential risks to through the food chain, p [12]. Therefore, the removal of TCN from water is very ecosystems and h crucial and valuable.

To date, methods for the efficient removal of antibiotics mainly include adsorption, photocatalytic, Fenton-like, filtration, bio-degradation and electrochemical degradation [13-22]. In contrast, adsorption proved the advantages of its low-cost effectiveness, environmental friendliness and simple operation, which has been considered as one of the most practical techniques to remove antibiotics from water. A variety of adsorbents have been applied to TCN adsorption [23-30]. Among many

candidates, metal-organic frameworks (MOFs) have received more attention because of their controllable structure, good thermal stability and high adsorption capacity to TCN (up to 303 mg g^{-1}) [3, 31]. Up to now, thousands of MOFs have been developed and have wide application prospects [32, 33]. Iron-based metal-organic framework MIL-53(Fe), a subclass of MIL-53(M) family, is featured with chemical versatility, abundant active sites, breathing feature and water stability [34]. Besides, the Fe³⁺ metal center with common carboxylate-type ligands can be synthesized with the characteristic of water stability [35]. Generally, MIL-53(Ee) water mainly used for advanced oxidation and adsorption [34, 36-38]. For example, (IL-53(Fe) and its magnetic hybrid have been used for adsorption representation representation and the second se and small organic molecules [34, 36]. MIL 53 Fe powders have exhibited great potential to accelerate photocatalytic degradation of organic pollutant by activating 8]. However, the crucial issues of easy persulfate and hydrogen perox recovery, regeneration and efficient adsorption are in urgent need of solution for practical applicate

Fortunately, the derivatives of MOFs prepared under thermolysis conditions have some special properties, such as magnetism, large specific surface area and pore volume, which have attracted widespread research interest [39-44]. Zhan and her co-workers have reported that porous Fe_2O_3 with preserved morphologies were synthesized by the pyrolysis of the MIL-53(Fe) and applied in catalytic selective oxidation of H₂S to sulfur [39]. Han et al. have reported that the high-performance electrocatalyst synthesized by one-step carbonization of Co-doped NH₂-MIL-53(Fe) was used for oxygen evolution reaction (OER) [40]. MnO-doped Fe₃O₄@C composite particles were prepared by a one-step thermal treatment of Mn-doped MIL-53(Fe) in Ar atmosphere and exhibited lithium-storage performance [41]. Some researchers reported that N-doped Fe/Fe₃C@C electrocatalysts were synthesized by the pyrolysis of the hexamethylenetetramine incorporated MIL-100-Fe under N₂ atmosphere and had great potential to replace Pt/C for ORR in practical use [42]. Luo et al. synthesized Fe/Fe₃C@NC by using functional carbon black combined with MIL-101(Fe) as a precursor, followed by carbonization [43]. Other research results also synthesized Fe-based catalysts by one-step Fe-NAC 383 pyrolyzation to hydrogenate conversion of carbon dioxide to valuable hydrocarbons [44]. However, the aqueous adsorption of organic pollutant only the derivatives of MOFs has not been investigated.

Within this context, we have exerct magnetic carbon- α Fe/Fe₃C derived from Zn-doped MIL-53(Fe) to the an idvanced adsorbent for highly effective removal of antibiotics from aqueus solutions. Therefore, this study provided magnetic carbon- α Fe/Fe₃C by the pyrolysis of Zn-doped MIL-53(Fe) at different temperature (810-1010 °C) under N₂ atmosphere. Morphologies, thermostability, magnetism and elementary composition were evaluated by characterization analysis. A series of adsorption experiments were carried out under different key parameters, such as ionic strength, humic acid, pH, adsorption kinetics and isotherms. Besides, the mechanism was studied in depth during the adsorption experiments. Finally, the reusability of magnetic carbon- α Fe/Fe₃C and its application in actual water environment were

discussed. It is the first time that magnetic carbon- α Fe/Fe₃C has been successfully applied to remove antibiotics, especially in real samples. We hope that our work can motivate more researches on derivatives of MOFs to be applied in the removal of water environmental pollutants.

