Improvement of Drinking Water Properties Using a Novel Biosorbent Material

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KEYWORDS:
Coagulation, Adsorption, Modified flaxseed husk (MFH), Flaxseed husk ash (FHA), Turbidity, Natural organic matter, Iron Kinetics Isotherm

Abstract—In the present study, a new material has been discovered in the world of natural coagulants and adsorbents to improve drinking water properties. Flaxseed husk was used as a natural coagulant and adsorbent to remove turbidity, natural organic matter and iron from underground water for several wells in Al-Jouf region in the north of the Kingdom of Saudi Arabia. To remove turbidity and natural organic matter, modified flaxseed husk (MFH) was prepared by addition of aluminum sulfate (AS) to flaxseed husk (FH) after removing mucilage and oil. The characterization of FH shows high specific surface area (125.18 m²/g) with average pore diameter of 26.28 μm. Moreover, FTIR spectra analysis indicated existence of large number of amino groups and another functional group indicated that FH efficiency about removal of turbidity and natural organic matter. The optimum ratio MFH:AS for removal was (80%:40%). Batch experiments were performed as function of process parameters such as slow mixing time, FH dosage and Initial NOM. The removal of turbidity and natural organic matter (NOM) by modified flaxseed husk (MFH) were investigated using a FH dosage of 40 mg/l-1 and 10 mg/l-1 AS which attained a removal efficiency of 98.0 % for turbidity and 99.3 % for NOM at the highest NOM concentration. Also, reduce the iron concentration in drinking water by flaxseed husk ash (FHA) which prepared by burning flaxseed husk at certain temperature was investigated. Batch experiments were performed as function of process parameters such as FHA type, contact time (CT), FHA dosage and initial Fe concentration. The FHA showed a high removal of Fe³⁺ from aqueous solution. The maximum Fe removal was 90% at FHA200, FHA dose 0.75 g/l and contact time 45 min, The pH value from 7 to 7.8 and the temperature value from 20 to 24 ⁰C. The adsorption equilibrium has been studied by the pseudo first order kinetics model, the pseudo second order kinetics model, Langmuir and Freundlich isotherm, all these models demonstrated the adsorption efficiency of the adsorbent surface behaved in a favorable manner for Fe adsorption.

I. INTRODUCTION

The existence of turbidity and natural organic matter (NOM) in drinking water has received considerable attention in recent decades because it can lead to odor, color and taste problems in addition to formation of disinfection by-products (DBPs) [1]. NOM is widely detected in most water sources such as lakes and rivers [2]. It has been reported that most of NOM in surface and ground water is in the form of humic and fulvic acids [3]. Humic and fulvic acids react with chlorine-based materials during the disinfection process and hence DBPs are formed. DBPs are carcinogenic
and cause many health and environmental problems to human and aquatic life [4,5].

Chemical coagulation using aluminum sulphate followed by sedimentation and filtration is the most prevalent sequence in water treatment plants. These basic processes are efficient for the removal of colloidal or suspended particles, but the removal of NOM is limited [6]. Some researches have addressed improving the coagulation process to remove turbidity and NOM by adjusting the pH of the water and increasing the amount of coagulant, but the high amount of chemical addition was a concern [7]. Lieknes et al. (2004) investigated the micro-filtration by metal membranes as a post process after coagulation for NOM removal [8]. However, the cost of micro-filtration was not considered.

Many researchers have used the adsorption by activated carbon and advanced oxidation processes for removing or degradation of different organics from water [9–12]. While, other researchers have developed low cost adsorbent materials that prepared from agricultural residues such as wheat straw [13], rice husk [14], peanut hull, coconut husk [15], black gram husk [16], Sawdust [17], sugarcane bagasse [14], banana pith [18], and pine grape stalk [16].

Globally, surface water is the main source of water in the world by up to 73 percent, followed by groundwater by up to 18 percent. While indirect sources represent the remaining fraction of less than 9 percent [19]. The majority of freshwater is already underground, represented by soil moisture and aquifers. Humans can use both ground and surface water. Groundwater accounts for 19.3 percent of the world’s human consumption [20]. While in some countries such as Saudi Arabia, groundwater represents 40% of drinking water after seawater desalination, which represents the highest percentage [21,22].

