Enhanced Macroporous Cationic Chitosan Hydrogel by Freezing and Thawing Method with Superadsorption Capacity for Anionic Dyes

Zahraa H. Athab  
University of Babylon

Ahmed F. Halbus  
(ahmed.halbus@uobabylon.edu.iq)  
University of Babylon  
https://orcid.org/0000-0001-9060-7073

Ahmed S. Abbas  
University of Babylon

Jasim M. Salman  
University of Babylon

Abbas J. Atiyah  
University of Babylon

Research Article

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Abstract

In this work, we have developed a simple method to prepare cationic chitosan hydrogel with interconnected porous structure using freeze–thaw process and the obtained hydrogel was named FCS hydrogel. Scanning electron microscopy (SEM) imaging revealed that the synthesized hydrogel demonstrated interconnected porous structure in the range of 5–20 µm. We also showed that the FCS hydrogel exhibits pH responsiveness behavior, and demonstrated reversible swelling and de-swelling behaviors maintaining their mechanical stability. We demonstrate that the FCS hydrogel swelling capacity decreased at alkaline pH and increased with a decrease in the pH value. Besides, the FCS hydrogel presented specific surface area of 78.25 ±8.75 m²g⁻¹, due to the cryogenic treatment of glutaraldehyde cross-linked chitosan hydrogel could increase the surface area and permeability of composite hydrogel and then strongly increasing the adsorption capacity. Subsequently, the FCS monolithic hydrogel tested dyes removal, which provides a high removal efficiency towards anionic dyes including congo red (CR) and sodium fluorescein (SFL) dyes. Significantly, we show that the FCS hydrogel could be regenerated and reused as an adsorbent for wastewater treatment without significant loss of pollutants removal efficiency over a number of adsorption and washing cycles. This study offers a promising environmental friendly and sustainable interconnected porous hydrogel for anionic dye removal from wastewater.

