Adsorption of Cr(VI) on Montmorillonite modified by Starch and Lanthanum


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GRAPHICAL ABSTRACT

Abstract
Industrial production brings serious Cr(VI) pollution problems, so as to explore new treatment methods for Cr(VI), In this paper, modified montmorillonite was prepared by modifying montmorillonite with starch and lanthanum nitrate, and its adsorption mechanism and effect on Cr(VI) were explored. The modified mechanism and modified monundonerization properties were characterized by Zeta potential, X-ray diffraction (XRD), Fourier Transform infrared spectroscopy (FTIR), and scanning electron microscopy analysis (SEM). The results showed that: Lanthanum-modified montmorillonite and starch-modified lanthanum-modified montmorillonite have better adsorption effects on 10 mg/L Cr(VI) polluted solution, which are 73.9% and 58.6%, respectively, while Na-based montmorillonite Cr(VI) has no adsorption capacity. Both lanthanum and starch only exist on the surface of montmorillonite, which can reduce the overall
electronegativity of montmorillonite, so that montmorillonite has the ability to adsorb Cr(VI), at the same time, after the surface of montmorillonite modified by lanthanum nitrate is covered with hydroxyl groups, it is easy to form surface coordination complexes with anions, which further enhances its adsorption effect, and provides a new research idea and direction for the adsorption and removal of Cr(VI).

**Keywords:** Montmorillonite; Starch; Lanthanum; Cr(VI)

1. Introduction

With the continuous development of the human light industry, the chromium-related chemicals are widely used in industrial production such as ignition, electroplating, thereby producing a problem of chromium wastewater. Chromium mostly exists in the form of trivalent chromium [Cr(III)] and sexivalent chromium [Cr(VI)] in aqueous solution. Some studies have found that the toxicity of the latter is much higher than that of the former, and Cr(VI) has stronger migration ability and is easy to bio-accumulate, thereby causing human carcinogenic and mutagenic through the food chain (Katlego et al., 2014). Chinese《Comprehensive Wastewater Discharge Standard》clearly stipulates that the total chromium and Cr(VI) content of chromium-containing wastewater discharged by enterprises should be lower than 1.0 mg/L and 0.2 mg/L. With high concentration, the effect in actual industrial treatment is often unsatisfactory in terms of removal effect or economic cost (Jie Tang et al., 2019). Therefore, the development of an efficient and economical Cr(VI) water removal method has become one of the important topics in the field of water pollution in China.

At present, the treatment methods of chromium-containing wastewater mainly include chemical precipitation, ion exchange, electrolysis and adsorption. The chemical precipitation method will use a large amount of reducing agent and cause secondary pollution problems such as chemical sludge at the same time (Wan Jiang et al., 2021); The ion exchange method has a high technical management threshold in the actual treatment, and there is a risk of being oxidized and causing adsorption failure (Zah A et al., 2021); The electrolysis method cannot be used in actual wastewater treatment on a large scale, and the equipment and consumption are too expensive (Liming Ren et al., 2018); The adsorption method of separating and removing pollutants in sewage through porous adsorbents has gradually become an excellent choice in Cr(VI) wastewater treatment processes due to its advantages of low cost, simple operation, and easy management.