Underground water contains mineral substances. The presence of these minerals in groundwater varies from place to another depending on the nature of the soil and the depth of the aquifers extracted from it. Some of these minerals are iron. Some of these subterranean sources contain iron ions with values in excess of what is allowed in drinking water or human use [23]. There is a need for the emergence of many ways to remove the minerals in this water, including: Sedimentation or precipitation processes, ion exchange, membrane techniques, biological methods, electrochemical processes as well as adsorption [24]. The threshold values for iron are set by the Saudi Arabian government regulation according to Gulf Standard Specification for bottled drinking water No. 149 / 2000 is 0.3 mg.L⁻¹ which specifying the requirements on water for human consumption and the quality of water for human consumption and match with United States Environmental Protection Agency (EPA) [25-27], while the requirements of the Standardization Organization of the Gulf Cooperation Council states that the health value of iron in bottled water is zero [27,28]. Iron occurs in the binary form or occurs as single ions (Fe²⁺) when it is in a dissolved form, but mainly as Fe (OH)₃ in undissolved forms. It can also be occurred bound to humic substances in a colloid form. The form in which iron is occurred and its composition in nature depends on water temperature, pH value, hydrolysis, the oxygen concentration in the aqueous solution (probability of oxidation or not), the solubility of iron compounds in water and iron binding to organic compounds or non-binding [29].

According to the EPA, the presence of iron ions at high levels in water carries bacteria into stomach, causing severe damage during digesting including fatigue, weight loss, stomach problems, nausea, vomiting and joint pain. Advanced symptoms include hemochromatosis which can lead to liver, heart and pancreatic damage, as well as diabetes [30]. In water supply system, iron (II) ion is oxidized to higher forms and the result of this oxidation leads to the formation of hydroxide suspensions causing water discoloration and undesirable turbidity. In well water, when Fe(II) occurs at concentrations up to be more specific milligrams per liter in anaerobic groundwater, color change or turbidity does not occur in the water when pumped directly from a well, but this develops overtime to produce color and turbidity in distribution pipe system at iron level above 0.05-0.1 mg.L⁻¹ [31].

One of the damages caused by the association of bacteria with iron in the distribution pipes over time causes the growth of bacteria and a large change in the water smell. This causes the formation of a thin layer coating the inside pipes. Iron is at the point of consumption, in washing machines and boilers oxidized to the triple state when conditions are favorable in the presence of oxygen [29-31]. All of the previous problems cause a reduction in the cross-section of water flow in pipes, corrosion and deterioration of water quality. This leads to failure of operation of water supply systems [29].

In this paper, first: Chemically modified flaxseed husk (FMH) was used for drinking water treatment. MFH was prepared by addition of aluminum sulfate to flaxseed husk (FH) after removing the mucilage and oil. The removal of turbidity and NOM by coagulation and adsorption using MFH was deeply investigated. Second: Reduce the concentration of iron in drinking water by flaxseed husk ash (FHA) prepared by burning flaxseed husk at certain temperature to achieve the specifications in accordance with the specifications of the Gulf Cooperation Council Water No. 149 / 2000. The study was conducted on a group of wells for drinking water in several fields in Al-Jouf region in the north of Saudi Arabia. The by-product of the flaxseed husk ash after removing oil was used as an absorbent material for the iron element. However, it was found that the measurements of manganese ratios in the studied wells are 0.1 mg. l⁻¹, i.e. within the permissible range according to the World Health Organization, which specified that manganese should not exceed 5 mg per liter.
II. Materials and methods

A. Materials and raw water
Flaxseed was purchased from a local market and stored at room temperature, the by-product which is the result of pressing flaxseed from an Oil refinery in Jeddah city, Saudi Arabia. \((\text{Al}_2\text{(SO}_4)_3\) \text{16H}_2\text{O}, n-hexane, Whatman filter paper grade 1 and NaHCO\(_3\) were purchased from SOMAT-CO, El-Riyadh City. The raw water was collected from different wells in Skaka, Al Jouf, Saudi Arabia. The characteristics of raw water are illustrated in Table 1. All the experiments were conducted without prior modifications of samples.