2. Experimental section

2.1 Chemicals and Materials

Zinc (II) chloride (ZnCl₂, 99%) was received from Xiya Reagent Co., Ltd. (Shandong, China). Iron (III) chloride hexahydrate **LO.** 99%). N. N-dimethylformamide (DMF, 99.5%), 1, 4-benzenedicarbo id (1, 4-BDC, 99%) were purchased from Sinopharm Chemical Reager Co., Ltd. (Shanghai, China). Tetracycline hydrochloride (TCN) was obtained from bomei biotechnology Co., Ltd (Hefei, China). The ultra-pure water (18.25 M $\Omega \cdot \text{cm}^{-1}$, produced by ultra-pure water hroughout the whole experiments. manufacturing system, Ulpure) 2.2 Synthesis and characterization of Zn-MIL-53(Fe) and magnetic carbon-aFe/Fe₃C sis. Zn-MIL-53(Fe) was prepared by the previous reported In a typical vnth method [35]. After 0.674 g of FeCl₃ 6H₂O and 0.340 g of ZnCl₂ were dissolved in 56 mL of DMF, 0.415 g of 1, 4-BDC was added. The resulting mixture was stirred for 2

h and then transferred into a Teflon-lined bomb and heated at 170 $^{\circ}$ C for 24 h. The suspension after reaction was centrifuged. The powder product was washed with DMF and ethanol for several times, and dried at 100 $^{\circ}$ C in a vacuum. Magnetic carbon- α Fe/Fe₃C was synthesized by carbonization of Zn-MIL-53(Fe) under a nitrogen atmosphere with a heating rate of 5 $^{\circ}$ C min⁻¹ at different temperatures (810,

910, or 1010 °C) for 2 h. And then, the temperature cooled naturally to room temperature. These obtained black powder products were denoted as magnetic carbon- α Fe /Fe₃C-810, magnetic carbon- α Fe /Fe₃C-910 and magnetic carbon- α Fe /Fe₃C-1010, respectively.

The surface morphology of Zn-MIL-53(Fe) and magnetic carbon-αFe/Fe₃C-910 was examined by scanning electron microscope (SEM). The energy dispersive X-ray spectroscopy (EDS) of Zn-MIL-53(Fe) and magnetic carbon-aFe/Fe₃C-910 was obtained using an energy dispersive X-ray detector. The crystal pbe of MIL-53(Fe), Zn-MIL-53(Fe) and magnetic carbon-αFe/Fe₃C was ana X-ray diffraction (XRD). The Brunauer-Emmett-Teller (BET) spec c surface area and pore size distribution of MIL-53(Fe), Zn-MIL-53(Fe) and magnetic carbon-aFe/Fe₃C were studied by nitrogen adsorption/desorption nethod. The thermostability of magnetic al gravimetric analyzer (TGA). The surface carbon-aFe/Fe₃C was measured composition of MIL-53(F), Z-MIL-53(Fe) and magnetic carbon-αFe/Fe₃C was hot electron spectrum (XPS). The magnetic property of studied by an rav magnetic carbon-aP/Fe₃C-910 was measured by vibrating sample magnetometer (VSM).

2.3 Batch adsorption experiments

The stock TCN solution (150 mg L^{-1}) was obtained by dissolving TCN in ultrapure water, and the solutions (5-150 mg L^{-1}) with different concentrations of TCN needed in the subsequent adsorption experiments were made by diluting the stock TCN solution with ultrapure water. In order to explore the adsorption properties

of materials to TCN, 10 mg of MIL-53(Fe), Zn-MIL-53(Fe), magnetic carbon-αFe /Fe₃C-810, magnetic carbon-αFe/Fe₃C-910, magnetic carbon-αFe/Fe₃C-1010 were added to 20 mg L⁻¹ of TCN solution and then shaken at 25 $\,^{\circ}$ C with a speed of 200 rpm, respectively. Adsorption kinetic experiments were conducted with TCN concentration of 20 mg L⁻¹ in solution, while the amount of adsorbents was kept at 10 mg in 50 mL of TCN solution. The mixtures were shaken under room temperature (25 $^{\circ}$ C) with a speed of 200 rpm in a gyratory shaker. For adsorption isotherm studies, 10 mg of adsorbents were added into 50 mL solutions containing mg L⁻¹ different 5 150 concentrations of TCN and shaken at 25 °C, 35 °C and 45 speed of 200 rpm for 24 h, respectively. After the mixture was centritized at 5000 rpm for 5 min, the residual TCN concentrations in the solution were measured by a UV-vis spectrophotometer (UV-2700, Shimadzu, Jappe) at 357 nm.

