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Synthesis of human hair fiber-impregnated chitosan beads functionalized with citric acid for the adsorption of lysozyme

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Chitosan materials have been applied widely in protein separation and purification processes, however, pure chitosan materials generally suffer from inadequate mechanical properties and low densities of active adsorption sites, thereby leading to short service lives, limited actual re-usage counts and unexpected re-absorption capacities as adsorbents. Herein, a novel CA-CS/HHF composite was firstly constructed by impregnating human hair fiber into a CS matrix and then functionalized with citric acid. The structures and morphologies of the thus-produced composites were characterized by FTIR and SEM, respectively. A mechanical stability test showed that the impregnation of HHF into the chitosan matrix could effectively enhance the mechanical stability of the CA-CS/HHF beads in comparison with pure chitosan spheres. The adsorption of proteins and a protein mixture revealed that the product could adsorb positively charged proteins, as ascertained by the method of sodium dodecyl sulfate polyacrylamide gel electrophoresis. The maximum value for LYZ adsorption on the CA-CS/HHF composite was 42.1 mg g^{-1} with a fast equilibrium time of 3 h. Moreover, the resultant product exhibited good reversibility with respect to lysozyme. This simple and intriguing approach characterized by both environmental conservation and waste recycling strongly extends any future utilization of human hair fiber, and the resultant CA-CS/HHF can potentially be truly applied for the quick and effective adsorption of positively charged lysozyme.

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Introduction

In recent years, using absorptive approaches for the highly efficient purification of proteins has found widespread utilization in biochips and biosensors,1 foodstuffs,2 cosmetics,3 and the biomedical and pharmaceutical industries.^{4,5} For example, a great number of protein carrying adsorbents with remarkable selectivities, including nanosilica, TiO2 nanotubes, graphene, gra and carbon nanotubes,9 have been developed. It has been verified that these adsorbents possess high selectivities and binding capacities, but generally suffer from high production costs and difficulties in their preparation and separation.¹⁰ Alternatively, chitosan (CS), the second most abundant natural polymer after cellulose, has been proposed for applications in protein separation and purification processes because of its high affinity to proteins, good biocompatibility, and antibacterial and hydrophilic properties. 11-13 Several chitosan-based adsorbents including chitosan nanoparticles,14 chitosan films,15 chitosan microspheres,16 a magnetite/graphene oxide/ chitosan composite,17 and chitosan derivative beads18 have already been explored for protein adsorption applications. Chitosan beads are more extensively used than the film or powdered forms owing to their easy fabrication and scale-up, controllable particle size, excellent dispersion stability, and simple recovery process.19 Regardless of such interesting features, it has been found unfortunately that pure CS materials generally suffer from inadequate mechanical properties, thereby leading to short service lives, limited actual re-usage counts and unexpected re-absorption capacities as adsorbents.20 To address the abovementioned shortcomings, the impregnation of natural fibers, like silk fibers,21 rice straw fibers,22 cellulose fiber,23 and carbon fiber,24 into the CS matrix has been shown to act as an efficient way to achieve desirable chitosan beads with highly strengthened mechanical properties.

Human hair fiber (HHF), a nanocomposite biological fiber with a diameter of about $80~\mu m$, is a material considered useless in most societies and therefore the common management options for human hair include dumping or burning as waste, which are characterized by serious resource waste and significant environmental pollution. The chief constituent of human hair is mainly keratin protein, which comprises about 65–95%, whereas of the other components about 32% is water and lipid pigments. The existence of cortex keratin endows

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HHF with surprising strength and the rupture load of a healthy hair fiber ranges from 50 to 100 g.²⁵ By virtue of its high tensile strength, HHF has been used for reinforcing clay-based constructions, and it has been verified that the addition of hair significantly enhances the structural strength and prolongs the life of these clay structures.²⁸ More importantly, as a potential material resource, human hair possesses other supplementary advantages including being completely biodegradable, clearly renewable, naturally abundant and available in every locality. Considering these attractive characteristics, HHF holds great promise as a reinforcing agent to improve the mechanical properties of brittle chitosan bead materials, yet this has never been reported before.

Conventionally, neat chitosan materials usually do not have high adsorption capacities or efficiencies as a result of their relatively small surface areas and low levels of active anchoring sites. To improve their protein adsorption performance, further surface functionalization or modification of the chitosan materials is urgently needed. All in all, the fundamental protocol for the modification of chitosan can be attributed to hydroxyl and amidogen groups inherited from chitosan, which have provided a versatile platform for secondary reactions with other desired molecules. The main goals of such modifications of chitosan were mainly focused on the enhancement of its protein binding capacity. For instance, Bayramoğlu et al. grafted poly (methacrylic acid) onto chitosan beads using grafting copolymerization initiated by ammonium persulfate under a nitrogen atmosphere, and the resultant products possessed relatively high adsorption capacities toward lysozyme.29 Hua et al. modified chitosan with ethylenediaminetetraacetic acid and the obtained materials could specifically bind his-tagged proteins.30 Sun et al. synthesized carboxymethyl chitosan magnetic nanoparticles by a chemical coprecipitation method and then treated these with various ion solutions for lysozyme separation and purification.31 Recently, Ding et al. have demonstrated another fascinating protein adsorbent by in situ functionalizing electrospun ethylene-vinyl alcohol nanofibrous membranes with critic acid (CA).32 It has been verified that the CA performs well as a natural grafting agent since it can introduce substantial carboxyl functional groups to the CS/HHF matrix, significantly improving the availability of active adsorption sites. Simultaneously, the relatively long carbon chains of CA can also serve as spacer arms to weaken the steric hindrance between the adsorption sites and target molecules, further facilitating the protein binding process.33 Such modification methods render chitosan adsorbents with improved adsorption performance, and consequently provide a new kind of functional material for the fast, effective, and low cost production of various highly purified protein products.