<table>
<thead>
<tr>
<th>Temperature (\degree\text{C})</th>
<th>pH</th>
<th>Turbidity (NTU)</th>
<th>UV(_\text{254}) (cm(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>20 - 24</td>
<td>7.0 - 7.8</td>
<td>4.0 - 9.5</td>
<td>0.05 - 0.44</td>
</tr>
<tr>
<td>Fe ((\text{mg.} lt^{-1}))</td>
<td>T.D.S ((\text{mg.} lt^{-1}))</td>
<td>Mn ((\text{mg.} lt^{-1}))</td>
<td>Conductivity ((\mu\text{S.cm}^{-1}))</td>
</tr>
<tr>
<td>0.13 - 1.72</td>
<td>975 - 989</td>
<td>0.1</td>
<td>1400 - 1408</td>
</tr>
</tbody>
</table>

B. Apparatuses and instruments

Stirrer 1: AGIMATIC AD-C magnetic Stirrer, Stirrer 2: ARE Heating magnetic Stirrer VELP scientific, Refrigerator Deodorizer SAMSUNG, Blender GEEPAS, Whatman filter paper grade 1 and Packers of different sizes from 200 ml to 1 liter. Digit heat JP Selecta serial 0494926 was used to burn samples. HACH SENSION3 pH meter was used to measure pH. Ultra-meter PI 6P MYRON L company was used to measure Conductivity \((\mu\text{S.cm}^{-1})\), T.D.S \((\text{mg.} lt^{-1})\) and Temperature \((\degree\text{C})\). Analytical Balance GR-120 MODEL A&D company was used to determine weight, 2100 P turbidimeter HACH company was used to measure turbidity (NTU). The bandgap of synthesized MFH was determined by using Spectrophotometer DR 5000 HACH company. A Brunauer-Emmett-Teller (BET) analysis was performed to determine the surface area and pore size for flaxseed husk (FH), which is done using the automatic Belsorp-max device using liquid adsorption N2 at a temperature of 77 K [32].

To determine the functional group, FITR Fourier transform infrared spectrometry test performed on A PerkinElmer "Spectrum BX" spectrometer in the range of 4000-400 cm\(^{-1}\) using KBr pellets for sample preparation was used. A Brunauer-Emmett-Teller (BET) analysis was performed to determine the surface area and pore size for flaxseed husk (FH), which was done using the automatic Belsorp-max device using liquid adsorption N2 at a temperature of 77 K [32].

III. Extraction and Experimental procedure

In this section, the process of the preparation modified flaxseed husk (MFH) from raw flaxseed was illustrated, including the removing of mucilage and oil and improving the flaxseed husk characteristic by blending it with aluminum sulphate (AS). The preparation of the flaxseed husk ash (FHA) by burning flaxseed husk at certain temperatures \((75, 100, 150, 200 \degree\text{C})\), and experimental procedures which were used to remove turbidity, natural organic matter and iron were illustrated.

A. Preparation of modified flaxseed husk (MFH)

1) Remove mucilage and oil from seeds
To remove the flaxseed mucilage, first, 50 milligrams of flaxseed were weighed and washed with distilled water to remove dust. Then, half a mole \((0.5 \text{ M})\) of sodium bicarbonate Na HCO\(_3\) \((1:8 \text{ w/v}, 40 \degree\text{C})\) was put in a half-liter baker. Then, the washed seeds were added to the sodium bicarbonate. After that, the baker was placed on the magnetic stirrer, set at 40 \degree\text{C} for an hour, then filtered with a cloth filter to separate the liquid (mucilage) for disposal. The solid which is left (flaxseed without mucilage) was washed several times with distilled water, dried and kept at room temperature \((24 \degree\text{C})\) for 24 hours. This new product (flaxseed without mucilage) is called (F-OUT m) [32].