The effect of pH on adsorburne CrCN was tested by performing adsorption experiments with the pH ratge of 2.0-12.0 at the TCN concentration of 20 mg L⁻¹ and adsorbent dosage Ω^2 g L⁻¹. The removal of TCN by magnetic carbon- α Fe/Fe₃C-91 was researched in the presence of different ionic strengths, humic acid and actual wastewater. To evaluate the reusability of magnetic carbon- α Fe/Fe₃C-910, 10 mg of adsorbent was added to 50 mL solution containing 20 mg L⁻¹ concentration of TCN. After shaking at 25 °C with a speed of 200 rpm, the employed adsorbent was transferred to ethyl alcohol (0.5 mol L⁻¹) and dried in a vacuum oven at 150 °C for 24 h. All the adsorption experiments were conducted in triplicates. The detailed equations and models were shown in the Table S1.

3. Results and discussion

3.1 Characterization of the adsorbents

The synthesized MIL-53(Fe) exhibited typical diffraction peaks at 9.28, 12.57, 17.68, 18.39, 25.57 and 27.36 in XRD patterns (Fig. 1), consistent with the literature [35, 45]. For the Zn-MIL-53(Fe) sample, a majority of characteristic peaks are good consistent with the main characteristic peaks of MIL-53(Fe), although there were some peaks that vary in height and strength. Besides, some new diffraction peaks, typical characteristic for zinc ion, were observed. Compared with MIL-53(Fe) and Zn-MIL-53(Fe), the typical diffraction peaks of magnetic abon-αFe/Fe₃C-810. magnetic carbon-aFe/Fe₃C-910 and magnetic carbon-aFe/Fe₃C-1010 disappeared because the high annealing temperature royed the crystal structure of dø Zn-MIL-53(Fe). The characteristic peaks at \$216, 43.03 and 77.82 are consistent with annealing at 810 °C for 2 h for the magnetic the Zn (JCPDS card No. 04-083 carbon- α Fe/Fe₃C-810. Additionally, the Zn peaks disappeared with the increase of the magnetic carbon-aFe/Fe₃C-910 and magnetic annealing temp ture carbon-αFe/Fe₃C-10.0 because of the volatilization of zinc. After the carbonization of Zn-MIL-53(Fe), the characteristic peaks at 37.83, 43.84, 46.11, 49.18 and 54.52 of Fe₃C species (JCPDS, No. 892867), as well as the characteristic peaks at 45 and 65 of α Fe (JCPDS, No. 870722), were observed in the XRD peaks of magnetic carbon-αFe/Fe₃C-810, magnetic carbon-aFe/Fe₃C-910 and magnetic carbon-αFe/Fe₃C-1010 [46]. Besides, the peak at 26.6 of graphitic carbon also can be observed, indicating that there is a special graphite structure.

"Here Fig. 1"

The typical SEM images exhibited the morphologies of the as-synthesized Zn-MIL-53(Fe) and magnetic carbon-αFe/Fe₃C-910 was shown in Fig. 2a, c and 2b, d. The element (C, O, Fe and Zn) composition of Zn-MIL-53(Fe) and magnetic carbon-αFe/Fe₃C-910 was confirmed by EDS associated with SEM in Fig. 2e-n. It was observed that the Zn-MIL-53(Fe) exhibited regular spindle structures. However, the morphology of magnetic carbon-αFe/Fe₃C-910 changed a lot and became cellular structures after carbonization, demonstrating that framework structures of Zn-MIL-53(Fe) were destroyed. The results of EDS mapping cated that zinc was successfully doped into Zn-MIL-53(Fe) but in small mounts and zinc didn't exist in this magnetic carbon- α Fe/Fe₃C-910 because of the volatilization of zinc, which was in accordance with XRD analysis. From the ICP-AES analysis, the doped amount of Zn ich confirmed the SEM result of low Zn into MIL-53(Fe) was only 0.46 content.

