Nano-Porous Composites of Activated Carbon–Metal Organic Frameworks (Fe-BDC@AC) for Rapid Removal of Cr (VI): Synthesis, Adsorption, Mechanism, and Kinetics Studies

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Abstract
Metal–organic frameworks (MOFs) are a group of porous materials that display potential in the elimination of toxic industrial compounds (TICs) from polluted water streams. However, their applications have so far been held up by issues due to their physical nature and cost. In this study, activated carbon (AC) is modified with an Fe-based MOF, iron terephthalate (Fe-BDC). A facile and cost-effective impregnation method is used for enhanced removal from aqueous solutions. The new adsorbent is characterized by SEM, FTIR, PXRD, and BET. The composite displays excellent uptake of Cr (VI) when compared to un-impregnated AC with a maximum monolayer adsorption capacity of 100 mg·g⁻¹. The experimental data shows a high correlation to the Langmuir adsorption model. The adsorption kinetic study reveals that the adsorption of Cr (VI) to Fe-BDC@AC obeys the pseudo-first-order equation. The composite shows high reusability after five cycles and high adsorption rates reaching equilibrium in just 50 min. Such properties make the nanocomposite promising for water decontamination on larger scales compared to powder-based alternatives, such as individual MOF crystals.

Keywords Metal–organic frameworks · Composite · Activated carbon · Adsorbent · Cr (VI) · Water decontamination

1 Introduction
Chromium(VI) is a hazardous contaminant categorized as a group ‘A’ human carcinogen by the U.S. Environmental Protection Agency (EPA) [1]. Chromium (Cr) concentrations in drinking water are limited to 0.1 ppm by the EPA [2–4]. Cr is considered to be one of the most toxic inorganic pollutants due to its carcinogenic effects on biological species [5]. Chromium is widely used in many industries and applications resulting in large quantities of hazardous waste. One application is hardening steel and manufacturing stainless steel, which is an essential raw material for construction, heavy machinery, automobiles, transportation, energy, and medical industries [6]. Chromium is also widely used in chrome plating processes, paint pigments, and the production of dyes, leather, plastics, and photographs. In an aqueous environment, chromium usually exists as trivalent Cr(III) or hexavalent Cr (VI) oxidation state. Ionic substances containing Cr (VI), such as CrO₄²⁻ or HCrO₄⁻, have higher solubility and fluidity than Cr(III) in nature, so substances containing Cr (VI) can more easily pass through cell membranes than Cr(III) [7]. It is estimated that in an aqueous environment, Cr (VI) is almost 100 times more toxic than Cr(III) [8]. Contaminants associated with the continental crust typically have a maximum Cr concentration of 100 ppm, but many studies test at high concentrations which may inflate adsorption capacity [9].

Many methods such as adsorption [10–12], ion exchange [13], membrane separation [14], coagulation [15], chemical precipitation [16], extraction [17], and electrochemical separation [18] have been proven to remove organic contaminants and heavy metal ions, such as hexavalent chromium [19], from sewage. Among these different methods, adsorption may be the most efficient, economically feasible, environmentally sustainable, and technologically promising process [20, 21]. Recently, various adsorbents such as activated carbon [20, 22], layered double hydroxides (LDHs) [23], polymers and biomass-based materials [24] have been...
used to remove Cr (VI). In general, these adsorbents suffer from low and slow Cr (VI) sorption, and limited selectivity [25, 26].

Metal–organic frameworks (MOFs) materials have been proven to be very effective in adsorbing Cr (VI), and their highly branched structures can be easily functionalized to selectively adsorb and remove different heavy metal ions [27–30]. One of the key issues concerning the potential usage of MOFs in wastewater treatment is their physical form. MOFs normally form fine powders, which hinder their application in batch and continuous adsorption processes [31]. Form-changing processes such as pelletization have been explored to overcome this problem. However, these processes increase engineering and synthetic costs beyond cheaper adsorptive materials such as activated carbon [32]. One alternative may be to merge a MOF and activated carbon into a porous composite material that displays positive features from both materials.