Introduction

Nowadays, water pollution caused by toxic organic materials are a considerable serious threat to the environment [1, 2]. Dyes are the main source of wastewater toxicity in different industrial activities such as paper, textile, plastics, food and pharmaceutical industries [3]. Dye wastewater have harmful effect to the human and animal health and all the natural ecosystems due to these materials contain many aromatic compounds, which are chemically stable and highly toxic [4, 5]. Therefore, developing new strategies and materials for removing these pollutants from wastewater has attracted important attention. Various techniques such as ion exchange, chemical precipitation, solid phase extraction, separation, biological treatment, and adsorption have been used for wastewater treatment [6–8]. Among these technologies, adsorption is considered the most effective, versatile and promising method for dyes removal owing to the flexible operation, convenient processing, cost-effective, and high removal efficiency [9]. Activated carbon has been a common adsorbent material, being used for removing many types of dyes [10–14]. However, sometimes the preparation of activated carbon is comparatively expensive and the separation and regeneration from treated water is complicated, which limited its application for water purification and thus increases requirements for substitutes adsorbent materials being economical and sustainable [15]. Hydrogels are considered novel materials used as adsorbents with rapid removing of dyes owing to their unique three-dimensional porous network structure, which can absorb large amounts of water with lack of solubility in water due to the chemical or physical crosslinking between the polymeric chains [16, 17]. Furthermore, hydrogels materials provide hydrophilic function groups that can be acted as an active site for organic contaminations removal [18]. In addition, the hydrogel simply
separates from treated solution and have the ability to recover and reuse a number of times, which meets the demand for sustainability and eco-friendly development [19]. Some types of hydrogels are named smart materials, which often respond to the stimulus of the surrounding environment, such as pH, temperature, light, and magnetic and electric field owing to their functional groups presented along the polymer chains [20]. For instance, the swelling and deswelling behavior of these materials may be sensitive to any change in the pH, temperature of the solution or even to the presence of the light [21]. Polysaccharides are desired candidates for preparing hydrogels material as adsorbents for wastewater treatment due to they are biodegradable, non-toxic and available in nature [22]. In particular, chitosan is a widely used biopolymer for preparing hydrogel for pollution remediation owing to its the second most abundant biopolymer in nature after cellulose and have high adsorption capacity due to containing a large number of functions group such as hydroxyl and amine groups, which make them efficient adsorbent for different organic pollutants removal [23, 24]. Chitosan considerable to be an excellent example of a pH-responsive natural polymer [25]. Moreover, it has been found that chitosan hydrogel has a better adsorption capacity than that chitosan polymer with non-crosslinking specially in acidic medium due to chitosan can be dissolved in acidic environment [26]. Chitosan hydrogels designed via either chemical or physical cross-linking using different crosslinker agents such as glutaraldehyde, glyoxal, formaldehyde, ethylene glycol diglycidyl ether, epichlorohydrin and oxalic acid were reported [27–30]. The mechanical properties and the chemical stability of chitosan are improved after cross-linking process [31]. However, unmodified chitosan hydrogel has a low surface area and a lake of porosity, which resulting low adsorption capacities [30]. To overcome these restrictions, there are different physical and chemical methods have been used for modifying chitosan hydrogel in order to increase the surface area and to create desirable porous structure and improve the mechanical properties, which are very important in our application [32, 33]. Chitosan has been chemically modified by grafting with nanoparticles or crosslinking with other polymers. Ehsan and co-workers have grafted chitosan with polyacrylamide matrix and dispersed TiO$_2$ nanoparticles in the network structure (TiO$_2$-PAM-CS) using in situ technique. This composite material was applied for the adsorption of Sirius yellow K-CF dye from aqueous solution and it was showed high removal efficiency of this anionic dye [3]. Hu Tu et al. enhanced the chitosan hydrogel by constructing chitosan with different ratios of cellulose by using the environmentally friendly method and materials. The prepared hydrogels were not only low cost, but also possessed high mechanical properties. Besides, the hydrogels had outstanding high adsorption ability towards anionic dyes in aqueous solution and preformed a surface area 33.9 m$^2$g$^{-1}$ when the ratio between chitosan:cellulose was 9:1 [34]. Mahmoodi et al. have studied the use of GO for modification of chitosan hydrogel (GO–CTS) and amine graphene oxide–chitosan (AGO–CTS) hydrogel which boosting the adsorption capacity. The removal of diclofenac GO–CTS hydrogel was investigated and it was exhibited high adsorption efficiency with maximum adsorption capacity reach to 132.11 mg g$^{-1}$ [35]. Alves D.C.S. et al. synthesised chitosan hydrogel modified with carbon nanotubes (HyCNT) and tested it adsorption efficiency towards phenol. They showed that HyCNT demonstrated high specific surface area, which is interesting characteristics for adsorption application [36]. However, these chemical modifications for chitosan hydrogel suffer from a number of disadvantages including synthesis these materials can be costly and time consuming [37]. In addition, the regeneration of these modified materials for adsorption
application were very complicated [38]. Besides, added another polymer with different function groups led to a lack the selectivity in the adsorption process. Furthermore, there are added concerns over the environmental impact of CNT and GO, which if leached from the chitosan composites, can enter the aquatic system and have adverse effects on the aquatic ecosystem [39]. Moreover, GO and CNT particles may be blocking the pores of chitosan hydrogel, which is impact on the diffusion of the pollutants through the porous structure and led to decrease the adsorption capacity [40]. Therefore, various researches have been used physical modifications of hydrogels to control the porosity to form superporous structure with high swelling properties such as gas blowing or freeze drying techniques, which can increase its adsorption capacity [41–43]. The freeze-drying technique has its own disadvantage as well such as high energy required, time consuming and may led to shrinkage the network structure because the interfacial tension resulted by solvent evaporation thus limiting their usage [44, 45]. This work provides new ideas for physical modification of chitosan hydrogel was used. Freezing and thawing method have been carried out on glutaraldehyde cross-linked chitosan hydrogel in order to produce porous chitosan hydrogel to increase the diffusion properties. It is worth noting that freeze–thaw process preferred because it provides open the pores in the structure of the CS matrix thus penetration from the external surface of the sorbent to the intraparticle pores and adsorption on both the external and internal available surfaces via interactions of the solute. Therefore, increase of the specific surface area due to the expansion of the chitosan network structure by changing the pores from close or blind pores to interconnected pores, which are desirable in wastewater treatment field. This is shown schematically in Fig. 1A,B. To the best of our knowledge, since there have no studies about use this type of modification for chitosan hydrogel by freezing-thawing treatment and used it for adsorption of anionic dyes.

**Materials And Methods**

**Materials**

Chitosan was obtained from across organics with a medium molecular weight (Mwt 100000-300000). Glutaraldehyde (50%) was purchased from Sigma-Aldrich. Congo red, Methylene blue dye (MB) and Fluorescein dye (acid yellow 73) (FL) were also purchased from Sigma-Aldrich. Rhodamine (6G) was obtained from Chem. Supply. Acetic acid was purchased from macron fine chemicals. All other chemicals used in this research were used directly without further purification. All solutions in this study were prepared with distilled water.

**Preparation of chitosan hydrogel**

The chemically cross-linked chitosan hydrogel was prepared based on a modified version of the method described by Xinhong Chang et.al. using glutaraldihyde as cross linker agent [27]. 1.0 g of chitosan was dissolved in 50.0 mL of 1.0 vol % acetic acid solution under magnetic stirring at room temperature (solution 1), and 1.0 mL glutaraldehyde (50 wt %) was dissolved at room temperature in 20.0 mL distilled water (solution 2). The two solutions were mixed at room temperature under vigorous stirring for 2 min and after that immediately placed into the plastic syringes of an internal diameter of 1 cm then gradually
transformed into wet gels within 10 min and were aged for 24 h. To form a porous chitosan hydrogel for wastewater treatment, the obtained chitosan hydrogels were frozen in a freezer at −20 °C for 48 h and then melted at room temperature. After cooling down the frozen chitosan hydrogel was washed with 0.01 M solution of hydrochloric and then with distilled water to remove all unreacted material. Finally, the obtained porous monolithic chitosan hydrogel was named FCS and it was stored in distilled water (DW) at room temperature until use.