After removing the mucilage, the oil was removed from the flaxseed (F-OUT M) by the following steps: The seeds (F-OUT M) were crushed by a laboratory mill and the n-hexane was added in a 1-liter baker \((1:5, \text{w/v})\) and placed on the magnetic stirrer for 6 hours. Regeneration of n-hexane every 2 hours by paper filters. After final filtration the resulting material is left in the dark to dry at room temperature exposed to air and then stored at 4 \degree\text{C} and called (FH-OUT M, O) [32].

To determine the functional group, FITR Fourier transform infrared spectrometry test performed on A PerkinElmer "Spectrum BX" spectrometer in the range of 4000-400 cm\(^{-1}\) using KBr pellets for sample preparation was used. A Brunauer-Emmett-Teller (BET) analysis was performed to determine the surface area and pore size for flaxseed husk (FH), which was done using the automatic Belsorp-max device using liquid adsorption N2 at a temperature of 77 K [32].

2) Modification of \((FH-OUT M, O)\)
\((FH-OUT \text{ M, O})\) was then used with aluminum sulphate (AS), as a mixed coagulant with a certain dosage. 25 mg. \Lt^{-1}\) for (AS) and 50 mg. \Lt^{-1}\) for (FH-OUT \text{ M, O}) were used as a coagulants dosages specified on (Table 2) [33-35]. Operational conditions of the coagulation / flocculation and sedimentation process was used in seed correlation (FH-OUT M, O) and alum coagulants were presented in (Table 3). After the coagulation / flocculation and sedimentation process was completed, samples which were taken 3 cm under the water surface was filtered by Whatman paper filter grade 1 (pore size of 0.6 \text{μm}). Finally, the residual turbidity and NOM were quantified by 2100 P Turbidimeter Hach Co, and spectrophotometer DR 5000 Hach Co respectively [36,37].
Fig. 1. The FTIR analysis of FH

**TABLE 2**

<table>
<thead>
<tr>
<th>POINT</th>
<th>% Coagulant (Al₂(SO₄)₃·16H₂O/FH-OUTM₀)</th>
<th>Coagulant dosage (mg.l⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0%, 100%</td>
<td>Al₂(SO₄)₃·16H₂O</td>
</tr>
<tr>
<td>2</td>
<td>20%, 80%</td>
<td>5</td>
</tr>
<tr>
<td>3</td>
<td>40%, 60%</td>
<td>10</td>
</tr>
<tr>
<td>4</td>
<td>60%, 40%</td>
<td>15</td>
</tr>
<tr>
<td>5</td>
<td>80%, 20%</td>
<td>20</td>
</tr>
<tr>
<td>6</td>
<td>100%, 0%</td>
<td>25</td>
</tr>
</tbody>
</table>

**B. Experimental procedure and the analytical measurements of (MFH)**

The hybrid process including coagulation and adsorption was carried out as follows: first: coagulation followed by adsorption, second: adsorption followed by coagulation.

Coagulation and subsequent adsorption tests were performed after collection of samples after sedimentation, and then different doses of MFH were added. Adsorption followed by coagulation experiments were executed by adding a certain dosage of coagulants with mixing stirring similar to coagulation experiments. Then, a sample was taken using the same procedure [38].

1000 ml of raw water was transferred into a beaker with a certain dosage of MFH. The process included 3 min of the rapid mix at 350 rpm, 30 min of slow mixing at 60 rpm for flocculation, and 30 min for settling. Adsorption experiments were executed by mixing MFH with raw water using a magnetic stirrer at 150 rpm for 120 min without heat. Subsequently, MFH was removed by filtration through a what man filter paper grad 1 (pore size of 0.6 μm) for measurements, according to [39,40] with modification.

**V. Results and discussion**

**A. Characterization of the flaxseed husk (FH)**

The specific surface area of FH was 125.18 m²/g. The average pore diameter of FH was 26.28 μm indicating reasonable adsorption capacity [41].