Beverly S developed an activated carbon adsorption model based on artificial neural network, which can effectively predict the removal effect of activated carbon on Cr(VI) (Beverly S et al., 2019); Şeyma Yüksel used almond shells and walnuts to prepare activated carbon, and under optimum conditions, the adsorption capacity of chromium reached 6.67 mg/g (Şeyma Yüksel et al., 2019); Panya Maneechakr explored the preparation of activated carbon from four-cornered windmill as raw material to remove 4mg⁻¹ of Cr(VI) polluted solution, and finally obtained an adsorption capacity of 1.32 mg/g, which proved that the adsorption method can effectively remove Cr₂O₇²⁻ in the liquid. Although activated carbon as an adsorbent has a good adsorption effect on Cr(VI) and other heavy metals, it is not suitable for large-scale promotion due to its non-renewability and high cost (Panya Maneechakr et al., 2017). As montmorillonite, which can be seen everywhere in nature and is easy to obtain, it also has a stable structure and a large
specific surface area, and is widely used as an adsorbent for the adsorption of Cr(VI). Anthony improved the processing efficiency of Cr(VI) to 100% by thermal synthesis of cerium/montmorillonite (Anthony Eric T et al., 2020). Li T achieved an adsorption effect of 156.25 mg/g on Cr(VI) through montmorillonite-supported carbon nanospheres (Li T et al., 2014). Yang Jing modified montmorillonite with tea polyphenols to improve the regeneration activity of Fe, so as to fix Cr(VI) in the soil more friendly (Yang Jing et al., 2021). Zhang Dading prepared a new montmorillonite material loaded with carboxymethyl cellulose, and the adsorption rate of Cr(VI) was increased to 424.6 mg/kg (Zhang Dading et al., 2021). Shengrong Liu used CPC to modify montmorillonite, which reduced the interlayer distance between CPC and montmorillonite and improved the adsorption rate of Cr(VI) (Shengrong Liu et al., 2020).

On the basis of the research on the adsorption of Cr(VI) by montmorillonite, some researchers use specific substances to modify and replace the low-valent cations in montmorillonite to achieve the effect of changing the properties and electrical properties of montmorillonite itself. A montmorillonite modified system was constructed to obtain better adsorption effect. Mirle Vinuth used Fe(II) modification to enhance the adsorption of Cr(VI) in Fe(II)-Mt system under acidic conditions (Mirle Vinuth et al., 2019); B.S. Krishna used polymer materials to improve the positive charge of montmorillonite to improve its adsorption effect. Montmorillonite is electronegative as a whole, and Cr(VI) often exists in the form of dichromate ions in water. Starch is a natural polymer material, and it is electropositive (B.S. Krishna et al., 2000). It has a wide range of sources and is inexpensive. It is possible to coexist with clay minerals such as montmorillonite in nature. Lanthanum (La), as a light rare earth material, can replace low-valent cations in the montmorillonite system, thereby improving its electrical properties. Can a coexisting system of two modifiers and montmorillonite produce a modified material? What is the adsorption effect of the composite modified material shared by two single modified materials and two adsorbents? In this study, montmorillonite was modified individually and in combination with starch and lanthanum nitrate, and the physical and chemical characterization of the modified materials was carried out to explore the adsorption performance of Cr(VI) in aqueous solution, and to reveal the relationship between montmorillonite and natural montmorillonite. The interaction between organic polymers and light rare earth materials, and their possible environmental governance effects, also provide new research ideas and directions for the adsorption and removal of Cr(VI).

2. Materials and methods

2.1. Reagents and instruments

Electronic balance, UV-VIS spectrophotometer, magnetic stirrer (78-1), pH acidity meter (Raymag PHS-3E), water bath constant temperature oscillator (SHZ-82A), desktop low-speed centrifuge (TDA-5A), Electric heating blast drying oven, X-ray diffractometer (Empyrean), Fourier transform infrared spectrometer, high sensitivity Zeta potential and particle size analyzer (nano ZS), field emission scanning electron microscope (SEM).

Montmorillonite (purity>97%), produced in Shijiazhuang City, Hebei Province, its main phase is calcium-based montmorillonite, contains a small amount of quartz impurities, and its chemical components are mainly Si, O, Al, Ca, etc., hardness is 1, the density is about 2 g.cm\(^{-3}\), and the
purity is 97%; Acetic acid, experimental corn starch, sodium hydroxide, sodium chloride, potassium dichromate, all purchased from Shanghai Aladdin Biochemical Technology Co., Ltd.; concentrated Sulfuric acid and concentrated nitric acid were purchased from Shanghai Standard Technology Co., Ltd. All the above reagents except montmorillonite are of analytical grade. The test water was deionized water.