**Characterization**

The structural morphology of the fabricated chitosan hydrogel was performed by using scanning electron microscopy (JEOL JSM-6480 LV SEM instrument). The sample was fractured to create a fresh surface, then adhered onto an SEM stub. The sample was then sputter coated with gold via sputter coater and then transferred into the SEM chamber. The analysis of the functional groups of the powder chitosan and the synthesised chemical cross-linked chitosan were carried out via examined using Thermo Scientific Nicolet 380 FT-IR (Thermo Scientific, Hemel Hempstead, UK), supplied with attenuated total reflectance (ATR). Each sample was analyzed in the wavenumber range of 4000–500 cm\(^{-1}\) using 4 scans. The specific surface area, pore size distribution and total pore volume for the freeze dried chitosan hydrogel material was measured by using Micromeritics Tristar II 3000 (USA) BET-N\(_2\) surface area analyser using N\(_2\) gas adsorption analysis by stabilising at -196 °C. Chitosan hydrogel was freeze dried using (VirTis SP SCIENTIFIC) to obtain the monolithic chitosan aerogel material and then was dried at 100 °C for 12 hours for degassing the samples before carrying out the surface area analysis. The pore size distributions were calculated using the Barett-Joyner-Halenda (BJH) model. A Zetasizer Nano instrument (Malvern Instruments, Malvern, UK) was used to measure the zeta potential on the surface of the FD-chitosan hydrogel. Chitosan hydrogel porosity percent (P\%) were calculated from the density of wet FCS hydrogel (\(\rho_{\text{FCS wet}}\)) and the density of dry FCS hydrogel (\(\rho_{\text{FCS dry}}\)). The swelling degree (SD) of FCS hydrogel was measured by using gravimetric analysis, from pH 2 to 10. Before measurement, the hydrogels were left in DW for 24 h. They were then transferred in a water at the desired pH for 24 h. The excess water on the surface of the sample was gently removed using filter paper before measurement. The weight of the hydrogel at a given pH (Ws) and the weight of the dehydrated hydrogel at pH 10 (Wd) were recorded and used to calculate SR according to the following equation:

\[
\text{SD} = \frac{W_s}{W_d}
\]

The swelling kinetics and the water uptake ratio (WU) of FCS hydrogel were also measured by gravimetry analysis. Briefly, the hydrogels were first plunged in pH 2 solution, brought to equilibrium, and weighed (Wg). They were then plunged in pH 10 solution for 24 hours (Wg), until equilibrium in the dehydrated state was reached. They were then immersed again in pH 2 and then the weight of the hydrogels was recorded (Wt) at different time through one hour. As mentioned previously, before weighing the excess of water on the surface of the hydrogels were absorbed with filter paper. The water uptake ratio WU is calculated using the following equation:
The deswelling kinetics and the water retention ratio (WR) of the FCS hydrogel were also studied by gravimetry analysis. The FCS hydrogels, initially stored in distilled water at pH 2 for 24 h then the weight of the hydrogels was recorded (Wg), were then transferred in pH 10 solution. Then, the weight of the hydrogels was recorded (Wt) at the same predetermined time intervals for swelling study. As mentioned above, before measuring the weight the excess of water on the surface of the hydrogels were absorbed carefully with filter paper. The water retention ratio WR was calculated using the following equation:

\[
WR = \left( \frac{Wt}{Wg} \right) \times 100
\]

The zeta potential of FCS hydrogel was evaluated by dynamic light scattering (DLS) using Zetasizer Nano ZL instrument (Malvern, UK). FCS hydrogel was crashed into small parts and then dispersed in deionized water using an ultrasonic probe (Branson LTD). The measurement was carried out in triplicate at room temperature.

**Adsorption studies**

CR, MB, R6G and FL were selected to examine the ability of adsorption and removal capacity of the chemical cross-linked CS hydrogel. The experimental data were recorded using a UV–Vis spectrophotometer at 498 nm, 664 nm, 525 nm, and 490 nm for CR, MB, R6G and FL respectively. The prepared FCS hydrogels were put into 100 mg/L dye solutions. Then, the solutions were shaken in the dark condition regularly for 3 hours using a platform shaker (Gemmy orbit shaker model: VRN-480). The absorbance at the maximum absorption wavelength was measured before and after adsorption. The batch adsorption tests were carried out to evaluate the percentage of dye removal (R%) and the adsorption uptake quantity \(Q_e\) of CR or SFL on FCS hydrogel whereby the monolithic hydrogel (~0.05 g of FCS, 3 mL volume) were plunged into isolated conical flasks with 100 mL serial solutions of CR or SFL dyes (10-60 mg L\(^{-1}\)). The samples were gently shaken in the dark condition at room temperature (25 ± 2 °C) in the dark condition with a shaker speed of 100 rpm until reached the equilibrium. All experiments were carried out in triplicate and the concentration of anionic dyes solutions before and after the adsorption process were calculated using double beam UV–Vis spectrometry (1650 PC-UV-Visible spectrometer Shimadzu.) at the maximum absorbance of CR dye (498 nm) and FL dye (490 nm). The dye removal percentage (R%) of the dyes and the adsorption quantity of the dye adsorbed per unit mass of adsorbent at particular time \(Q_t\) and at equilibrium \(Q_e\) (mg. g\(^{-1}\)) were calculated according to the following equations:
Where $C_0$ and $C_t$ are the initial dye concentration (mg·L\(^{-1}\)) before adsorption and is the concentration of dyes after adsorption at a given time $t$. $C_e$ (mg L\(^{-1}\)) is the concentration at equilibrium, $m$ and $V$ are the dose of FCS hydrogel (g) and the volume of dye solution (L).