Thus, in the present paper, novel CA-CS/HHF composite beads were fabricated by impregnating HHF into a CS matrix as a reinforcing filler through an instantaneous gelation method and subsequent heat treatment to graft citric acid molecules under mild conditions. Benefiting from the tough and inexpensive HHF filler and the abundant carboxyl groups of CA, the resultant CA-CS/HHF composite beads exhibit the integrated properties of a high number of active sites and robust

mechanical strength. Such fantastic advantages endow the asprepared CA-CS/HHF with brilliant adsorption performance, including efficient separation, a high adsorptivity towards lysozyme (LYZ), a fast adsorption equilibrium, and good reversibility, implying that this material could potentially be truly used in protein adsorption fields.

Experimental

2.1. Materials

Human hair fibers (HHF) were collected from a barbershop (Shaanxi, China). Chitosan (CS) with a molecular weight of 200 000 and an 88% degree of deacetylation was supplied by Tianjin Chemical Reagent Factory (Tianjin, China). Before use, the raw human hair was cleaned three times with distilled water, dried in an oven at 60 °C for 24 h, and finally cut into pieces with lengths of less than 1 mm. Citric acid, sodium hydroxide, acetic acid and acetone were supplied by Tianjin Chemical Reagent Factory (Tianjin, China). All the chemicals used in the present study were of reagent grade and were used without further purification.

2.2. Preparation of the CA-CS/HHF composite beads

Typically, 0.2 g of chitosan was dissolved in 10 mL of 2% (v/v) acetic acid solution, then a specific amount of human hair fiber (0-50%) was immersed in the solution. After magnetic stirring at 250 rpm for 20 min at room temperature, a homogenously dispersed chitosan-HHF dispersion was achieved and dropwise added into a precipitation bath containing a 5 wt% NaOH solution through a 0.8 mm gauge syringe needle, and subsequently chitosan/human hair fiber (CS/HHF) beads were obtained. Blank CS beads were obtained by a similar preparation process without adding HHF. The beads were soaked in deionized water for 24 h and then rinsed with deionized water five times. The immobilization of citric acid on the CS/HHF intermediates' surfaces was performed according to the method reported in previous work with some modification.34 The prepared CS/HHF beads (1 g) were dispersed in methanol solutions with various citric acid concentrations (0, 0.1, 0.2, 0.3, 0.4, 0.5 and 0.6 mol L^{-1}) at 50 °C for a certain time. Next, to completely eliminate excess citric acid, the beads were washed with deionized water and the residues were retained in an aqueous environment for future use.

2.3. Testing mechanical performance

Compression strength testing is an easily performed and visualized way to evaluate the mechanical strength of a bead. The prepared samples were analyzed by an Instron 3345 materials testing machine with a 10 N force transducer. A wet bead was placed onto a flat platform and compressed to a deformation of 40% by a flat-tip probe at a speed of 0.5 mm min $^{-1}$. Five beads from each sample were compressed in order to provide statistically representative measurements. Applied force (F) versus displacement (H) data was generated from the compression measurements, and was employed to determine the Young's modulus (E) according to the Hertz model.³⁵

$$F = \frac{4R^{1/2}}{3} \frac{E}{1 - v^2} \left(\frac{H}{2}\right)^{3/2} \tag{1}$$

where R is the radius of a bead, and ν (taken as 0.5) is the Poisson ratio.36

2.4. Determination of carboxyl content

The carboxyl content of the CA-CS/HHF composite beads was determined by using an acid-base titration method reported by Salam.37 1.0 g of the obtained sample was firstly dissolved in excess 0.1 mol L^{-1} NaOH (pH 12.5) for 1 h, and then 0.1 mol L^{-1} HCl was added to the mixture using phenolphthalein as an indicator. The carboxyl content (meq./100 g) in milliequivalents of acidity per 100 g is obtained by:

Carboxyl content =
$$\frac{(V_b - V_a)}{M} \times C_{HCl} \times 100$$
 (2)

where V_a and V_b (mL) represent the volume of HCl used to titrate in the presence and in the absence of the product, respectively. $C_{\rm HCl}$ is the normality of the used HCl, and M(g) is the mass of the dry specimen.

2.5. Testing adsorption performance

To investigate the static adsorption performance of the CA-CS/ HHF composite beads toward different types of separate proteins, 0.5 g of freeze-dried samples were soaked in 100 mL of protein solution (1 g L⁻¹) for a certain time; subsequently the absorbance intensity change was detected using a UV-visible spectrophotometer to determine the protein concentration. The amount of adsorbed protein $q_t \text{ (mg g}^{-1})$ at various times (t) was calculated by the following (eqn (3)):

$$q_t = \frac{(C_0 - C_t)V}{m} \tag{3}$$

where m is the amount of sample (g), V is the volume of the protein solution (L), and C_0 and C_t are the initial concentration of the protein solution and its concentration at time t (g L⁻¹). Additionally, the method of sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE)38 was employed to analyze the selective adsorption ability of CA-CS/HHF composite beads toward a protein mixture (containing LYZ, bromelain, BSA, albumin, and pepsin).

To test the reversible adsorption property of the as-prepared CA-CS/HHF composite beads, consecutive adsorption-desorption experiments were conducted with 0.5 M NaCl for ten cycles. Briefly, the LYZ-loaded beads were immersed in 50 mL of NaCl solution for a period of 20 min to elute the adsorbed protein, then the composites were rinsed with fresh deionized water 3 times to remove the NaCl. After that, the regenerated composite beads were dipped into the lysozyme solution to reuse for another cycle.