XPS was used a further explore the elemental composition and chemical states of absorbent materials. As shown in Fig. 3a, the survey spectrum indicated that zinc was successfully doped into Zn-MIL-53(Fe) and Zn peaks disappeared for the magnetic carbon- α Fe/Fe₃C-910 because of the volatilization of zinc, which was also consistent with XRD and SEM analyses. The spectrum of Fe 2p (Fig. 3b) showed four main peaks at 707.65, 711.12, 720.15 and 724.39 eV, which were assigned to Fe⁰ 2p_{3/2}, Fe³⁺ 2p_{3/2}, Fe⁰ 2p_{1/2} and Fe³⁺ 2p_{1/2} respectively, demonstrating the presence of Fe and

"Here Fig. 2"

Fe₃C [42, 46]. The C1s spectrum (Fig. 3c) exhibited one peak at 384.78 eV, which corresponded to C-C/C=C. The magnetic property of magnetic carbon- α Fe/Fe₃C 910 was measured by VSM system at room temperature, with the field sweeping from -20000 to +20000 Oe in Fig. 3d. The magnetization saturation values (Ms) of magnetic carbon- α Fe/Fe₃C 910 was 55.51 emu g⁻¹. The curve also showed that the magnetic carbon- α Fe/Fe₃C 910 exhibited a hysteretic behavior. The magnetic performance of magnetic carbon- α Fe/Fe₃C 910 was shown on the upper inset image of Fig. 3d. As displayed in the bottom inset image of Fig. 3d, wis thorption material of magnetic carbon- α Fe/Fe₃C 910 was quickly attracted where magnet. It was noteworthy that the coercivity (Hc) value for magnetic carbon- α Fe/Fe₃C 910 ensured that this adsorbent can be easily separated from the aqueous solution.



The N₂ aparption-disorption isotherms of MIL-53(Fe), Zn-MIL-53(Fe), magnetic carbon- α Fe/Fe₃C 810, magnetic carbon- α Fe/Fe₃C 910 and magnetic carbon- α Fe/Fe₃C 1010 were presented in Fig. 4. The type IV isotherms with a type H₃ hysteresis were also obviously shown in Fig. 4a, which demonstrated the existence of macroporous and mesoporous structure [47]. The pore size distributions of MIL-53(Fe), Zn-MIL-53(Fe), magnetic carbon- α Fe/Fe₃C 810, magnetic carbon- α Fe/Fe₃C 910 and magnetic carbon- α Fe/Fe₃C 1010 were mainly concentrated in 8.812, 5.336, 7.721, 8.687 and 9.520 nm (Fig 4b), and the BET surface areas of adsorption materials were 52.177, 46.990, 147.293, 171.715 and 174.484 m² g⁻¹, respectively. These adsorption materials exhibited pore volume distribution of mesoporous and macroporous, which was in accordance with the results in Table 1. The magnetic carbon- α Fe/Fe₃C 1010 with relatively large specific surface area and pore volume was involved in facilitating material transfer and provided additional adsorption sites to enhance adsorption capacity. The thermal stability of the magnetic carbon- α Fe/Fe₃C 810, magnetic carbon- α Fe/Fe₃C 910 and magnetic carbon- α Fe/Fe₃C 1010 was also researched in Fig. 5. The weight loss of the magnetic rarbon- α Fe/Fe₃C 810, magnetic carbon- α Fe/Fe₃C 910 and magnetic carbon- α Fe/Fe₃C 910 was 8.23%, 7.03% and 3.65% respectively, owning to the volational adsorption of water and residual organic components. It is provise mat the thermal stability of the carbon- α Fe/Fe₃C is better with the increase of carbonization temperature.



3.2 Adsorption studies

MIL-53(Fe), ZMIL-53(Fe) and adsorbents synthesized at different carbonation temperatures (magnetic carbon- α Fe/Fe₃C 810, magnetic carbon- α Fe/Fe₃C 910 and magnetic carbon- α Fe/Fe₃C 1010) showed different adsorption capacities for TCN in aqueous solution. In order to determine the optimal carbonization temperature, the adsorption experiment results are as follows. As seen in Fig. S1, the adsorption capacity of magnetic carbon- α Fe/Fe₃C 810, magnetic carbon- α Fe/Fe₃C 910 and magnetic carbon- α Fe/Fe₃C 1010 was much greater than MIL-53(Fe) and Zn-MIL-53(Fe), owning to the BET surface area, pore size and total pore volume increasing significantly after the carbonization of Zn-MIL-53(Fe). Compared the magnetic carbon- α Fe/Fe₃C 910 and magnetic carbon- α Fe/Fe₃C 1010, the adsorption capacity of magnetic carbon- α Fe/Fe₃C 1010 was slightly higher but the cost was high. Therefore, magnetic carbon- α Fe/Fe₃C 910 as the optimal adsorbent was used in whole experiment.