**Adsorption isotherm**

The adsorption isotherms were investigated using both Langmuir and Freundlich isotherms for adsorption of CR and FL dyes on the FCS hydrogel. The Freundlich model is an empirical model and more applicable at low adsorbate concentrations. In the theoretical Freundlich model, it is assumed that adsorption of dye occurs on a heterogeneous surface of the adsorbent, it suggests multilayer adsorption and there is an interaction between the adsorbate and adsorbent molecules depend on the electrostatic interaction between negative and positive charges. The Freundlich isotherm parameters were calculated according the following equation:

$$\log Q_e = \log K_F + \frac{1}{n} \log C_e$$

Where, $Q_e$ (mg/g) is amounts of dyes adsorbed at equilibrium, $C_e$ (mg/L) is the concentration the dye at. $K_F$ and $(1/n)$ are Freundlich isotherm constants. The Langmuir model, it is assumed that the dye undergoes monolayer adsorption around the homogenous surface of the adsorbent and that there is no interaction between the dyes molecules and the adsorbent. The Langmuir isotherm parameters were calculated using the following equation:

$$\frac{C_e}{Q_e} = \frac{1}{Q_{\text{max}} K_L} + \frac{C_e}{Q_{\text{max}}}$$

Where $C_e$ (mg L\(^{-1}\)) is the equilibrium concentration of CR and FL dyes in solution, $Q_e$ (mg g\(^{-1}\)) is the equilibrium adsorption capacity of CR and FL dyes on the FCS hydrogel, $Q_{\text{max}}$ (mg g\(^{-1}\)) is the maximum amount of dyes on the FCS hydrogel to form monolayer on the surface, $K_L$ (L mg\(^{-1}\)) is the Langmuir constant that relates to the affinity of adsorption active sites.
Results And Discussion

Characterization of FCS hydrogel

The morphology of the FCS hydrogel of the fracture surface was analyzed by SEM, and the results were shown in Fig. 2. In the cross-sectional images of the FD-CS hydrogel, there are a highly porous matrix with interconnected pores extending from the core to the surface of the monolithic hydrogel, indicating the successful crosslinking of chitosan polymer chains to form chitosan composite hydrogels. Furthermore, the prepared chitosan hydrogel with freeze-thawing treatment exhibited open pores with high interconnectivity structure. In contrast with other study which presented chitosan hydrogel with rough surface but with no pores were observed on the surface [25]. This is due to the freeze-thawing method has been used to create porous chitosan hydrogel with interconnected pores [42]. During cooling the prepared chitosan hydrogel, the nucleation and the ice crystals was grown and moves into non-nucleated solution leading to change the morphology and form interconnected pores [46]. At a temperature below the freezing temperature (room temperature ), these ice crystals melt and gives rise to interconnected macroporous networks [47, 48]. This network structure can improve the specific surface area and then increase the adsorption capacity of the dye molecules from aqueous media due to promoted diffusion of the dyes molecules within the hydrogel structure and makes contact between the functional groups of the chitosan hydrogel with dye molecules more efficient [49]. The specific surface area and pore size distributions of FDCS hydrogel was determined by employing the nitrogen sorption technique. The $N_2$ adsorption–desorption isotherms of FDCS are represented in Fig. 3A.

The $N_2$ adsorption–desorption isotherm for FCS belongs to type H3 hysteresis loop according to IUPAC classification, which proves interconnected pore structure with a parallel plate shape [50]. The synthesized FCS in this work displayed a specific surface area of $78.25 \pm 8.75 \, \text{m}^2\text{g}^{-1}$. The BET surface area of the prepared FCS chitosan compared with previous studies in the literature and it was found that the surface area for the FCS hydrogel was larger value [50–52]. These results strongly support the fact that the surface area and porosity of FCS were increased after freezing the CS hydrogel. Obviously, cryogenic treatment for the synthetic glutaraldehyde cross-linked chitosan hydrogel played an impressive role in the increasing of the surface area of this hydrogel. As expected, after freezing the CS hydrogel convert the porous structure from close pores to interconnected pores, which can allow more active sites in the hydrogel to interact with dye molecules and led to improve adsorption efficiency [53, 54]. Therefore, FCS hydrogel with larger specific surface area and more active groups are a promising adsorbent materials due to these characteristics are particularly important for adsorption purposes for wastewater treatment [55]. The BET results well proved the SEM images as the pores development can be seen obviously in the surface morphology of FCS hydrogel. For a better evaluation of the impact of the initial CS concentrations on the porosity % of FCS hydrogel, four different CS concentrations were used by fixing the crosslinking agent concentration constant. Increasing the CS concentration decreased the porosity of final FCS hydrogel as shown in Fig. 3B. The average porosity of FCS composite hydrogels decreases from 93.49 % to 80.05 % when the initial concentration of CS increases from 0.5 to 2.5%
w/v. This is due to the fact that increasing the concentration of the polymer increases the cross-link densities of the network structure of the prepared hydrogel, which is lead to decrease the porosity [56].