**TABLE 3**

| OPERATIONAL CONDITIONS OF COAGULATION/FLOCCULATION AND SEDIIMENTATION PROCESS FOR POINT 3,4 - RMV: RAPID MIXING VELOCITY; RMT: RAPID MIXING TIMES; SMV: SLOW MIXING VELOCITY; SMT: SLOW MIXING TIMES; ST: SEDIMENTATION TIMES. |
|-----------------------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| **ASSAY** | 1 | 2 | 3 | 4 | 5 | 6 |
| **RMV (rpm)** | 200 | 300 | 350 | 250 | 300 | 350 |
| **RMT (min)** | 1 | 1 | 1 | 1 | 3 | 3 |
| **SMV (rpm)** | 60 | 60 | 60 | 60 | 60 | 60 |
| **SMT (min)** | 15 | 20 | 30 | 15 | 20 | 30 |
| **ST (min)** | 15 | 15 | 15 | 30 | 30 | 30 |

**IV. Preparation of flaxseed husk ash (FHA)**

**A. Preparation of flaxseed husk ash (FHA)**

C. The sample of by-product flaxseed oil removal was dried in direct sunlight for one week, then washed with distilled water, dried, ground and taken to burn at 75,100,150 and 200 °C for 2 hours.

**Fig. 2.** shows the functional groups which were done by FTIR spectrometry test, detected by the band intensity. Ketone and alkane groups (C-H) are at band intensity 2850 cm⁻¹ and 2925 cm⁻¹, and aromatic cyclic groups are at band intensity 1650 cm⁻¹ and 1750 cm⁻¹. The intense vibration at 1350 cm⁻¹ gives evidence of a large number of amino groups in flaxseed husk (FH) structure [38].

**B. Characterization of the (FHA)**

**Fig. 2.** shows the functional groups which were done by FTIR Fourier transform infrared spectrometry analysis point out several main peaks at 3420, 2920, 1750 and 1170 cm⁻¹ were associated with -OH groups, -CH, C O bands and -OCH₃ groups [42]. The presence of these active groups on the surface of flax seeds indicates their participation in the removal and absorption of iron from drinking water.
C. Removal of Turbidity and NOM

1) Effect of different concentration of AS / FH-OUTM, O

The removal efficacies of turbidity and NOM at different percentages of AS / FH-OUTM, O are shown in Fig. 3. The higher percentage of AS leads to a better turbidity removal due to the abundance of dissolved positively charged aluminum ions which neutralize the suspended particulates and increase the potential of agglomeration and floc formation [43]. On the other hand, the optimal ratio of (AS/FH-OUTM, O) for NOM removal was (40/60) achieved a removal efficiency of 81%. The higher percentage of AS leads to smaller surface area of coagulant/polymer which inhibit the adsorption of NOM on FH-OUTM, O. In addition, the lower percentage of AS decreases the efficiency of agglomeration and precipitation of suspended particulates which reduces the NOM removal.

2) Effect of slow mixing time

The effect of slow mixing time on the removal of turbidity and NOM is shown in Fig. 4. It is clear that increasing the slow mixing time improved the removal of both turbidity and NOM. This improvement was achieved because the particles have adequate time to agglomerate and form stable flocs [44].

The maximum removal efficiency of turbidity was achieved after about 20 min of slow mixing, and additional time did not improve the removal of turbidity because the coagulant dose may be consumed at this stage and additional dosage may be required [45].

The optimum removal of NOM was achieved after 30 min of mixing, while additional time did not significantly improve the removal. Such a trend could be attributed to the consumption of aluminum ions and the occupation of all active sites on FH-OUTM, O surface [46]. Accordingly, the optimum slow mixing time is considered 30 min in this study. The adsorption of organic molecules may be attributed to the electrostatic attraction forces with amino groups of flaxseed husk which have been detected in FTIR spectra [47].

3) Effect FH-OUTM, O Dosage

Different dosages of FH-OUTM, O were tested to investigate the optimum dosage for NOM and turbidity removal as shown in Fig. 5. It is clear that increasing the dosage of more than 40 mg L⁻¹ slightly improve both turbidity and NOM removal. This result is due to agglomeration of most of suspended particulates and/or adsorption of dissolved organic matter on MFH particles [48]. Accordingly, excess FH-OUTM, O amount didn't achieve significantly efficiency improving.