2.6. Characterization

The functional groups of the obtained products and human hair were confirmed using a Perkin Elmer FTIR System 2000 in the 600-4000 cm⁻¹ range at a resolution of 4 cm⁻¹ using the KBr wafer technique. A scanning electron microscope (JSM-6490LV, and 10 kV) was employed to observe the morphological changes of the obtained products and human hair. Before SEM analysis, the obtained beads were dried via lyophilisation and the human hair was dried at 60 °C, then all the samples were coated with gold using an Emitech K 550 Coater. The concentrations of the protein solutions were detected using a Shimadzu-18A UVvisible spectrophotometer. Mechanical stability testing was conducted by an Ultrasonic cell disrupter (KH-160TDV) with frequency 20 kHz and power 100 W.

3. Results and discussion

3.1. Synthesis and characterization of CA-CS/HHF composite beads

The detailed functionalization and fabrication processes of the CA-CS/HHF composite beads are schematically illustrated in Scheme 1.

According to previous reports, the incorporation of natural fibers is an effective and frequently used approach to reinforce the physico-chemical properties of a chitosan matrix.21-24 In the current study, the interactions between the HHF fillers and the chitosan matrix may include covalent and non-covalent interactions such as hydrogen bond interactions and electrostatic interactions. These can be observed during the synthesis by the apparent change of the morphology of the chitosan-HHF dispersion from a suspension to beads when drops of the mixture were neutralized and gelated in an alkaline solution. In other words, the HHF filler has been successfully buried in the CS matrix, and millimetric CS/HHF hybrid beads with free -NH₂ and -OH groups were constructed by the dropping and pHprecipitation method. These groups are of great importance since they can afford secondary reactions with many citric acid molecules. In the subsequent citric acid immobilization process, lots of free CA molecules were anchored through hydrogen bond linkages on the surfaces of the CS/HHF beads, which contain numerous hydroxyl (-OH) and amino groups (-NH₂). Under the subsequent heating conditions at 50 °C, two adjacent carboxylic acid groups of CA were easily dehydrated and lost one water molecule to yield a cyclic anhydride.39 The reactive anhydrides subsequently reacted with other hydroxyl or amino groups on the chitosan surface. Hereby, the overall surfaces of the CS/HHF beads were decorated with lots of citric acid molecules and numerous carboxylic groups were introduced for binding proteins.

From the above analysis, it can be supposed that the chitosan, citric acid and human hair fiber have made their own extraordinarily important contributions to the formation of the CA-CS/HHF hybrid beads. Namely, (1) for the chitosan component, the native entanglement or crosslinking properties of the chitosan polymer chains help the discrete HHFs unite together, forming uniform spherical structures, which greatly simplifies the isolation procedure and speeds up the assay process. Also, the inherent and ample hydroxyl and amidogen groups on the CS substrates can be used for further derivatization, making the grafting reaction of the CA molecules occur facilely on the surface of CS/HHF. (2) As with classical fillers, the utilization of

Scheme 1 Proposed mechanism for the formation of CA-CS/HHF hybrid beads.

HHF as an inner skeleton can also strengthen the mechanical stability of CA-CS/HHF composite beads. Of course, it is obvious that the partial replacement of the internal parts of the CS beads with HHF fillers will have avoided unnecessary waste of the chitosan resource and reduced the production costs of these absorbent materials greatly. (3) The CA molecules used as modifiers in this research are natural and environmentally friendly. They offer a large number of active sites (carboxyl groups) for binding protein molecules and decrease the steric hindrance between the adsorption sites, and consequently help the CA-CS/HHF composite beads capture proteins easily, which significantly improves the adsorption performance of CA-CS/ HHF composite beads. From these points of view, this intriguing and economical strategy opens a suitable route to prepare a protein adsorbent with an improved absorption capacity and mechanical strength by using the abundance, biodegradability and renewability traits of CS and HHF.

To analyse structure and surface composition changes, the valuable and well established testing technique of FTIR spectroscopy was employed and the spectra of the intermediates and the final product are exhibited in Fig. 1.

For HHF, the broad and strong peak around 3425 cm⁻¹ corresponds to –OH and –NH stretching vibrations, and the medium absorption peak at 2924 cm⁻¹ represents the –CH stretching of the methylene groups in human hair. The absorption bands at 1644, 1544, 1157 and 1070 cm⁻¹ are assigned to C=O stretching (amide I), –NH bending (amide II), antisymmetric stretching vibrations of C–O–C bridges, and skeletal vibrations involving C–O bonds stretching separately, which are the characteristic absorption peaks of human hair.⁴⁰ As for chitosan, the peak at 3425 cm⁻¹ is attributed to the stretching of free amine (–NH₂) and hydroxyl (–OH) functional groups in the ring of CS. The peaks at 1544 and 1424 cm⁻¹ are related to –NH₂ and –OH bending vibrations respectively, suggesting the existence of amido and hydroxyl groups. In the IR

spectrum of the CS/HHF intermediate, the above characteristic absorption peaks belonging to HHF and CS were almost retained. The above results provide preliminary evidence that HHF and chitosan were intertwined and that together they formed a stable CS/HHF hybrid bead, which can be further confirmed by the subsequent scanning electron microscopy (SEM) step. Following the subsequent reaction between the CS/HHF beads and CA, the final product shows some changes in its absorption peaks. For instance, the absorption band near 1544 cm⁻¹, corresponding to the bending vibrations of -NH₂ in the CS ring, vanished in the CA-CS/HHF spectra, suggesting that some of the amine groups in the CS/HHF beads had already reacted with the -COOH groups of CA. Moreover, it can be distinctly observed that two intense peaks appear at 1257 and 920 cm⁻¹ in (d) which are ascribed to the C=O stretching