In order to further confirm the successful crosslinking and the structural changes of composite hydrogel after crosslinking, the infrared spectra were investigated for powder chitosan and cross-linked chitosan hydrogel as shown in Fig. 3C. Comparison of the FTIR results reveals that the FCS cross-linked chitosan hydrogel shows spectra that are different from those of non-cross-linked chitosan. Non-cross-linked chitosan demonstrate peaks at 949 and 1104 cm\(^{-1}\) are attributed to C–O–C bridge of chitosan unit. The absorption bands at 2911 and 2873 cm\(^{-1}\) can be corresponds to C–H symmetric and asymmetric stretching. A broad band in the region of 3240- 3400 cm\(^{-1}\) attributed to the associated of the vibrations of -OH and –NH\(_2\) groups. Band at about 1564 cm\(^{-1}\) are attributed to the bending of the primary amine's N–H bond [57]. After chemical crosslinking of chitosan using glutaraldehyde, these band present a change in it is intensity and width, this is occurring due to the bond generated by overlapped the N-H, O-H, and C-O bonds of CS with the GA during the crosslinking reaction. This proven that some of the hydrogen bonds in the chitosan structure are broken and new hydrogen bonds between GA and chitosan polymer chains are created as NH\(_2\) and OH functional groups engaged in hydrogen bonds [58]. A new peak at about 1646 cm\(^{-1}\) was observed, which is most probably attributed to the imine bond (C=N). This bond produces and observed in the spectrum of FCS hydrogel due the chemical crosslinking reaction between GA and CS amino groups [26]. The FTIR results confirms that GA cross-linking takes place by the reaction of amine groups of chitosan with aldehyde groups of glutaraldehyde. Fig. S1 shows change of the chitosan hydrogel color from transparent light yellow to orange this was also confirms the formation of cross-links between CS polymer chains by GA. This is most likely due to the formation of a chromophore as stable crosslinking products, which is not present in the starting precursors [59]. The Z-potential of FCS hydrogel in deionized water was measured by the DLS of suspensions that were prepared by dispersing 0.05 g of the vacuum dried FCS hydrogel in 50 mL of deionized water by a digital sonicator. The zeta potential of FCS hydrogel was found to be +26.5 ± 3 mV as shown in Fig. 3D. At this point, the positive surface charge of FCS hydrogel contributed to increase the adsorption capacity towards the anionic contaminated and this result agrees with previous studies [60–62].

It is important to highlight that the positive value of ZP for FCS hydrogel indicating that the high protonation of the NH\(_2\) groups, which are did not take part in the crosslinking reaction and suggests that the interactions between the FCS hydrogel and anionic dyes molecules may occur by hydrogen bonding and/or electrostatic interactions [63].

The swelling behavior of the FCS hydrogel was analyzed via gravimetric method. In detail, a small piece of vacuum dried hydrogel (~0.05 g) was immersing in D.I. water for 24 hours at room temperature (25±3 °C) and neutral pH. The FCS hydrogel shows remarkable and quick volume change due to the water pass through the hydrogel via capillary forces. This was owing to its interconnected porous structure and the presence of abundance hydrophilic functional groups on backbone [64]. Swelling behavior of FCS
hydrogels as a function of pH was evaluated as shown in Fig. 4. As can be seen in Fig. 4A the swelling ratio of FCS hydrogels with decreasing the pH value from 12 to 3 was determined. The swelling ratio of FCS hydrogels in pH 3 was 26.3, which doubled compared to the same sample in pH 12 (11.7). It is worth noting that, FCS hydrogel in acidic environment showed higher swelling ratio than the same sample in basic and neutral environment. In order to explain this phenomenon, it was due to in acidic media ionized the $-\text{NH}_2$ groups in the FCS hydrogel to ammonium ion ($\text{NH}_3^+$) and these cationic charges made repulsive forces between polymer chains [65]. Fig. 4B shows the swelling deswelling reversibility of FCS hydrogel between pH 3 and pH 10. The results demonstrated that the reversibility toward the pH changes was retained even after more than ten cycles. FCS hydrogel water uptake were studied by measuring the degree of swelling with time. Fig. 4C depicts the degree of swelling as a function of time for FCS hydrogels in acidic media. Fig. 4D shows deswelling kinetics of the FCS hydrogel at pH 10. FCS hydrogels exhibited quicker loss most of water after change the pH value from 3 to 10 in comparison with water uptake of the same FCS sample. This results are reasonable since both swelling and deswelling processes start from the surface of the hydrogels to inside the hydrogel through its porous structure. During swelling, water will take longer time to penetrate inside the sample, while it will take a shorter time for the transportation from the hydrogel during deswelling. Fig. 4E demonstrated schematic represented the swelling and deswelling properties of FCS hydrogel in acidic and alkaline conditions respectively.