2.2. Sample preparation

2.2.1 Preparation of Na-based montmorillonite (Na-Mont)

Dissolve 10 g of 97% high-purity montmorillonite [Montmorillonite (hereinafter referred to as Mont)] and 1 mol/L NaCl in a 250 mL volumetric flask, stir in a magnetic stirrer for 24 h, remove the supernatant by centrifugation and deionize washed with water, dried in an electric blast drying oven at 65°C for 24 hours, and crushed to obtain Na-based montmorillonite (Na-Mont).

2.2.2 Preparation of La-Modified Montmorillonite (La-Mont)

Weigh 4g of lanthanum nitrate hexahydrate [La(NO)₃·6H₂O] and 10g of prepared Na-Mont to dissolve in a 250 mL volumetric flask, stir in a magnetic stirrer for 8h, and remove the supernatant by centrifugation and washed with deionized water, dried in an electric heating blast drying oven at 65°C for 24 hours, and crushed to obtain Na-based montmorillonite (La-Mont).

2.2.3 Preparation of Starch Modified La Montmorillonite (St-La-Mont)

Weigh 1 g of test starch (St), dissolve it in 50 mL of acetic acid (5%), add 10 g of the prepared La-Mont, stir in a magnetic stirrer for 24 h, remove the supernatant by centrifugation and wash with deionized water, dried in a high-temperature drying oven at 65°C for 24 hours, and crushed to obtain a starch-modified montmorillonite (St-La-Mont).

2.3. Test process

Weigh an appropriate amount of K₂Cr₂O₇ and dissolve it in a 1 L volumetric flask to prepare a solution with a concentration of 30 mg/L K₂Cr₂O₇ for use. The contaminated solution was shaken in a water bath thermostatic oscillator at 30°C for 8 hours (200 r/min), centrifuged and the supernatant was taken. The concentration of Cr(VI) was measured by UV-Vis spectrophotometer, and Cr was obtained according to formula (1). (VI) removal rate. Three groups of parallel samples were set up in the experiment and the average value was taken. The removal rate of Cr(VI) is shown in the following formula.

\[ \eta = \left( \frac{C_0 - C_e}{C_0} \right) \]  \hspace{1cm} (1)

In the formula: initial concentration of Co—Cr(VI), mg/L; Ce—concentration of Cr(VI) in solution after the adsorption reaches equilibrium, mg/L; \( \eta \)—Cr(VI) adsorption rate.

2.4. Sample Characterization

The spatial structure of the material was characterized by X-ray diffractometer (XRD), and the state between the material layers was analyzed. Fourier transform infrared spectroscopy (FTIR)
was used to characterize the types of surface functional groups and chemical bonds. Zeta potential was characterized by high-sensitivity Zeta potential and particle size analyzer (nano ZS), and the amount of charges on the surface before and after montmorillonite modification was analyzed. The morphology of montmorillonite before and after modification was observed by field emission scanning electron microscope (SEM).

3. Results and Discussion

3.1. X-ray diffraction characterization

The X-ray diffraction characterization of Mont, La-Mont and St-La-Mont is carried out, and the results are shown in Figure 1. According to the diffraction patterns, it can be seen that Mont and La-Mont have almost the same diffraction peaks around 25~30°, indicating that the two have a certain similarity in surface structure. The modification of La did not bring a new structure to the surface of the Mont crystal, and there was no obvious numerical difference in the diffraction 2θ angles of the three, indicating that La-modified Mont and St-modified La-Mont both changed the interlayer domain of the two. Mont and La-Mont have the same diffraction peak at 29.36° (Wang Xin et al., 2021), but St-La-Mont has no diffraction peak at this position, indicating that the modification of La-Mont by St changes Mont and La-Mont through surface wrapping. The diffraction peaks shift to high angles, indicating that St modification reduces the interlayer spacing of montmorillonite and reduces the thickness of the interlayer domain of Mont. It is speculated that it may be because St does not enter the interlayer domain of Mont, but wraps it outside it. the diffraction peak at 29.36° has a certain influence on the adsorption efficiency of Cr(VI). At the same time, according to the literature review, the loading of La in this study is not large, and it is difficult to capture the diffraction peaks of lanthanum at a scanning speed of 10°/min by XRD (Frois S R et al., 2012).