In this study, the MOF selected is MIL-53 Fe (Fe-BDC). The impregnation of Fe-BDC within the AC enhances the adsorption efficiency of active carbon towards Cr (VI). Fe-BDC is chosen because Fe acts as an electron donor to form less toxic and less water soluble states such as Cr (III) [33]. The higher electron density and lower water solubility make the material easier to remove by chemisorption or physisorption with AC. Fe is also comparatively lower cost than other metals, such as copper.

This material has been used in the photocatalytic removal of organic contaminants [34]. One limitation is that pristine Fe-BDC does not have a high surface area compared to most MOFs. However, Fe-BDC can open its pores only in the presence of guest particles owing to its breathing feature and flexible structure [35]. There are a limited number of studies on MOFs-activated carbon nanocomposites used for Cr (VI) sorption. In this study, we synthesize and characterize a novel Fe-BDC@AC nanocomposite. Subsequently, we test the adsorption interaction between Fe-BDC@AC and Cr (VI) ion in an aqueous solution and study the adsorption mechanism and characteristics of Cr (VI) ion onto Fe-BDC@AC.

2 Experimental Procedure

2.1 Materials

Cinnamon sticks were obtained from the local market, as they are widely used in Egypt. Iron chloride (FeCl₃·6H₂O, 98%, Sigma Aldrich), 1,4 benzene dicarboxylic acid (BDC, 98%, Sigma Aldrich), acetic acid (98%, Sigma Aldrich), and N, N-dimethylformamide (DMF, 98%, Sigma Aldrich), are used for the synthesis of Fe-BDC. HCl (37%, Alfa Aesar) were used for regeneration purposes. Absolute ethanol, acetone, and deionized water (DIW, all from Alfa Aesar) are used for washing purposes.

2.2 Synthesis of Fe-BDC

Samples of Fe-BDC are synthesized through sequential addition of FeCl₃·6H₂O (0.27 g, 1 mmol) 1,4-benzene dicarboxylic acid (BDC) (0.166 g, mmol), and acetic acid (1 mL) into a beaker. Afterwards, 20 mL of N, N-dimethylformamide (DMF) is added to completely dissolve all reagents [36]. The mixture is poured into a closed vial and placed in an oven at 100 °C for 6 h. The resulting powder, Fe-BDC, is recovered by centrifugation at 6000 rpm for 30 min and rinsed rigorously three times with each of DMF, ethanol, acetone, and water. Finally, the recovered crystals are placed into an oven set at 140 °C to dry overnight.

2.3 Preparation of Cinnamon Sticks-Based Activated Carbon

The obtained raw materials (Cinnamon sticks) were washed with distilled water to remove the contaminants after dissolving the undesirable substances, dehydrated at 110 °C for 24 h, grounded, and sieved to achieve particle sizes below 2 mm in diameter. The dehydrated cinnamon granules were pyrolyzed to 900 °C, with a heating rate of 10 °C/min and a dwell time of 2 h under constant nitrogen gas flowing at 150 cm³/min. The activation process was done by using carbon dioxide as an activating agent. The carbonized samples were activated at 900 °C with a holding time of 120 min. The heating rate was 20 °C/min, and the flow rate of CO₂ was 150 cm³/min. Additional details about the fabrication process of AC follow previous work [37].

2.4 Synthesis of Fe-BDC@AC

For the preparation of Fe-BDC@AC, a mixture of terephthalic acid (0.166 g, 1 mmol) and AC (150 mg) is mixed and sonicated in 10 mL DMF for 5 min. A separately prepared solution of FeCl₃·6H₂O (0.27 g, 1 mmol) in 5 mL DMF and 1 mL acetic acid is then added. The vial is capped and the mixture is stirred for 8 h at 100 °C. The product is filtered and then dried in an isothermal oven at 120 °C overnight.