**Adsorption study**

The synthesized three-dimensional FCS hydrogel with unique pore structure were evaluated for the removal of some organic dyes. In the present work, Congo Red and Sodium Fluorescent dyes as an anionic dye model and Methylene Blue and Rodamine 6G as cationic dye model to test the ability of FCS hydrogel to remove organic dyes. It is also important to note that the preparation process of our FCS hydrogel requires only mixing of chitosan and the chemical cross linker in acidic media and then freeze-thaw treatment to generate the pores, with no further drying step required to form the hydrogel absorbent. This low energy fabrication method, together with the use of the green solvent, water, renders our approach relatively environmentally friendly. Further, we described the dye adsorption process utilizes monolithic hydrogel, with no required for the additional steps of centrifugation or adsorbent separation, unlike methods that utilized particulate-based absorbents. Fig. S2 show a photograph of the adsorption process of CR and SFL by FCS monolithic hydrogels. After immersing the FCS monolithic hydrogel by 24 hours into CR and SFL dyes solutions separately, the red and yellow solutions containing CR and SFL dyes become very light and almost clear and transparent, whereas the hydrogels turned red and dark yellow due to the adsorption of CR and SFL dyes respectively. These images indicate that the composite FCS pores hydrogels possess highly efficient of anionic dyes adsorption. In contrast, FCS hydrogel demonstrates very low removal towards MB and R6G as shown in Fig. S3. However, the monolithic FCS hydrogel turned to blue and dark orange after immersing these hydrogels into MB and R6G solutions by 24 hours. The reason for this is due to the interconnected porous structure of FCS hydrogel make easily filled by these dyes solutions without being adsorbed [66]. The adsorption capacities of the FCS hydrogel toward CR and SFL dyes were evaluated by using the UV−Vis spectra to measure the concentrations of
the dye solution before and after adsorption as shown in Fig. 5 A and B. Removal and adsorption of CR and SFL dyes onto the FCS monolithic hydrogels from the aqueous solution was observed. UV-Vis spectroscopy shows low absorption bands for these two dyes in the solution after adsorption. This phenomenon may be explained by electrostatic attraction between NH$_3^+$ group protonated in the acid medium of FCS hydrogel and anionic dye molecules formation (Dye-NH$_3^+$), which is cause the captured of this type of dyes on FCS hydrogel [3]. We discovered that the hydrogen bonding and electrostatic interactions played a significant role in the adsorption of the anionic dyes [67]. Similarly, FCS hydrogel adsorption capacity were tested towards MB and R6G dyes aqueous solutions as cationic dyes. In contrast to CR and SFL dyes, no significant adsorption was observed toward MB and R6G (less than 10%), with little change in the UV-Vis spectrum of the dyes solutions after adsorption as shown in Fig. 5C and D. This confirmed that FCS monolithic hydrogel has no obvious adsorption of cationic dyes on its surface due to the electrostatic repulsion interactions between the dyes and the cationic FCS hydrogel [67]. However, it should be noted that the poor removal efficiency of cationic dye due to the hydroxyl groups in the FCS may also participate to chelation with cationic dyes [68]. In order to study the adsorption behavior of the FCS porous monolithic hydrogel towards CR and SFL dyes batch adsorption method was also performed. The batch adsorption of these two anionic dyes were determined at different initial concentrations with constant weight of FCS hydrogel (0.05 g) for different adsorption times intervals at a pH value of 6. As shown in Fig. 6A, demonstrates the adsorption process of both CR and SFL dyes (the concentrations were 60 mg/L) onto the FCS monolithic hydrogel were progressed rapidly in the initial stage and the adsorption rate decreased gradually moving toward equilibrium after 12 hours. The best possible explanation for this phenomenon is that at the beginning of the reaction, the higher available active site with the higher driving force on the FCS hydrogels surfaces provides better opportunity of collision between anionic dyes molecules and the adsorbent [69]. As the adsorption reaction reached equilibrium the adsorption capacity was remained constant, which is may be attributed to saturated the available adsorption sites of the FCS hydrogel by dyes molecules [70]. The adsorption capacity at equilibrium of CR and SFL dyes were calculated and it were found 104.9 and 99.6 mg·g$^{-1}$ respectively. The removal percentage of CR and SFL dyes were up to 96.73% and 95 % at equilibrium as shown in Fig. 6B. The study of the adsorption isotherm play a significant role to describe how adsorbates molecules interact with adsorbents surface [71]. In the present work, to investigate the equilibrium condition of anionic dyes (CR and SFL) onto FCS hydrogel, two types of isotherms including Freundlich and Langmuir isotherm models were evaluated. The Langmuir isotherm model can be regarded as a single layer adsorption system onto a homogeneous surface, while the Freundlich model are considered as a heterogeneous multi layers adsorption system [72]. As we can see in Fig. 6 C and D the plots of Langmuir and Freundlich isotherm for CR and SFL dyes. The fitted parameters and data are shown in Table S1. The results indicate that the Freundlich adsorption isotherm showed a better value of correlation coefficient ($R^2$). Therefore, the adsorption of both CR and SFL dyes onto FCS hydrogel obeyed with the Freundlich model, and is mainly multilayer adsorption. Through the Langmuir model fitting, the maximum adsorption capacity $q_m$ values for CR and SFL dyes onto FCS hydrogel were obtained and it
were found 136.69 and 142.857 mg.g$^{-1}$ respectively. The small difference in maximum adsorption capacities of these dyes was owing to their different affinities for adsorbance.