3.2.1 Adsorption kinetics and isotherms

Adsorption kinetics of different concentrations by magnetic carbon-αFe/Fe₃C 910 was researched and the results were n Fig. 6a, b. It can be found by fitting the experimental data with the odels that the fitting degree of pseudo-second-order model is higher, indicating that chemisorption between TCN and magnetic carbon-aFe/Fe₃C 910 controlled the adsorption rate. Besides, active sites ion capacity [48]. The calculated kinetic played an important role in t parameters of two modes wre summarized in Table 2. The determination cord-order model ($0.9961 < R^2 < 0.9999$) were higher than that coefficients of ps doof pseudo-first-order model $(0.9145 < R^2 < 0.9753)$ and the calculated $q_{e,cal}$ of pseudo-second-order equation was greater alignment with the experimental q_{e,exp}, which implied that the adsorption of TCN mainly depended on chemical interactions. In order to further elucidate the diffusion mechanism in the whole process of adsorption, the intra-particle diffusion model was utilized. It was obvious that the intra-particle diffusion in different concentrations of TCN presented in two distinct regions (Fig. 6c), which was mainly attributed to two factors such as external mass

transfer and internal particle diffusion [49]. The first linear region was the fast adsorption process, owning to macropore diffusion on the magnetic carbon- α Fe/Fe₃C 910. Besides, the second linear region was a relatively slow and gradual equilibrium adsorption process, this may be attributed to micropore diffusion [50]. As seen in Table S2, all linear portions didn't go through the origin, suggesting that intra-particle diffusion was not just a rate-limiting step [51].

"Here Fig. 6"

"Here Table 2"

Adsorption isotherms of magnetic carbon-aFe/Fe₃C ferent temperatures are shown in Fig. 6d, e, f. All the experimental data fitted the Langmuir, Freundlich and Temkin models well in Table ne determination coefficients 3. $(0.9957 < R^2 < 0.9987)$ of Langmuir model w re much higher than that of Freundlich <R²<0.8825) models, which indicated that $(0.8354 < R^2 < 0.8434)$ and Temk the homogeneous adsorptio of 7 N occurred on the magnetic carbon- α Fe/Fe₃C 910. we used for a more in-depth study on Langmuir isotherm. A separation fact $(\mathbf{R}_{\mathbf{I}})$ As seen in Table 3, II the values of the calculated parameters (RL) were between 0 and 1, indicating that the adsorption process of Langmuir isotherm was favorable. Moreover, the lower R_L values demonstrated that the interaction between TCN and magnetic carbon- α Fe/Fe₃C 910 was comparatively strong [52]. It is well-known that absorbent materials with a large affinity coefficient (KL) could have a large adsorption capacity (q_e) [53]. The affinity coefficient of magnetic carbon- α Fe/Fe₃C 910 increased with the increase of temperature, indicating that high temperature could increase the adsorption capacity of magnetic carbon- α Fe/Fe₃C 910. For further analysis of the affinity coefficient (K_L) depending on the Langmuir model at different TCN concentrations, two single point sorption coefficients (K_{0.03}, K_{0.3}) were calculated [54]. As shown in Table 3, the values of K_{0.03} and K_{0.3} increased with the increase of temperature. The higher affinity coefficients indicated the higher adsorption capacity of magnetic carbon- α Fe/Fe₃C 910 at low TCN concentrations, which was very significant for the actual application of magnetic carbon- α Fe/Fe₃C 910. In addition, the adsorption performance of TCN on magnetic carbon- α Fe/Fe₃C 910 has been compared with other samples reported, and the results were morninin Table S3. It was easy to see that the adsorption performance of magnetic carbon- α Fe/Fe₃C 910 was better than that of other adsorbents.