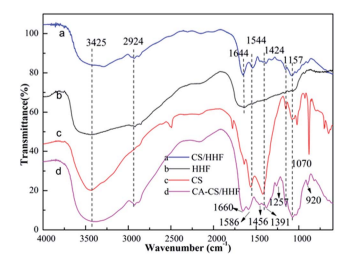


Fig. 1 FTIR spectra of HHF fillers (b), chitosan (c), CS/HHF beads (a), and CA-CS/HHF composite beads (d).

vibrations of carboxyl groups and -OH bending vibrations in COOH, further demonstrating the introduction of additional carboxyl groups onto the CS/HHF surface during the modification with CA.41 Besides this, the peak at 1660 cm⁻¹ is assigned to carboxamide, caused by the superposition of the C=O in amide I and C=O in COOH, and the bands at 1586, 1456 and 1391 cm⁻¹ are due to C=O asymmetric and symmetric stretching in a carboxylate anion. 42 Comparing the FT-IR spectra of CS/HHF and CA-CS/HHF suggested that the CA was anchored on the surface of the CA-CS/HHF hybrid beads.

The incorporation of HHF into the CS matrix can be further verified from photographs (Fig. 2a) and the influence of the HHF filler on the microstructure of the dried CA-CS/HHF beads is observed in the SEM images (Fig. 2b-d). Fig. 2a shows the appearance of prepared wet CA-CS/HHF composite beads with different weight ratios of HHF. As can be seen, with a larger proportion of HHF, the spherical beads, with diameters of 3.0 mm, have a more significant black color in the wet state. This appearance change provided further support that CS was embedded with HHF. In marked contrast with the flat surface of neat CS (the inset image of Fig. 2b), the pictured CA-CS/HHF bead was highly rugged with abundant papillae and interstitial spaces (Fig. 2b). Obviously, the flat surface of the CS was destroyed by the introduction of HHF, which has a regular rodlike structure (Fig. 2d). Fig. 2c is the cross section of a CA-CS/ HHF bead; it is obvious that lots of rod-like HHF was implanted into the interior of the pure CS bead, giving further evidence that the internal linkage between HHF and CS provides superior mechanical strength. Moreover, there are also some HHF granules at the outer surface of CA-CS/HHF, indicating that the HHF fillers are immobilized fully by CS beads inside and outside. In analogy, the incorporation of HHF could contribute to improving the mechanical properties of CA-CS/HHF hybrid beads and the unique surface morphology of CA-CS/HHF, with large quantities of carboxylic groups, is favorable for providing sufficient space for protein adsorption.

3.2. Enhanced mechanical stability of CA-CS/HHF composite beads

Since favorable mechanical properties can endow protein adsorbents with long service lives and good endurance under high pressure, it is extremely important to improve the mechanical properties of the CA-CS/HHF composite beads by impregnating HHF into the CS matrix. Good mechanical properties have been previously identified as being correlated with the presence of well-distributed fibers or nanomaterials in the chitosan matrix.43 Thus, in order to investigate the effect of HHF fillers on the mechanical properties of CA-CS/HHF beads, a series of hybrid beads with various HHF contents (0-50%) were studied using compression testing (Fig. 3).

Fig. 3a reveals that the applied force values for CA-CS/HHF hybrid beads were generally higher than those for pure CS beads and presented an apparent increasing trend upon increasing the weight ratio of HHF and CS from 10 to 50%, and these values varied from 0.39 to 0.84 N with 40% deformation. Moreover, the variation in the force values between the 40 and 50% beads was small (<0.05 N), indicating that CA-CS/HHF

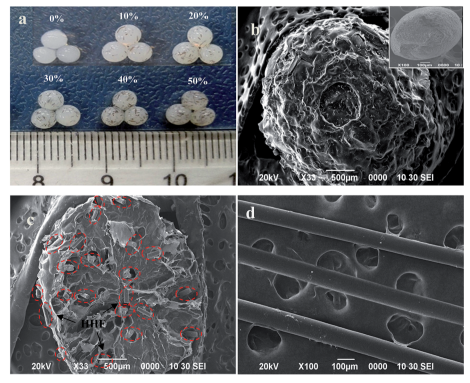
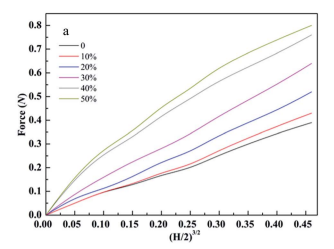


Fig. 2 Appearance of CA-CS/HHF hybrid beads with different HHF content (a), and SEM micrographs of CA-CS/HHF and CS beads (b), the cross section of CA-CS/HHF (c) and HHF (d)

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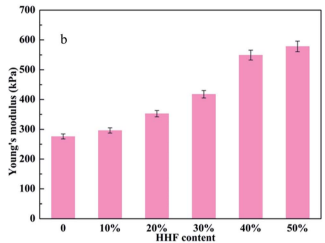