2.5 Characterization and Instrumentation

The synthesized Fe-BDC and Fe-BDC@AC are characterized using several techniques. Fourier transform infrared spectroscopy (FTIR, Jasco FT/IR 4100) using a KBr pellet method and Raman (Bruker Sentera) are used to identify the chemical structure. Powder X-ray diffraction (PXRD, Shimadzu XD-1) is used to verify the crystalline nature. SEM/EDX (Zeiss EVO-10 microscopy) are used to determine surface morphology and
elemental composition, respectively. The X-ray photoelectron spectrum (XPS, Thermo Fisher Scientific K-ALPHA) was acquired with a monochromatic Al-Kα (1486.7 eV). Nitrogen adsorption tests (NOVA Station A) are used to calculate BET surface areas.

### 2.6 Adsorption Studies

Adsorption experiments are accomplished in a set of 150 mL Erlenmeyer conical flasks filled with 50 mL solutions of DIW and different initial concentrations of Cr (VI) (5–50 mg/L). The solution's initial pH was adjusted to ≈ 5.5 by (0.1 M) HCl and/or NaOH. The solution was continually stirred by an orbital shaker (SI-300R). 50 mg samples of impregnated active carbon (Fe-BDC@AC) with mesh size (18*30) are used in each adsorption study. Treated solutions of Cr (VI) samples are regularly withdrawn every 5-min to measure the concentration with a double beam UV–Vis spectrophotometer (Shimadzu, Japan). A wavelength of 375 nm ($\lambda_{\text{max}}$) is implemented for all absorbance measurements. Eq. (1) is utilized to calculate the Cr (VI) removal efficiency ($\%R$) where $C_0$ and $C_t$ (mg·L⁻¹) indicate the initial Cr (VI) concentration and the concentration at time interval $t$.

$$\%R = \frac{C_0 - C_t}{C_t} \times 100$$  

(1)

All experiments are conducted in temperature-controlled environments at 25 °C. The adsorption capacities at equilibrium ($q_e$, mg·g⁻¹) are measured by applying Eq. (2) where $C_e$ (mg·L⁻¹) implies the concentration after equilibrium for the Cr (VI) solutions, $w$ denotes the mass of the adsorbent (g), and $V$ is the volume of Cr (VI) solution.

$$q_e = \frac{(C_0 - C_e)V}{w}$$  

(2)

### 3 Results and Discussion

#### 3.1 Characterization

Freshly prepared Fe-BDC and AC, as well as the nanocomposite Fe-BDC@AC, are characterized using SEM, PXRD, FTIR, and BET. The SEM image of AC, as seen in Fig. 1a, shows its highly porous nature. Figure 1b shows the SEM images of Fe-BDC with a diamond-like crystalline structure, similar to previous literature reports [38, 39]. This confirms the successful synthesis of Fe-BDC. The nanocomposite elemental mapping is shown in Fig. 1(c, d, e, f) where Fe-BDC covers the outer surface and macropores of the AC. This confirms the successful fabrication of Fe-BDC@AC.

The XRD patterns of Fe-BDC and Fe-BDC@AC are shown in Fig. 2a. The recorded 20 values are consistent
with the single crystal simulation data reported in literature [40]. The primary peaks are identified at $2\theta = 9.4^\circ$, $12.6^\circ$, $16.4^\circ$, $18.8^\circ$, $19.5^\circ$, $22.0^\circ$ and $28.5^\circ$, corresponding to (002), (101), (102), (103), (200), (202) and (211) planes, respectively [41]. Substantial alterations in the peaks can be noted between Fe-BDC and the functionalized Fe-BDC@AC. A broad C peak corresponding to the (002) plane can be observed centred at $23.3^\circ$. Another carbon peak can also be observed at $43.1^\circ$ corresponding to the (101) plane.