4) Effect of Initial NOM

The removal efficiency of NOM at different initial concentrations was investigated and exhibited in Fig. 6. The removal efficiency results ranged from 98.4% to 100% using the optimum dosage of MFH. The removal efficiency was between 98.4 % and 100 % for initial concentration of UV254 between 0.05 to 0.18 cm⁻¹, and between 98.4 % and 99.3 % for
initial concentration of UV254 between 0.19 and 0.44 cm\(^{-1}\). The high removal efficiency was achieved due to the attractive of NOM functional groups for the high surface area of MFH which suggests a high adsorption capacity \[49\]. This finding indicates that the MFH could be used for removal of high concentration of NOM from water.

![Image](image1.png)

**Fig. 5.** The effect of different FH dose, initial turbidity = 9.5 NTU, initial UV254 = 0.4 cm\(^{-1}\), and MFH: (40% AS, variable FH-OUTM, O).

**D. Removal of Iron**

1) **Effect of FHA different types on iron removal (Fe).**

*Fig. 7 and table 4.* show the effectiveness of FHA150 and FHA 200 on iron removal, which were 72% and 85% respectively. The removal process was accomplished by adding 250 mg of FHA with 250 ml of fresh raw water sample which was taken from EL Taleem well, after 30 min stirrer operating.

2) **Effect of FHA200 dosage.**

In order to study the effect of the burned dose on iron removal efficiency, doses ranging from 0.1 to 1 g and the concentration of iron in the raw water sample of the well 0.4 mg.l\(^{-1}\) liter were used. The results of these experiments were represented in *Fig. 8.*, and the different values were expressed. From *Fig. 8.* several conclusions can be read. At first: the removal efficiency was increased rapidly up to dose 0.4 g, whereas the removal efficiency was increased slightly from dose 0.4 g to 0.75 g until reaches a roughly constant value begging from dose 0.75 g to 1 g.

![Image](image2.png)

**Fig. 6.** The effect of different concentrations at optimal condition, MFH: (40% AS 60% FH-OUT M, O).

![Image](image3.png)

**Fig. 7.** Effect different types of FHA in removal iron (Fe).

**TABLE 4**

<table>
<thead>
<tr>
<th>FHA</th>
<th>FHA 75</th>
<th>FHA 100</th>
<th>FHA150</th>
<th>FHA 200</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe values (mg.l(^{-1}))</td>
<td>0.49</td>
<td>0.32</td>
<td>0.11</td>
<td>0.06</td>
</tr>
<tr>
<td>% Removal</td>
<td>–</td>
<td>20%</td>
<td>72%</td>
<td>86%</td>
</tr>
</tbody>
</table>

This trend was expected due to the increased number of particles that have been absorbed with an increased dose of FHA200 which results in more iron attached to the surface area of FHA200. It has been observed that the largest removal efficiency is 90% starting from the dose 0.75 g of the burner (FHA200). Any other FHA200 dose was added more than 0.75-0.8 gm. l\(^{-1}\) did not cause any significant change in removal efficiency.

3) **Effect of contact time.**

The efficiency of iron removal efficiency was measured, to samples of several concentrations which were taken from different wells. After studying the results shown in *Fig. 9.* specifically, from one minute to 60 minutes, it was founded a several explanations, at the beginning of contact, the rate of iron removal increased, which gradually decreased until it
reached the stage of equilibrium, within 30 to 45 minutes. Hence, it noticed that the removal rate was become fixed from 45 minutes to 60 minutes of contact time, see Photo 1. and Photo 2. The efficiency of removal was found to be 89% to 90% after 45 minutes of contact. This explains the efficiency of the active surface of the FHA, which can adsorb iron on the outside surface. Over time, the absorbed material was moved from the outer FHA surface to the inner surface result in adsorption of an additional amount of iron.