Fig. 3 (a) The force (N) and $(H/2)^{3/2}$ curves of CA-CS/HHF composite beads with 40% final deformation; H is the displacement. (b) Effect of the HHF content on the Young's modulus of CA-CS/HHF composite beads

hybrid beads with high mechanical performance can be obtained by the incorporation of 40% HHF into the CS matrix. Since the space within the CA-CS/HHF composite beads was occupied by the tough HHF, the composite beads became more rigid compared with the pure CS beads, and consequently beads with a higher feed ratio of HHF fillers required a larger force to produce a given degree of deformation. Additionally, the stiff HHF fillers impeded the movement of the CS chains necessary for shape change under external stress, also resulting in a larger force being required to deform the CA-CS/HHF composite beads. Most importantly, the HHFs dispersed within the hybrid beads could form a string-like matrix owing to their slenderness. Hence, compared to CS beads, the CA-CS/HHF hybrid beads could transfer compression stress more evenly from the soft chitosan matrix phase to the rigid HHF filler, resulting in enhanced mechanical strength.44 Hence, the mechanical strength of the CA-CS/HHF hybrid beads can be enhanced by simply impregnating HHF into the CS matrix. Moreover, judging from the relationships between the Young's modulus and HHF content, the Young's moduli of the CA-CS/HHF hybrid

beads were significantly higher than that of the control sample (CS beads) and increased with an increasing feeding ratio of HHF (Fig. 3b). Specifically, the Young's moduli of the CA-CS/ HHF beads were 298 \pm 8 kPa, 352 \pm 10 kPa, 417 \pm 12 kPa, 549 ± 16 kPa, and 578 ± 17 kPa at ratios of 10%, 20%, 30%, 40% and 50%, respectively, whereas the corresponding value for neat CS beads was only 276 \pm 8 kPa. For comparison, Eng Seng Chan et al. verified that the Young's modulus of alginate beads increased from 430 \pm 5 kPa to 582 \pm 7 kPa after embedding 100 g L⁻¹ of halloysite nanotubes. 45 All of these results indicate that it is more effective to improve the mechanical stability of the CA-CS/HHF hybrid beads by embedding HHF into the CS matrix, and the weight ratio of 40% was selected to perform the subsequent protein adsorption experiments.

3.3. Optimizing protein adsorption performance of CA-CS/ **HHF** composite beads

The adsorption capability of CA-CS/HHF composite beads is mainly dependent on electrostatic attraction between ionized carboxyl groups on the CA-CS/HHF hybrid beads and the positively charged residues on proteins; that is, the adsorption performance of composite beads can be significantly improved as the amount of carboxyl groups on CA-CS/HHF increases. Thus, it was necessary to determine the number of carboxyl groups introduced on CA-CS/HHF composite beads and to optimize lysozyme adsorption by adjusting the concentration of CA (from 0.1 to 0.6 mol L^{-1}). The experimental results are depicted in Fig. 4.

As Fig. 4 shows, the carboxyl content of CA-CS/HHF composite beads was dramatically increased in comparison with that of native CS/HHF. Specifically, the carboxyl content increased from 68 meq./100 g to 407 meq./100 g upon increasing the CA concentration from 0 to 0.4 mol L⁻¹, signifying that increasing numbers of CA molecules were fixed on CA-CS/HHF. Above this CA concentration range, however, the carboxyl content negligibly changed, indicating that 0.4 mol L^{-1} of CA was sufficient to saturate the carboxyl groups on the composite beads. Correspondingly, the adsorption capacities of

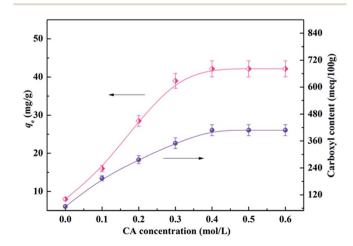


Fig. 4 Effect of the CA concentration on the carboxyl content and adsorption capacity of CA-CS/HHF composite beads.

experiments.

the CA-CS/HHF composite beads present a typical sustained increasing trend upon increasing the CA concentration, and the maximum capacity was obtained with a concentration of 0.4 mol L⁻¹. A further increase in the CA concentration did not result in an enhancement of the adsorption behavior, demonstrating that the adsorption process had reached an equilibrium state because of the limited number of carboxyl groups on the composite beads' surfaces. It is obvious that the changes in the lysozyme adsorption capacity are consistent with those of the carboxyl group content. Conforming to the maximum

lysozyme adsorption capacity, CA-CS/HHF composites modified with $0.4 \text{ mol } L^{-1}$ CA were employed to perform the following

3.4. Tunable protein adsorption study of CA-CS/HHF composite beads

Considering that the electrical charge on the proteins and CA-CS/HHF composite beads could be tuned by the ionization of acidic or basic groups associated with the proteins and composite, the pH value of a protein solution plays a critical role in testing the adsorption capacity of charged CA-CS/HHF composite beads. As the solution pH is changed, the ionized acidic or basic groups contribute to attractive or repulsive interactions between the protein and the composite. 46,47 To study the adsorption of protein as a function of the solution pH, positively charged lysozyme (LYZ) was selected as a model protein. The adsorption of LYZ (at a concentration of 1 g L⁻¹) at different pH values was performed and the obtained values are illustrated in Fig. 5a.

As can also be seen, LYZ absorption on the samples is low under acidic conditions, but sharply increases and almost reaches equilibrium in the pH range from 7 to 8. This phenomenon can be attributed to the abundant carboxyl groups on the surface of the CA-CS/HHF composite beads. In theory, most of the -COO groups fixed on the structures of the CA-CS/HHF materials could combine with hydrogen ions (H⁺) in acidic media to form -COOH groups, decreasing the negative surface charges of the samples. With the reduction of the -COOgroups, the electrostatic force between the positively charged LYZ (LYZ⁺) and anionic groups would be weakened; consequently, the adsorption capacity was limited. The electrostatic attraction between lysozyme and CA-CS/HHF strengthened when the pH value was increased, since the number of negatively charged -COO groups increased, promoting the binding of more LYZ onto the CA-CS/HHF composite beads. When the LYZ solution pH was higher than 8, the observed adsorption of LYZ on the CA-CS/HHF composite beads gradually reduced. This phenomenon can be understood considering the fact that a positively charged self rejecting layer of LYZ molecules was formed on the CA-CS/HHF surface via electrostatic attraction between the positively charged LYZ (LYZ⁺) and the negatively charged composite surface. 48,49 Long range exclusion, by a positively charged self rejecting layer of LYZ, of similarly charged LYZ molecules in solution could be responsible for the further decline in the adsorption capacity. Therefore, the CA-CS/HHF composite beads could be used to realize the