FTIR spectra of AC and Fe-BDC@AC are shown in Fig. 2b. Both show two strong peaks at 1377 and 1585 cm$^{-1}$ that are associated with the characteristic antisymmetric and symmetric O–C = O stretching frequency vibrations, respectively. AC shows a weak peak at 873 cm$^{-1}$ corresponding to C-H vibrations on aromatic rings that are shifted to 748 cm$^{-1}$ in the composite, as a result of the newly introduced benzene rings. By comparison, the Fe-BDC@AC sample exhibits a broad peak at 3422 cm$^{-1}$ attributed to the O–H stretching mode of the benzene ring in the BDC molecule. The metal–oxygen bond vibration ($\nu[M–O]$) at 543 cm$^{-1}$ is attributed to the interactions between the carboxyl group of terephthalic acid and Fe atoms [42]. The specific surface area of the AC decreased from 780 to 496 m$^2$/g when FE-BDC is loaded onto the AC. Figure 2c shows the substantial reduction in N$_2$ adsorption as a result of Fe-BDC covering the surfaces and filling the pores of the AC. However, the impregnation of Fe-BDC within the pores of AC greatly enhances the specific adsorption of Cr (VI) due to the presence of FeO$_6$ metal centers [36].

Figure 3a,b represents the XPS spectra of Fe-BDC@AC before and after the adsorption of Cr (VI). The presence of the principal Fe2p, Cl1s, and O1s core levels before and after adsorption with relatively similar abundances suggest good chemical stability of Fe-BDC@AC during the adsorption process. Moreover, residual chlorine was observed before adsorption tests from the precursor FeCl$_3$. It was completely detached and substituted with chromium after adsorption tests with Cr (VI).

The Cl2p core-level spectral envelope was adequately decomposed into three spin–orbit split doublets (Cl2p3/2 and Cl2p1/2) as shown in Fig. 3c, indicating three different electronic environments of chlorine. The binding energy values for the Cl2p3/2 peaks resided at 197.7, 199.0, and 200.1 eV. The first and last components suggest the presence of the ion (Cl$^-$) [43–45] and covalent (–Cl) [43, 45–48] chlorine species, respectively. The chlorine species with intermediate binding energy values (Cl*) is more appropriately associated with charge-transfer interactions between the coordinated chlorine and metal [49–52]. Further, this binding energy contribution may also be referred to as the free Cl$^-$ ion [53]. The various Cl components were expected since Cl is mainly bonded/koordinated with metallic Fe. Interestingly, all chlorine species have completely vanished after successive adsorption processes of Cr (VI). The newly observed Cr2p peaks (Fig. 3d) at 577.78, 582.31, 579.65, and 588.09 eV are attributed to Cr(III) [54–60] and Cr (VI) species [57–61], respectively. Cr (VI) was instantaneously adsorbed onto the surfaces of the Fe-BDC@AC by electrostatic attraction and ion exchange due to the positively charged surfaces, hydroxyl, and ironoxy groups of the Fe-BDC@AC. The Cr(III) could appear due to Cr (VI) being reduced by the AC directly [62]. Alternatively, FeO$_6$ metal cluster nodes could be acting as redox mediators to enhance the reduction of Cr (VI) to Cr (III) [63].

### 3.2 Effect of Contact Time and Initial Cr (VI) Concentration on the Adsorption Capacities of (Fe-BDC@AC) at Equilibrium

The effect of contact time on Cr (VI) adsorption is investigated at a concentration of 25 mg/L by measuring the decrease in absorbance at a wavelength of 375 nm ($\lambda_{max}$). The results are shown in Fig. 4a. All the experiments are
Fig. 3  XPS spectra of Fe-BDC@AC (a) before and (b) after adsorption of Cr (VI). Deconvoluted regions of the XPS spectra are shown for (c) Cl and (d) Cr.