As a results, the high adsorption capacity of FCS hydrogel due to the adsorption mechanism is affected by the interconnected porous structure of the hydrogel, which is allow better access of dye molecules through the internal structure of the hydrogel and increase the available active adsorption sites [73].

Additionally, the strong electrostatic force between the positively charged amine groups (-NH$_3^+$) in the hydrogel FCS interact with the negatively charged groups on anionic dyes and the like hydrogen bonds interaction encourage anionic dyes adsorption on hydrogels resulting high adsorption capacity [16, 74]. To preform, the adsorption behavior of the FCS hydrogel, kinetic models containing pseudo-first order (PFO) and pseudo-second-order (PSO) models, which are widely used in the fitting of experimental kinetics data as shown in Table S2 and S3 for CR and SFL dyes respectively. The fitting plots of the adsorption kinetic model is shown in Fig. 7. It was observed that the correlation coefficient ($R^2$) corresponding to the PSO model was significantly higher than that corresponding to the PFO model. Furthermore, the results showed that the simulation adsorption capacity from the plot of PSO model was found to be well fit with the experimental results than that calculated from PFO. Therefore, PSO model better describe the adsorption mechanism for both CR and SFL dyes on FCS hydrogel. It is worth to note that the adsorption reaction of CR and SFL dyes on to FCS hydrogel is more tending towards physical adsorption. Basically, the adsorption mechanism of dye molecules on a porous hydrogel adsorbent material cannot be described by only the PFO and PSO models due to the adsorption on these materials may include a multi-stage process [75]. As a result, an intraparticle diffusion model was employed for further interpretation to show the movement of the dye molecules from the aqueous solution into the interconnected pores of the FCS hydrogel. Fig. 7E and F show that the linear plots do not pass through the origin for both CR and SFL dyes, indicating that the intraparticle diffusion models is based on the migration of the adsorbates into the hydrogel pores and controlling the rate adsorption process and the results of adsorbent parameters for intraparticle diffusion model were listed Table S2 and S3 for CR and SFL dyes respectively. This model represents two stages, the first stage where there is a rapid adsorption of dye molecules on the external surface of the hydrogel and the second stage where the intraparticle diffusion is the rate-limiting step [76]. Therefore, in addition to PSO, the intraparticle diffusion model plays a significant role to interpret the adsorption mechanism.

**Reusability and desorption studies**

In order to evaluate the reusability and regeneration of FCS monolithic hydrogel as adsorbents to be a cost effective material, desorption experiments were carried out using FCS hydrogel as adsorbent and CR and SFL dyes as adsorbates. So the desorption of adsorbed CR and SFL dyes from FCS monolithic hydrogel was investigated in a batch desorption system. Because the FCS is a monolithic hydrogel, the collection of the adsorbent was fast and easy. The FCS hydrogel was recovered by immersing in a 0.01 M NaOH solution for 6 h to desorb CR and SFL dyes that had adsorbed on the hydrogel. Then the hydrogel was protonated in excess 0.01 M HCl for 1 h to regenerate cationic active sites to use in the next
adsorption cycle. The regenerated FCS was further washed by deionization water to reach neutrality. The reusability of FCS hydrogel was evaluated over five cycles of adsorption/desorption by using NaOH solution as the desorption agents with 0.05 g FCS monolithic hydrogel in 100 mL dye solution (10 ppm) at 25°C for 24 h (each cycle), and the results are shown in Fig. 8. As seen, no significant losses in the dye adsorption performance was detected even after five cycles of reusability. It was observed that about 9% and 3% reduction in removal efficiency of FCS hydrogel for CR and SFL dyes respectively.

Thus, the results depict that the electrostatic attraction is dominating driving force in the adsorption process in this study. It is important to highlight that the positively charged amino group in FCS hydrogel was deprotonated in alkaline conditions, which is affecting the electrostatic interactions between FCS hydrogel and anionic dye and then desorbed the loaded dye from adsorption active site [77]. Therefore, it can be concluded that FCS monolithic hydrogel promising ecofriendly adsorbent material due to can be regenerated easily and used repeatedly as an efficient adsorbent for water purification.