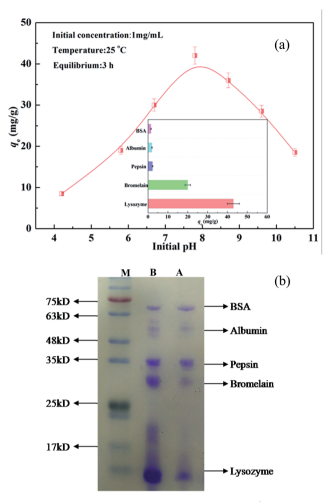


Fig. 5 (a) The observed adsorption of LYZ (1 mg mL⁻¹) onto CA-CS/HHF composite beads. Inset image: adsorption capacities toward different types of separate proteins. (b) SDS-PAGE analysis of the purification of a protein mixture: M, protein marker; lanes B and A, protein mixture before and after adsorption on CA-CS/HHF hybrid beads.

selective adsorption of positively charged proteins by regulating and controlling the buffer pH values. Given that protein solutions with near neutral pH values are more stable and more appropriate in practical applications, a pH value of 7.8 was selected to conduct the subsequent tests.

From the inset image in Fig. 5a, it is apparent that the adsorption of positively charged proteins (LYZ, bromelain) was much greater when compared to that of other proteins (BSA, albumin, pepsin), and this result is connected with the surface charge characteristics of the experimental subject. The composites' adsorption of positively charged proteins was mainly controlled by electrostatic forces between the positive charges on the LYZ and bromelain and the negative charge on the composite, and therefore they are attracted each other, indicating that more positively charged proteins than negatively charged molecules can make contact with a composite with the same surface area. Hence, the variation in the adsorption capacity towards differently charged proteins suggests that the

resultant materials can be a potential candidate for the easy adsorption of positively charged proteins. Moreover, the adsorption ability of the obtained CA-CS/HHF composite beads toward a protein mixture was also assessed at a buffer pH of 7.8, as visually illustrated in Fig. 5b. Apparently, the bands for lysozyme and bromelain presented color fading after their adsorption by the CA-CS/HHF composite beads. Conversely, the change of the band colors for BSA, albumin, and pepsin can be hardly observed, since the proteins have the same surface charges as the carboxyl groups on the CA-CS/HHF composite beads. Accordingly, the CA-CS/HHF composite beads could be used an intriguing tool for the adsorption of positively charged proteins by adjusting and controlling the pH of a protein mixture.

3.5. Adsorption kinetics

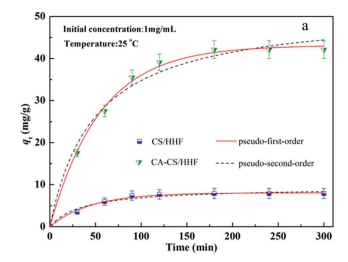
Fast adsorption kinetics are one of most crucial factors for protein adsorbents since they have a direct influence on the actual efficiency of the adsorbents. The existence of citric acid molecules on the surfaces of the CA-CS/HHF composite beads improves the binding capacity towards positively charged lysozyme, which can be verified by comparing the adsorption capacities of CA-CS/HHF and CS/HHF beads. Therefore, the adsorption kinetics performances of CA-CS/HHF and CS/HHF composite beads were investigated by the determination of the residence time needed for the completion of the lysozyme adsorption process, as shown in Fig. 6. It can be easily observed that the adsorption by CA-CS/HHF abruptly increased during the initial stage and the increasing trend did not stop until a state of equilibrium was achieved after approximately 3 h, indicating the high adsorption efficiency of CA-CS/HHF for lysozyme molecules. Furthermore the adsorption capacity of CA-CS/HHF composite beads is up to 42.1 mg g⁻¹ while the CS/ HHF beads without citric acid decoration have a lower binding capacity (7.9 mg g⁻¹). The enhanced adsorption capacity is attributed to the considerable availability of carboxyl groups derived from the citric acid layer on the CA-CS/HHF composite bead's surface, which provides numerous external adsorption sites for effectively binding lysozyme molecules through the electrostatic attraction between anionic CA-CS/HHF and cationic lysozyme.51

Kinetics is one of the most fascinating characteristics responsible for the efficiency of absorption. Significantly, the experimental data at 25 °C were fitted by pseudo-first-order and pseudo-second-order kinetic models, as shown in Fig. 6. These two kinetic models were defined by the following formulas:^{52,53}

$$\ln(q_{\rm e} - q_t) = \ln q_{\rm e} - k_1 t \tag{4}$$

$$\frac{t}{q_t} = \frac{1}{k_2 q_e^2} + \frac{t}{q_e} \tag{5}$$

where q_e and q_t (mg g⁻¹) represent the adsorption capacities at equilibrium and at any time (min), respectively. k_1 (min⁻¹) and k_2 (g g⁻¹ min⁻¹) are the rate constants and can be evaluated from the intercepts and slopes of linear plots of $\ln(q_e - q_t)$ versus t and t/q_t versus t, respectively. The corresponding kinetic parameters including k_1 , k_2 , correlation coefficients (R^2), and