Fig. 4  (a) Effect of initial Cr (VI) concentration on the equilibrium adsorption capacity; (b) removal percentages at different initial Cr (VI) concentrations; (c) effects of different Fe-BDC@AC doses on removal percentage for Cr (VI) at 5 ppm; (d) removal efficiency of Fe-BDC@AC against organic dyes at 5 ppm. All tests are conducted with 50 mg of adsorbent, a pH of 5.5 and room temperature (25 ± 1 ºC) conditions.
conducted at static conditions of carbon dose (50 mg), pH (5.5), room temperature (25 ± 1 ºC), and 100 rpm agitation speed. The absorbance of Cr (VI) at a wavelength of 375 nm (λ_max) decreases with an increase in contact time, indicating an increase of adsorption capacity and percent removal. No noticeable change in adsorption capacity or percent removal can be observed between 50 and 55 min, suggesting saturation of the material. The percent removal after 55 min was 85%. Therefore, the optimum contact time is approximately 55 min. This is also the equilibrium time of the batch adsorption experiments because adsorption is not changed beyond a contact time of 55 min. Equilibrium is attributed to an increase in the accumulation of Cr (VI) and the saturation of active sites and pores. The AC alone shows substantially lower adsorption capabilities, as shown in Fig. 4a. Therefore, the primary adsorption mechanism is hypothesized to be the breathing mechanism of the Fe-BDC. However, the powder-like nature of MOFs such as Fe-BDC makes it unfeasible for large-scale application in water decontamination. Secondly, the adsorption of Cr (VI) by (Fe-BDC@AC) is studied at different initial concentrations of 5, 10, 15, and 25 ppm.

The final percentage of Cr (VI) removed decreases as the initial concentration increases, as shown in Fig. 4b. A relatively low removal efficiency of 61% can be observed at 25 ppm owing to the limited pore volume and surface area of the nanocomposite. Fe-BDC@AC doses (25–200 mg) have also been studied against 5 ppm of Cr (VI) concentrations. As expected, the removal percentages have been increased with increasing absorbents doses, as shown in Fig. 4c. More active sites and vacancies are available with increasing the doses of Fe-BDC@AC, resulting in higher adsorption capacity and removal percentages.

The removal efficiency of the nanocomposite was checked against two different dyes: negatively charged methyl orange and the positively charged methylene blue, as shown in Fig. 4d. The removal efficiency has been dramatically decreased with methylene blue dye which may be due to the positively charged open metal center in the Fe-BDC. However, the molecular size of methyl orange (1.2 nm) is higher than the dichromate ion (calculated as 0.65 nm) [65]. The larger molecular size limits physical adsorption leading to lower removal efficiency than Cr(VI).

According to the classification of adsorption isotherms in an aqueous phase, the adsorption of Cr (VI) by the prepared (Fe-BDC@AC) sample is classified as class L (Langmuir-type) [66]. This type of isotherm refers to circumstances when there is no strong competition on the quantity of the solvent molecules in the coverage of the active sites on the adsorbent surface (Fe-BDC@AC) by the adsorbate (Cr (VI)). Here the adsorbate molecules are arranged horizontally in the adsorption layer [66]. Maximum sorption capacity was observed at 25 ppm. The influence of the initial Cr (VI) concentration is governed by the immediate relation between the Cr (VI) concentration and the available active binding sites on an adsorbent surface [67]. Commonly, the percent of Cr (VI) removal decreases with an increase in initial concentration, which may be due to the complete occupation of adsorption sites on the adsorbent exterior [68]. At low Cr (VI) concentrations, there will be empty and available active binding sites on the (Fe-BDC@AC). When the initial Cr (VI) concentration increases, the active sites needed for adsorption of the Cr (VI) molecules will close [69]. Still, an increase in Cr (VI) concentration will increase the adsorption capacity. This may be due to the excessive driving forces as a result of excess mass or chemical potential at higher initial Cr (VI) concentrations [70]. At lower concentrations, the ratio between Cr (VI) molecules to the free adsorption sites is small. Consequently, the adsorption shifts are independent of the original concentration. However, the available sites of adsorption decrease at high concentrations. Hence the percent removal of Cr (VI) is dependent upon initial concentration.