![Fig.1 XRD Phenogram](image)

3.2. Fourier transform infrared spectroscopy FTIR characterization

In order to determine whether there are chemical changes between montmorillonite and starch
during the modification process and whether the modification affects the surface functional groups of montmorillonite, Mont, La-Mont and St-La-Mont were characterized by FTIR, respectively as shown in picture 2. When the peak positions are 3412 cm\(^{-1}\) and 1478 cm\(^{-1}\), peaks appear in the Mont, La-Mont and St-La-Mont curves. In contrast, the peak of St-La-Mont is weakened, which may be caused by the stretching vibration of water molecules between La-Mont and St interlayers (O-H). At the same time, it can be seen that La-Mont has obvious vibration enhancement in the peak area at 1478 \(\text{cm}^{-1}\) compared with Mont, which indicates that La reacts to Mont mainly with O-H (Larraza I et al., 2012). When the wave number is 2897 \(\text{cm}^{-1}\), the three peaks also appear, which is due to the fact that both Mont and La-Mont contain a large number of CH\(_2\) structures, which are caused by the C-H symmetrical stretching vibration between the CH\(_2\) structures; When the peak position is 1526 \(\text{cm}^{-1}\), peaks appear in the curves of Mont, La-Mont and St-La-Mont, which are caused by the C-O stretching vibration. When the peak position is 876 \(\text{cm}^{-1}\) and 1200 \(\text{cm}^{-1}\), the common curves of Mont, La-Mont and St-La-Mont appear, which may be due to the fact that the three are modified based on Mont, so that their substances contain Si(Al)-O stretching vibration.

At the same time, it can be seen from the FTIR scan pictures of the three that there is not much difference in the infrared spectra of the three on the wave crest. The front and rear migrations are all within the error range, and no new characteristic spectrum is formed. If La enters the internal structure of Mont and participates in the skeleton vibration, the Si-O bond bending vibration of the modified La-Mont will become larger and the wave number will decrease. It can be seen from the figure that such a situation does not occur, indicating that the reaction with Cr(VI) is mainly due to the electrostatic adsorption between La on the surface of Mont (Yang-Min et al., 2018).

![Fig.2 FTIR spectra diagram of Mont, La-Mont, St-La-Mont](image)

**Fig.2** FTIR spectra diagram of Mont, La-Mont, St-La-Mont

### 3.3. Scanning Electron Microscope (SEM) Characterization

Through further scanning electron microscope image analysis of Mont, La-Mont and St-La-Mont, the combination of La and St with La-Mont was explored. It can be seen from the figure that the surface stacking phenomenon of the three shows the stacking difference of St-La-Mont, La-Mont and Mont from dense to scattered, It can be clearly seen that the particle size of Fig. 4 is smaller and the arrangement is neat and orderly, while the particle size of Fig. 6 is significantly larger and
denser, which indicates that La-Mont and St-La-Mont Some particles on the surface have agglomeration effect, and a certain degree of agglomeration occurs. La-Mont and St-La-Mont have obvious lateral synaptic structure compared with Mont, and the structure similar to lateral synapse with surrounding fine particles is La element (Sahu Sumanta et al., 2021). At the same time, it can be seen that after St modification of La-Mont, the La element distributed on the surface is obviously reduced, and replaced by uniformly distributed St particles. The FTIR scanning results show that La entering the surface of Mont is mainly loaded on its surface, During the modification process of La and Mont, the solution of Mont gradually formed metal oxides and hydroxides, and on the surface of these metal oxides, due to the coordinative unsaturation of its surface ions, it coordinates with water in an aqueous solution to form a hydroxylated surface. The surface hydroxyl groups can undergo proton migration in solution, showing amphoteric surface characteristics and corresponding charges. The modified Mont surface After covering the hydroxyl groups, it is easy to form surface coordination complexes with anions (You-Wei et al., 2017), which is also the reason why La modification greatly improves the adsorption effect of Cr(VI).