3.2.2 Thermodynamic analysis

The results of thermovynamics were listed in Table 4 and indicated that higher temperature as for c ossible would facilitate TCN adsorption on the carbon- α Fe/Fe₃C 911. All the results of Δ G were -30.548, -32.005 and -32.729 kJ mol⁻¹ at 25, 35 and 45 °C, respectively, demonstrating that TCN adsorption onto magnetic carbon- α Fe/Fe₃C 910 was spontaneous and thermodynamically favorable [55]. Moreover, the value of Δ H was 37.981 kJ mol⁻¹, which revealed a typical endothermic process. And this result was consistent with isothermal analysis [56]. The positive value of Δ S (0.240 kJ mol⁻¹) manifested that the randomness of the TCN adsorbed state [57, 58].

"Here Table 4"

3.2.3 The effect of ionic strength, humic acid and pH

The effect of ionic strength on TCN removal by magnetic carbon-αFe/Fe₃C 910 was exhibited in Fig. S2a. Sodium chloride (0.1, 0.2, 0.3, 0.4 and 0.5 mol L⁻¹) was added in TCN solutions (20 mg L⁻¹). Clearly, the adsorption capacity of TCN decreased with the addition of Sodium chloride, demonstrating that electrostatic interaction played a certain role in the adsorption process. Simultaneously, the significant decrease might be ascribed to the active sites compared ion on magnetic carbon-αFe/Fe₃C 910 between TCN and sodium [58]. Hu d containing many functional groups can interact with other materials rough π - π stacking interaction, which is widely found in the actual water environment [59]. As shown in Fig. S2b, the adsorbed TCN on magnetic carbon-aFeFe₃C 910 decreased with high HA concentration (30 mg L⁻¹), which attributed to complex surface complexation in the magnetic carbon-αFe Fe₃C

The pH was considered as an important factor for the removal of TCN, owning to its impact on the presence of TCN and properties of magnetic carbon- α Fe/Fe₃C 910. As shown in Fig. S3, the maximum adsorption capacity of TCN was 97.73 mg/g at pH 3, along with an appreciably decrease to 97.42 mg/g at pH 4. Whereafter, the adsorption capacity of TCN rapidly reduced with the increasing pH from 5 to 11. The adsorption trend of tetracycline was closely related to the zeta potential measurements of magnetic carbon- α Fe/Fe₃C 910 in TCN solution and the dissociation constant of tetracycline, which would be discussed in depth in the mechanism analysis part.

3.3 Mechanisms for TCN adsorption

In order to further get insight into the adsorption mechanism of magnetic carbon- α Fe/Fe₃C 910 toward TCN, physisorption and chemisorption were taken into account. Compared with pure MIL-53(Fe) and Zn-MIL-53(Fe) in Fig. 4 and Table 1, the specific surface area (171.715 m² g⁻¹) and pore volume (0.312 cm³ g⁻¹) of magnetic carbon- α Fe/Fe₃C 910 were increased to a certain extent, which provided more active adsorption sites. Correspondingly, increasing mesoporous volume (0.260 cm³ g⁻¹) of magnetic carbon- α Fe/Fe₃C 910 could weaken the teri bindrance effect and facilitate the adsorption process. Consequently, the physical adsorption was controlled by pore filling effect owning to the suitable pore size distribution.

As for the chemical adsorption, electrostatic interaction was regarded as the main chemical driving force for the removal of NN. It was well known that the different 3.32, 7.78 and 9.58, which determined the dissociation constants (pKa) of different forms of its existence in aqueous solution, including cation (TCN⁺), michs $(TCN^{-} \text{ and } TCN^{2-})$ [60]. On the other hand, the molecule (TCN) and zero-potential point of magnetic carbon-αFe/Fe₃C 910 was around 2.34 in Fig. S3. The magnetic carbon- α Fe/Fe₃C 910 has positive and negative surface charge at pH< 2.34 and pH> 2.34. Therefore, there was a strong repulsive force between the positive surface charge of magnetic carbon- α Fe/Fe₃C 910 and the cation (TCN⁺) particles at pH 2. The repulsive interaction decreased with the increased pH from 3 to 7, since TCN existed mainly in molecular form. The decrease rate of qe increased with the increase of pH (from 8 to 11), attributing to increasing repulsive force between magnetic carbon- α Fe/Fe₃C 910 and TCN. In summary, the electrostatic effect caused by the change of pH value can effectively affect TCN adsorption on magnetic carbon- α Fe/Fe₃C 910. The magnetic carbon- α Fe/Fe₃C 910 has a good adsorption performance in a wide pH range, which is conducive to the application in practical wastewater treatment.