### 3.3 Adsorption Equilibrium Isotherms

Three isotherms were verified for their capability to define experimental results, namely the Langmuir, Freundlich, and the Temkin isotherm. Equation (3) expresses the Langmuir model where \( q_e \) (mg·g⁻¹) and \( C_e \) (mg·L⁻¹) is Cr (VI) adsorbed per unit mass of Fe-BDC@AC and Cr (VI) concentration at equilibrium, respectively. The maximum quantity of the Cr (VI) per unit mass of Fe-BDC@AC to form a whole monolayer on the surface-bound at high \( C_e \) is \( q_{\text{max}} \). The constant \( K_L \) is referred to as the affinity of the binding sites (L·mg⁻¹).

\[
\frac{C_e}{q_e} = \frac{1}{q_{\text{max}} \cdot K_L} + \frac{C_e}{q_{\text{max}}}
\]

(3)

The outline of specific adsorption (\( C_e/q_e \)) against the steadiness concentration (\( C_e \)) illustrates that the adsorption conforms to the Langmuir model, as shown in Fig. 5a. The Langmuir constants \( q_{\text{max}} \) and \( K_L \) are determined from the plot’s slope and intercept, respectively. The results are tabulated in Table 1. The separation factor, \( R_L \), is a critical characteristic of the Langmuir isotherm calculated by Eq. (4). \( C_0 \) is the uppermost initial Cr (VI) concentration (mg·L⁻¹), and \( K_L \) (L·mg⁻¹) is the Langmuir constant.

\[
R_L = \frac{1}{1 + K_L C_0}
\]

(4)
The $R_L$ value determines if the isotherm is unfavorable ($R_L > 1$), linear ($R_L = 1$), irreversible ($R_L = 0$), or favorable ($0 < R_L < 1$). The experimental value of $R_L$ is found to be 0.10 which demonstrates that the adsorption of Cr (VI) on Fe-BDC@AC is favourable. In addition, the Fe-BDC@AC sample shows excellent hexavalent chromium mono-molecular uptake of 100 mg·g⁻¹ compared to other adsorbents available in literature [10, 15, 24, 25], as shown in Table 2. Activated carbon in this work is primarily used as high porosity solid support for Fe-BDC to enable enhanced Cr (VI) adsorption efficiency in industrial applications and increase the sorbent–solute contact time compared to non-impregnated samples which show lower adsorption potential to Cr (VI).

$$q_e = \frac{R_T}{b} \ln K_f + \frac{R_T}{b} \ln C_e$$

The Temkin isotherm is extended to determine whether it is physical or chemical sorption, as shown in Eq. (6). It has commonly been applied in the linear form as described in Eq. (6), where $b$ is a constant linked to the sorption heat (J·mol⁻¹), $K_f$ is the Temkin isotherm constant (L·g⁻¹), $R$ is the gas constant (8.314 J·mol⁻¹·K⁻¹), and $T$ is the temperature (K).

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absolute temperature (K). The constants $K_T$ and $b$ can be estimated by plotting $q_e$ versus $\ln(C_e)$ in Fig. 5c. If the sorption heat is less than 1.0 kcal·mol$^{-1}$, then physical adsorption occurs. If the value is between 20 and 50 kcal/mol, then chemical adsorption is occurring. If the sorption value is between 1.0 and 20 kcal/mol, then both chemical and physical adsorption are occurring. As seen in Table 1, the Langmuir isotherm matches quite well with the experimental data, as can be seen from a correlation coefficient $R^2 > 0.99$. From the Langmuir adsorption model, the monolayer adsorption capacity is approximated as 100 mg·g$^{-1}$. This value is slightly higher than values observed experimentally (~90 mg·g$^{-1}$) but is within reasonable range if the initial Cr (VI) concentration or adsorption time is further increased. Fe-BDC active sites are assumed to have homogeneous distribution within the Fe-BDC@AC nanocomposite because the Langmuir equation assumes that the surface is similar or homogenous.

### 3.4 Adsorption Kinetics

Kinetic adsorption outcomes for different Cr (VI) concentrations were examined using a pseudo-first-order model and a pseudo-second-order model. For the pseudo-first-order kinetic model, $k_1$ values are obtained from the slopes of the linear log plots ($q_e - q_t$) versus $t$, as shown in Fig. 5d. The correlation coefficient value is determined to be 0.990. The experimental $q_e$ values are well-aligned with the estimated values obtained from the linear plots, as seen in Table 3. This shows that the adsorption of Cr (VI) to Fe-BDC@AC obeys the pseudo-first-order equation. A linear plot of $t/q_t$ versus $t$ is generated for the pseudo-second-order kinetic model, as illustrated in Fig. 5e. A weak relationship between the experimental and measured $q_e$ values can be observed, as verified in Table 3. Moreover, the correlation coefficient for the second-order kinetic model is only 0.85 for the adsorption of Cr (VI). This indicates the second-order kinetic model is less suitable to explain the Cr (VI) adsorption mechanism on the prepared Fe-BDC@AC sample.