**Conclusion**

In the present study, macroporous FCS hydrogel was successfully prepared using a new and simple freeze-thaw strategy as physical modification of glutaraldehyde cross-linked chitosan hydrogel with the aim of fabrication interconnected porous hydrogel. This porous structure in the hydrogel materials improves the accessibility of adsorbates in the hydrogel active sites and increases the dye adsorption capacity of the FCS hydrogel. To the best of our knowledge, there are no studies regarding the use of the freeze-thaw method for the fabrication of macroporous chitosan hydrogel with interconnected pores. The FCS hydrogel was characterized and the results demonstrated that FCS hydrogel had a highly porous matrix with interconnected pores extending from the core to the surface of the monolithic hydrogel as revealed by SEM. The swelling-deswelling behavior of FCS hydrogel was investigated and the results indicated that the FCS hydrogel exhibited good swelling properties in acidic media than the same sample in basic media. The reason behind that the amine groups (-NH$_2$) of chitosan in acidic medium protonation to ammonium ion (NH$_3^+$) due to the presence of hydrogen ions and these cationic charges made repulsive forces between polymer chains and increase the capillary force. The swelling deswelling reversibility results of FCS hydrogel showed a significant pH dependent, which demonstrated that the reversibility toward the pH changes was retained even after more than ten cycles. The detailed evaluation of adsorption behavior of different organic dyes by FCS hydrogel revealed that this hydrogel possesses the unique adsorption toward anionic dyes with a large adsorption capacity, but very low adsorption capacity toward cationic dyes. The electrostatic interactions dominated the interaction between the FCS hydrogel and the dye molecules regardless to their charge states. The adsorption isotherms data of CR and SFL onto FCS showed a better fit with the Ferunditich model. The adsorption kinetics analysis exhibited that the adsorption process could be described well with the pseudo-second-order model. The obtained FCS provided not only selective absorption towards anionic dyes with a high removal efficiency but also exhibited excellent adsorption efficiency with no significant loss even after five cycles of reusability. From the obtained results, the synthesized FCS hydrogel by the freeze-thaw method is
considered to be a promising candidate for the removal of anionic dyes from an aqueous solution due to its high adsorption capacity and cost-effective properties.

**Declarations**

**Notes**

The authors declare no competing financial interest.

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Figures

Figure 1

Schematic illustration of (A) the synthesis of cationic chitosan with interconnected hydrogel using glutaraldehyde as a crosslinking agent and treated by freezing and thawing method (B) its application for removal of anionic dye by adsorption (Congo Red dye as typical model for anionic dyes).

Figure 2

SEM cross-sectional images of the FCS hydrogel dried by freeze drying (A-D) at different magnifications and scales.

Figure 3

(A) $\text{N}_2$ adsorption/desorption isotherms of FCS hydrogel (B) FTIR spectrum of powder chitosan and glutaraldehyde cross linked chitosan (C) Bar charts showing porosity of FCS hydrogel with different CS concentration (D) The zeta potential of FCS hydrogel was dispersed in deionized water at pH 7.
Figure 4

(A) Swelling degree (SD) as a function of pH of the FCS hydrogel at room temperature (B) Reversibility cycle of swelling–deswelling of FCS hydrogel as a function of time by varying the pH from 3 to 10 (C) Water uptake ratio (WU) for the FCS hydrogel as a function of time, when plunged in a pH 3 after initial equilibration at pH 10. (D) Water retention ratio (WR) of FCS hydrogel as a function of time, when plunged in a pH 10 after initial equilibration at pH 3. (E) Schematic illustration of swelling and deswelling behavior of the FCS hydrogel in wet states at different pH and at 25°C. Corresponding optical image of the shrunk and swollen states of FCS hydrogel in basic and acidic media.

Figure 5

Selective adsorption behaviors toward CR, SFL, MB and R6G dyes: UV-Vis spectra of CR (A), SFL (B), MB (C) and R6G (D) before and after adsorption by using FCS hydrogel for 24 h. Insets are the images of CR, SFL, MB, and R6G dyes solutions before and after adsorption for 24 hours.

Figure 6

(A) The effect of adsorption time on adsorption capacity of CR and SFL by FCS hydrogels (the initial dye concentration: 60 mg/L, adsorbent dosage: 0.05 g in 100 mL of dye solutions, pH of 6, t= 25 °C, shaking rate is constant at 100 rpm). (B) The effect of contact time on the removal percentage of CR and SFL dyes of dye (C) and (D) Freundlich and Langmuir adsorption isotherms respectively.

Figure 7

Kinetic models for adsorption of CR and SFL dyes using FCS hydrogel (A) and (B) pseudo-first order model for CR and SFL dyes respectively (C) and (D) pseudo-second order model for CR and SFL dyes respectively (E) and (F) intraparticle diffusion model for CR and SFL dyes respectively.
Reusability of FCS monolithic hydrogel for CR and SFL dyes (FCS dosage = 0.05 g; dye concentration = 10 mg/L in 100 mL; pH = 6; t = 25 °C; shaking rate is constant at 100 rpm).

Supplementary Files

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