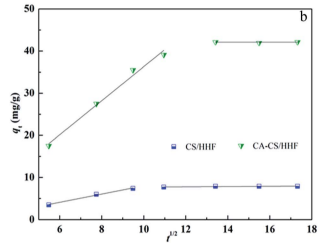


Fig. 6 (a) Adsorption kinetics of both CS/HHF and CA-CS/HHF samples in 1 mg mL $^{-1}$ lysozyme solution; (b) intraparticle diffusion plots.

calculated $q_{\rm e,cal}$ values are determined by a non-linear fitting method, as tabulated in Table 1. For the CS/HHF and CA-CS/HHF samples, the R^2 values for the pseudo-first-order model

Table 1 The kinetic parameters and coefficients of LYZ adsorption onto CS/HHF and CA-CS/HHF composite beads

	Kinetic constant	Samples	
Kinetics type		CS/HHF	CA-CS/HHF
Pseudo-first-order	$q_{\mathrm{e,exp}} \left(\mathrm{g} \; \mathrm{g}^{-1} \right)$	7.91	42.10
	$q_{\rm e,cal} (g g^{-1})$		
		8.077	43.10
	$k_1 (\mathrm{min}^{-1})$	0.022	0.018
	R^2	0.9911	0.9969
	χ^2	0.073	0.7096
Pseudo-second-order	$q_{\rm e,cal} ({\rm g g^{-1}})$	9.38	51.59
	$q_{ m e,cal}~({ m g~g^{-1}}) \ k_2~({ m g~g^{-1}~min^{-1}})$	0.0029	0.004
	R^2	0.9829	0.9672
	χ^2	3.929	0.2708
Intraparticle diffusion	$K_{\mathrm{id},1}$	0.9791	4.041
	$K_{\mathrm{id,2}}$	0.0297	0.012
	C	7.426	42.11
	R^2	0.9831	0.9854
	χ^2	3.082	0.9297

(0.9911 and 0.9969, respectively) are obviously higher than those for the pseudo-second-order model (0.9829 and 0.9672, respectively). Additionally, the theoretical values $(q_{e,cal})$ obtained from the pseudo-second-order model seemed to be

clearly higher than the experimentally observed data $(q_{e,exp})$, whereas the values from the pseudo-first-order kinetic model were quite consistent with the experimental values. This signifies that the swelling process could best be described by the pseudo-first-order model.

The steps of the adsorption process are analyzed thoroughly with an intraparticle diffusion model which is represented in egn (6).54

$$q_t = K_{\rm id} t^{1/2} + C {6}$$

where k_{id} (mg g⁻¹ min^{-1/2}) is the intraparticle diffusion rate constant and C (mg g^{-1}) is the intercept. The values of the characteristic constants k_{id} and C were obtained from the slope and intercept, respectively, of a plot of q_t versus $t^{1/2}$, and the results are displayed in Fig. 6b and Table 1. In the present case, for CS/HHF and CA-CS/HHF, the data points are related by two straight lines during the LYZ adsorption process. The first one represents the immediate occupation of the most readily available adsorption sites on the surfaces of the CA-CS/HHF composite beads. The second adsorption step may be attributed to very slow diffusion of the protein from the external surfaces into the inner pores. Furthermore, a fit of q_t versus $t^{1/2}$ is found to be linear, but the plots have nonzero intercepts. This phenomenon indicates that the adsorption process may be of a complex nature involving both surface adsorption and intraparticle diffusion, but that these are not the only controlling factors in determining the kinetics of the process.55,56

3.6. Adsorption isotherms

Adsorption isotherms are important for describing the adsorption behavior and are also important for providing information about the surface properties of the adsorbent and the design of adsorption systems. Here, commonly employed models including the Langmuir, Freundlich and Temkin isotherm models were developed to test the adsorption data. These models can be described in nonlinear form as follows:57-59

Langmuir isotherm:

$$q_{\rm e} = \frac{q_{\rm m} C_{\rm e} K_{\rm L}}{1 + C_{\rm e} K_{\rm I}} \tag{7}$$

Freundlich isotherm:

$$q_{\rm e} = K_{\rm F} C_{\rm e}^{1/n} \tag{8}$$

Temkin isotherm:

$$q_{\rm e} = A \ln K_{\rm T} C_{\rm e} \tag{9}$$

where $q_{\rm e}$ and $q_{\rm m}$ represent the equilibrium and theoretical maximum adsorption amounts (mg g^{-1}), respectively, and C_e is the equilibrium dye concentration (mg mL⁻¹). K_L , K_F and K_T refer to the constants of the Langmuir, Freundlich and Temkin models, respectively. The parameters n and A are the Freundlich and Temkin coefficients related to the adsorption intensity and surface heterogeneity of the adsorbent, respectively. The values of the parameters as well as the regression coefficients (R^2) can be determined from the above three isotherm equations and are displayed in Table 2. The R^2 value (0.9872) for the Langmuir model is much larger than those for the Freundlich model (0.9465) and the Temkin model (0.9741), reflecting that the adsorption process better fits the Langmuir model, and that the adsorption of lysozyme molecules on the CA-CS/HHF adsorbent took the form of monolayer adsorption due to the existence of strong electrostatic attractions between the carboxylic ion groups of the adsorbent and the lysozyme molecules. This result can be further reinforced by the comparison of the experimental points and fitted isotherm plots in Fig. 7.