The adsorption of Cr (VI) occurs relatively rapidly in our experiments. As can be seen from Table 2, the synthesized Fe-BDC@AC reaches max adsorption uptake or adsorption equilibrium between 50 and 55 min. By comparison, many literature values and commercially activated carbons take approximately three times this duration to reach adsorption equilibrium.

For the possible assessment of the performance of adsorbent systems, regeneration and reusability are important influencing factors. The desorption of Cr (VI) could be achieved by using a solution of 1 M HCl in the system to allow for ions exchange. Reusability experiments are conducted for five adsorption/desorption cycles. The percent removal of 5 ppm Cr (VI) is plotted in Fig. 5f. The results show that the removal efficiency decreases slightly from 85 to 80%. This confirms that the Fe-BDC@AC can retain its efficiency to an acceptable extent, and can be reused effectively. This shows the promise of MOF-AC
nanocomposites for commercial applications where both adsorption rate and reusability are extremely important parameters.

3.5 Adsorption Mechanism of Cr (VI)

The results showed that Langmuir isotherm matches well with the experimental data (correlation coefficient $R^2 > 0.99$). The adsorption kinetic study reveals that the adsorption of Cr (VI) to Fe-BDC@AC obeys the pseudo-first-order equation. These results suggest that the uptake of Cr (VI) is most likely due to physical adsorption. Therefore, the surface area is one important characteristic of the adsorbent. However, the adsorption capacity increases substantially after impregnating with Fe-BDC despite substantially reducing the BET surface area. Therefore, surface properties must play a larger role in the physical adsorption process. Electrostatic attraction between the positive surface charge of the adsorbents (AC) due to its impregnation with Fe-BDC and negatively charged Cr (VI) is critical to the mechanism, as indicated in Fig. 6. A similar trend is reported on methyl orange sorption on zirconium-based MOF [75] and chitosan/nanodiamond composite [76]. There have been previous studies that report carbonaceous adsorbents act as electron donors to reduce Cr (VI) to Cr(III)[77]. Furthermore, the presence of Cl within the MOF act as an avenue for ion exchange with the heavy metal Cr (VI)[78]. It is expected that these mechanisms further enhance the adsorption kinetics and increase the favourability of physical adsorption.

4 Conclusions

In conclusion, this study shows the synthesis of MOF-carbon nanocomposite materials. The facile functionalization of Fe-BDC onto AC generated from waste cinnamon bark is a promising, low-cost material for Cr (VI) adsorption. SEM imaging shows that the Fe-BDC covers the outer surface and fills the macropores of the AC. The removal efficiency of the new composite for Cr (VI) reaches 85% in just 50 min which is a substantially shorter time than most sorbents, such as commercial AC. One limitation of this work is higher concentrations of Cr(VI). The removal efficiency is reduced to 61% at an initial concentration of 25 ppm, suggesting more than 0.05 g of adsorbent may be required. The adsorption of the Cr (VI) on Fe-BDC@AC adsorbents could be described by the pseudo-first-order kinetic and the Langmuir isotherm model. The adsorption capacity reached up to 100 mg·g$^{-1}$ which is substantially higher than AC alone. The adsorption mechanism includes electrostatic attractions and ion exchange between negatively charged Cr (VI) and the positive surface charge of Fe and AC. The materials may be tailored to remove a broader range of contaminants than the individual components alone by controlling surface area, pore size and surface properties. This approach can be extended to the construction of other MOFs on activated carbon granules, as well as other solid supports with relative ease allowing potential application of powder-based MOFs.
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