4) Effective of initial iron concentration

The experiment was conducted at an initial iron concentration in raw water ranged from 0.13 to 1.72 mg per liter. Photo 3. shows a sample used from Morog well at several iron concentration. It was found that the higher iron concentration, the greater removal efficiency. Based on these results, can say that, the high initial concentration of iron in raw water enhances the absorption process and also enhances the interaction between the burner (FHA200) and the iron. It was also apparent that the removal efficiency has increased steadily in the low concentration of iron when comparing a several concentrations (from 0.13 to 0.72 mg. l\(^{-1}\)). Whereas, in high iron concentrations (from 0.73 to 1.72 mg. l\(^{-1}\)), the removal efficiency increases slightly. This was shown by comparing the slope of the imaginary straight-line AB with the slope of the straight-line BC in Fig. 10. The Fe removal efficiency percentage was found about 72% for 0.13 mg. l\(^{-1}\) Fe and 89% for 1.72 mg. l\(^{-1}\) Fe after 30 minutes of contact time (CT).

E. Adsorption kinetics.

Adsorption kinetics are one of the most important characteristics that determine adsorption efficiency on the surface of the adsorbent material, which describes the soluble absorption rate. Fe (II) adsorption kinetics were evaluated by applying two popular models: the pseudo- kinetics model of the first order [50] and the pseudo- kinetics model of the second order [51].

1) The pseudo- kinetics model of first order

The first pseudo-kinetic model assumes that the Fe (II) absorption rate with time is directly proportional to the number
of active sites available on the surface of the adsorbents. The first pseudo-kinetic model is given to the equation as follow:

\[ \ln (q_e - q_t) = \ln q_e - k_1 t, \]

(1)

Where \( q_e \) and \( q_t \) represent the amount of absorption (mg/g) at equilibrium case and \( t \) (time) respectively, and \( k_1 \) is the adsorption rate constant the pseudo-kinetics model of the first order (min\(^{-1}\)). The \( k_1 \) values can be specified empirically by \( \ln (q_e - q_t) \) against \( t \).

2) The pseudo-kinetics model of second order

The pseudo-kinetics model of the second order assumes that chemical absorption represents the phase of limiting absorption rate, which involves the forces of equivalence by exchanging or transferring electrons between adsorbate and absorbent materials. Pseudo-kinetics model of the second order equation is expressed as follow:

\[ \frac{t}{q_t} = \frac{1}{k_2 q_e^2} + \left( \frac{t}{q_e} \right). \]

(2)

Where: \( k_2 \) (g / mg min) is the rate constant of the second-order equation, \( q_t \) (mg / g) is the absorption time \( t \) (time) amount and \( q_e \) is the absorption equilibrium amount (mg / g).

The results showed that the most suitable model for expressing the absorption rate of Fe (II) is the pseudo-kinetics model of the second order. The correlation coefficients of adsorption using FHA for the pseudo-kinetics model of the second order (\( R^2 = 0.9999 \)) are closer to the unity than those for the pseudo-kinetics model of the first order. Fig. 12 shows the linear segment of \( t/q_t \) versus the pseudo-kinetics model of the second order for the absorption of Fe (II) ions on the FHA. The theoretical \( q_e \) values of the FHA correspond well to the experimental values as compared to those of the pseudo-kinetic model of the first order. While the theoretical values of the absorption of flaxseed husk for both ends of the two movements are well correlated with the experimental \( q_e \) values, kinetic models are well suited to the absorption process and Fe (II) chemisorption assay emphasizes by flaxseed husks.

F. Adsorption isotherm studies

The adsorption process is described by Adsorption isotherms and also describe the interaction between adsorbates and bio-sorbent. For analysis and design of adsorption systems

Photo 3. sample from Morog well Fe = 0.4 mg.l\(^{-1}\)

Fig. 10. Effect of initial concentration on the Fe removal by FHA200, (dose = 0.75 gm. per liter and CT = 45 min)

Fig. 11 shows the linear segment of \( \ln (q_e - q_t) \) versus the pseudo-kinetics model of the first order for the absorption of Fe (II) ions on the FHA. The correlation coefficients of adsorption using FHA for the pseudo-kinetics model of the first order (\( R^2 = 0.7987 \)) are closer to the unity.