Fig.4 Structural equation modeling spectra diagram of Na-Mont(a)

Fig.5 Structural equation modeling spectra diagram of La-Mont(b)
3.4. Zeta potential characterization

In order to determine the change of the overall electrical properties of La and St before and after the modification process, the Zeta potential characterization of Mont, La-Mont and St-La-Mont was carried out respectively. As shown in Figure 7, the potential of Na-Mont is -21.2mV, the La-Mont potential is -12.7mV, and the St-La-Mon potential is -10.9mV. La and St are both positively charged, so the modification effectively increases the potential of Na-Mont and La-Mont, that is, reduces their overall electronegativity. It is worth noting that the mass ratio of St-modified Na-Mont in this paper is 1:10, which fails to change Na-Mont to electropositivity.

3.5. Adsorption of Cr(VI) by Na-Mont, St, St-Mont

Experiments were carried out according to the experimental procedure of 1.3, and finally the
adsorption results of Mont, La-Mont and St-La-Mont on 10 mg/L Cr(VI) were obtained, as shown in Figure 8. The research results show that Na-Mont has almost no adsorption effect on Cr(VI), La-Mont and St-La-Mont have good adsorption effect on Cr(VI) (between 50% and 75%). It can be seen from the FTIR analysis that Mont, La-Mont and St-La-Mont have similar peak positions in the 0-4000 cm\(^{-1}\) peak position. But Sem analysis shows that La-modified Mont surface is covered with hydroxyl groups, and it is easy to form surface coordination complexes with anions, which can be removed by reacting with negatively charged Cr\(_2\)O\(_7^{2-}\) in solution (Jianjun Yuan \textit{et al.}, 2014). It is also because the modification of La and Mont only stays on the surface of Mont, so after St loading La-Mont, St particles wrap the shell of La-Mont, which reduces the chance of direct contact with Cr\(_2\)O\(_7^{2-}\), so although St is further reduced the electrical properties of La-Mont, however, it reduces the effect of complex adsorption between La-Mont and Cr\(_2\)O\(_7^{2-}\), so its adsorption effect is 58.6% less than that of La-Mont, which is 73.9%, but it also reduces the negative charge of Mont, so its adsorption effect is higher than that of Mont 1.5%.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{Fig8.png}
\caption{Chromium(VI) adsorption effect of Mont, La-Mont, St-La-Mont}
\end{figure}

Combined with the Zeta potential characterization results of Mont, La-Mont and St-La-Mont on Cr(VI), it can be seen that Mont is electronegativity as a whole, while Cr(VI) exists in the form of Cr\(_2\)O\(_7^{2-}\) ions in aqueous solution, and is in the Coulomb repulsion force. Under the action of, there is almost no adsorption reaction between Mont and Cr(VI). The electronegativity of La-Mont is reduced due to the modification of La, so that its adsorption capacity for Cr(VI) has been significantly improved. At the same time, since the surface of La-modified Mont is covered with hydroxyl groups, it is easy to form with anions. Surface coordination complex, the removal rate reached 73.9%. At the same time, according to FTIR and SEM analysis, St stays on the surface of La-Mont and has surface stacking phenomenon, which further reduces the negative charge of St-La-Mont and increases its adsorption rate to 58.6%. The encapsulation effect masks the original diffraction peak of La-Mont at 29.36°, which reduces the effect of La modification, and the overall adsorption efficiency is also lower than 73.9% of La-Mont.

1. Conclusion

La-Mont and St-La-Mont composites were prepared in this study. La and St modification may
only exist on the surface of montmorillonite, which reduces the overall electronegativity of Mont. The change of the overall electrical properties of St-Mont makes St-Mont have the ability to adsorb Cr(VI). At the same time, after the surface of La-modified Mont is covered with hydroxyl groups, it is easy to form surface coordination complexes with anions, which further enhances its adsorption effect.

In this study, environmentally friendly and inexpensive montmorillonite and starch, which are ubiquitous in nature, were selected as the base material and modifier, respectively, to study the adsorption effect of Cr(VI) ions in water. The modification and its adsorption of heavy metals have theoretical significance, which provides a new research idea and direction for the adsorption and removal of Cr(VI) ions.

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