To evaluate the feasibility of adsorption on the adsorbent, a separation factor known as the equilibrium parameter $(R_{\rm L})$ can be calculated by means of the Langmuir constant, K_L :

$$R_{\rm L} = \frac{1}{1 + C_0 K_{\rm L}} \tag{10}$$

where C_0 is the initial lysozyme concentration (mg L⁻¹). The values of R_L give information about the Langmuir adsorption process, be it irreversible ($R_L = 0$), linear ($R_L = 1$), favorable (0 < $R_{\rm L} < 1$) or unfavorable $(R_{\rm L} > 1)$. 60,61 $R_{\rm L}$ values for the adsorption of lysozyme are in the range of 0.223-0.801, at 298.15 K. This confirms that the lysozyme sorption onto CA-CS/HHF is favorable and therefore CA-CS/HHF can be referred to as a proper adsorbent for protein separation.

3.7. Reversibility of the CA-CS/HHF composite

From the perspective of economic plausibility, regeneration is an environmentally acceptable option which may be critical to decreasing process costs and providing the possibility of recovering the protein adsorbed from the solution. Generally, it is highly challenging but desirable to preserve the high adsorption capacity of such adsorption systems after repeating many cycles of adsorption and regeneration. Very recently, Ma and his colleagues described a NaCl treatment procedure for the regeneration of maleic anhydride-functionalized materials.62 The authors found that proteins could be directly

Table 2 Adsorption isotherm parameters for LYZ adsorption at 25 °C

•	·	
Models	Model parameters	R^2
Langmuir isotherm	$q_{ m max} = 56.12 \ { m mg \ g^{-1}}$ $K_{ m L} = 2.484 \ { m L \ g^{-1}}$ $R_{ m L} = 0.223 { m -} 0.801$ $\chi^2 = 3.054$	0.9872
Freundlich isotherm	n = 2.416 $K_{\rm F} = 39.00 \text{ L g}^{-1}$	0.9465
Temkin isotherm	$\chi^2 = 10.35$ $A = 13.34$ $K_{\rm T} = 19.69 \; {\rm L \; g^{-1}}$ $\chi^2 = 3.783$	0.9741

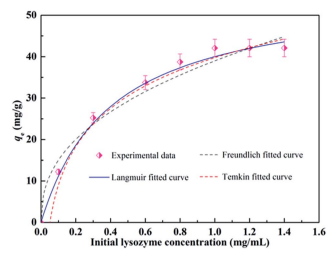


Fig. 7 Isotherm plots for the adsorption of LYZ onto CA-CS/HHF composite beads at 25 $^{\circ}\text{C}.$

eluted from the carbonyl group functionalized membranes, without the need for any other chemical reagents. Following a similar procedure, a NaCl solution was employed to elute the adsorbed lysozyme and facilitate the regeneration of the CA-CS/HHF composite for another cycle of adsorption. The adsorption capacity of the regenerated CA-CS/HHF is shown in Fig. 8.

As shown in Fig. 8, the adsorption capacity of the CA-CS/HHF composite beads was found to decline from 42.1 mg g $^{-1}$ to 38.4 mg g $^{-1}$ during 10 regeneration/reuse cycles, suggesting the significant chemical removal of binding lysozyme and the formation of activated CA-CS/HHF composite beads. Upon increasing the number of reuse cycles up to 10, no prominent difference in the lysozyme-absorbing ability could be observed between any pair of cycles, confirming that the CA-CS/HHF composite beads still retained a high adsorption activity toward lysozyme and can be effectively regenerated and

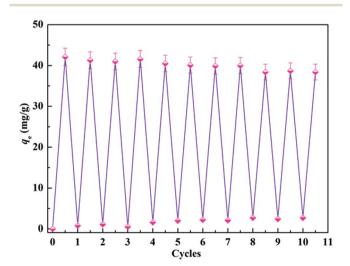


Fig. 8 Change in the adsorption capacity of CA-CS/HHF materials upon increasing the cycle number up to 10.

successively reused. Two major advantages of the CA-CS/HHF composite beads can explain this good reversibility. First, the robust and strong HHF fillers enhance the mechanical stability of the CA-CS/HHF composite beads, avoiding severe fracture during the adsorption–desorption process, and thus effectively prevent the CA molecules from shedding upon regeneration. Second, the large number of active functional groups (COOH) inherited from CA are not destroyed and are preserved after rinsing with natural salt solution repeatedly, endowing the regenerated composite with good adsorption performance. The excellent reversibility of the CA-CS/HHF composite beads can, significantly, achieve the demands of long service life for their practical application.

4. Conclusion

In conclusion, the present research was focused on developing a facile and eco-friendly strategy for the fabrication of novel carboxyl group functionalized composite beads (CA-CS/HHF) via the thermo-chemical modification of encapsulated coated human hair fiber (HHF) with citric acid. HHF and citric acid were introduced into the formulations as a reinforcing filler and protein catcher, respectively. SEM and FTIR analyses confirmed that HHF fillers were implanted into the interior of the chitosan beads and that abundant carboxylic groups derived from the citric acid are immobilized on the outer surface of CA-CS/HHF. The mechanical tests showed that the impregnation of HHF into the chitosan matrix with various weight ratios could effectively enhance the mechanical stability of the CA-CS/HHF composite beads in comparison with pure chitosan beads. The sodium dodecyl sulfate polyacrylamide gel electrophoresis method was used to test the adsorption of positively charged proteins. The obtained CA-CS/HHF exhibited a high lysozyme adsorption capacity of 42.1 mg g⁻¹ within 3 h, and the adsorption behavior followed pseudo-first-order kinetics and a Langmuir model. Most importantly, the CA-CS/HHF composites possessed good reversibility after 10 regeneration/ reuse cycles and the adsorption capacity was found to have declined to 38.4 mg g^{-1} . The fast adsorption equilibrium, high adsorption capacity and retention properties, as well as good reversibility, attained by these CA-CS/HHF composite beads would be fruitful for guiding future design of a composite for lysozyme adsorption.

Conflict of interest

The authors declare no competing financial interest.

Acknowledgements

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