3.4 Adsorption Behavior of magnetic carbon- α Fe/Fe₃C 910 on real samples and reusability

The investigation of magnetic carbon- α Fe/Fe₃C 910 advortion properties on TCN in tap water, river water and deionized water was control out. The quality parameters of tap water, river water and deionized water were shown in Table S4. Three kinds of tap water, river water and deionized water samples were used as the solution to provide the TCN solution (Σ^{0} mg L⁻¹), respectively. The removal efficiencies of tetracycline were Σ^{0} 10,027% and 99.56%, respectively, when the adsorption equilibrium was reached, which indicated that the dissolved organic matters and co-exeting one had adversely effect on the adsorption performance [61].

As shown in Fig. S4, magnetic carbon- α Fe/Fe₃C 910 exhibited great reusability after five cyclic adsorption experiments. Besides, the magnetic carbon- α Fe/Fe₃C 910 was easy to separate because of its magnetism and it can be readily regenerated by using ethyl alcohol on account of the solubility of TCN, which was potential in practical application in wastewater treatment.

4. Conclusion

In this work, magnetic carbon- α Fe/Fe₃C with carbonization under N₂ atmosphere

was prepared successfully with the obviously increased specific surface area and pore volume for highly efficient removal of TCN from aqueous solutions. The results indicated the properties of magnetic carbon-aFe/Fe₃C were significantly affected by different carbonization temperatures, and magnetic carbon-aFe/Fe₃C carbonized at 910 °C exhibited optimal adsorption capacity of TCN. It was found that the adsorption process was better fitted with pseudo-second-order and Langmuir isotherm models. Additionally, the adsorption mechanisms were dominated by electrostatic adsorption and pore filling effect. Future studies involved ct of pH, ionic strength, humic acid and the application in actual water en ent were taken into consideration. Most importantly, good reusability are magnetic separation property of magnetic carbon-αFe/Fe₃C 910 were mor brade to for potentially wider applications. We also sincerely expect that by changing functional metal ions and organic ligands with functional ferent kinds of MOFs can yield derivatives with different superior proerties for practical restoration of the water polluted with antibiotics.

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Figures:

Fig. 1. The XRD spectrum of the prepared adsorbents.

Fig. 2. The SEM images of Zn-MIL-53(Fe) (a and c) and magnetic carbon- α Fe/Fe₃C-910 (b and d), SEM-EDS elemental mapping images of Zn-MIL-53(Fe) (e-i) and magnetic carbon- α Fe/Fe₃C-910 (j-n).

Fig. 3. The XPS spectra of the prepared adsorbents: (a) the full XPS spectra of magnetic carbon- α Fe/Fe₃C-910, (b) Fe 2p, (c) C 1s; (d) room-temperature magnetization curve of magnetic carbon- α Fe/Fe₃C-910. Fig. 4. N₂ adsorption-desorption isotherms (a) and pore set distribution (b) of the prepared adsorbents.

Fig. 5. Thermogravimetric analysis profiles of magnetic carbon- α Fe/Fe₃C-910. Flow rate of N₂, 100 mL min⁻¹; ramping rate, 10 $^{\circ}$ min⁻¹.

Fig. 6. The pseudo-first order plots for TCN (a) adsorption; the pseudo-second order plots for TCN (b) adsorption the intra-particle diffusion model for TCN (c) adsorption. The transmuir sotherm model for TCN (d) adsorption; the Frendich isotherm model for TCN (e) adsorption; the Temkin isotherm model for TCN (f) adsorption. Reaction conditions: adsorbent loading = 0.2 g L⁻¹; temperature = 25 °C; initial pH = 7.

Tables:

Table 1. Parameters of the porous structure for adsorbents.

 Table 2. Adsorption kinetics parameters of TCN at different initial TCN concentration.

Table 3. Adsorption isotherm parameters of TCN at different solution temperature.

Table 4. The results of thermodynamic analysis of TCN adsorption.