![Graph](image-url)
must establish the most acceptable correlations for the batch
equilibrium data, or at equilibrium solution and constant
temperature, the Adsorption isotherms used to describe the
correlation between the number of species/ions adsorbed onto
the solid phase.

\[ \frac{t}{q_e} = \frac{1}{q_mK_L} + \left( \frac{C_e}{q_m} \right) \]  

Where: \( k_m \) is a constant that expresses the Convergence of
the obligated sites towards the adsorbent, called the Langmuir
constant (L mg\(^{-1}\)), \( q_m \) known as the monolayer uptake capacity
or the quantity of absorbed pores per unit mass of adsorbents
that form a complete monolayer on the outward surface (mg g\(^{-1}\)), \( q_e \) is equilibrium concentration of Fe(II) on the bio-
sorbent (mg g\(^{-1}\)) and \( C_e \) is the concentration of adsorbed substance
at equilibrium or adsorbate concentration (mg g\(^{-1}\)).

Langmuir isotherm plots for the bio-sorption of Fe (II)
showed in Fig. 13, which demonstrate the degradation plot of
\( C_e / q_e \) versus \( C_e \) the Langmuir isotherm curve shows R2:

\[ R^2 = 0.9982 \]

It can be expressed in the following equation:

\[ q_e = K_F \cdot C_e^{1/n} \]  

where \( K_F \) is Freundlich distribution coefficient or
Freundlich constant, represent uptake capacity of iron onto
FHA ((mg g\(^{-1}\) (L mg\(^{-1}\))1/n). Can express about uptake intensity by empirical coefficient 1/n, according to the 1/n value, can determine the favorability of adsorption. At 1/n
between 0 and 1 means favorable isotherm, at 1/n= 0 means irreversible, at 1/n > 1 means unfavorable uptake, the smaller value of 1/n means stronger interaction occurs between FHA and iron [57].

A Freundlich regression plot of ln(qe) versus ln (Ce) is shown in Fig. 14, in which the slope of the line (1/n) and the other value associated with it (Kf) is shown. From the figure it turns out that the slope value is 0.264 less than the correct one, so the absorption is classified as preferred (favorable isotherm). The slope value (1/n) also illustrates several considerations: 26.4% of the active adsorption sites retained the same energy level [58], and the smaller the slope value, the more effective the adsorption, the higher the adsorption strength and the heterogeneity of hard surfaces as well. [59]

![Fig. 14. Freundlich isotherm plots for the bio-sorption of Fe(II) onto FHA200 absorbent dosage: 0.1 g/100 mL; contact time: 45 min; pH: 7.8](image)

### G. Comparison study

The (Table 5) gives a comparison between metal ions sorbents, including iron, in terms of adsorption capacity, contact time, dissolved ions concentration and temperature range [60].

<table>
<thead>
<tr>
<th>No</th>
<th>Adsorbents</th>
<th>Adsorption capacity (mg/g)</th>
<th>Contact time (min)</th>
<th>Concentration range (mg/L)</th>
<th>Temp. range (K)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Cross-linked chitosan</td>
<td>64.1</td>
<td>60</td>
<td>3–9</td>
<td>—</td>
<td>[61]</td>
</tr>
<tr>
<td>2</td>
<td>Chitosan</td>
<td>57.5</td>
<td>40</td>
<td>3–9</td>
<td>—</td>
<td>[61]</td>
</tr>
<tr>
<td>3</td>
<td>Modified coir fibres</td>
<td>7.49</td>
<td>120</td>
<td>73.50–83.9</td>
<td>308</td>
<td>[62]</td>
</tr>
<tr>
<td>4</td>
<td>Rice husk ash</td>
<td>6.21</td>
<td>60</td>
<td>2–40</td>
<td>298</td>
<td>[63]</td>
</tr>
<tr>
<td>5</td>
<td>Flaxseed husk ash</td>
<td>6.19</td>
<td>45</td>
<td>0.4–1.72</td>
<td>296</td>
<td>Present study</td>
</tr>
<tr>
<td>6</td>
<td>Coir fibres</td>
<td>2.84</td>
<td>120</td>
<td>73.50–83.9</td>
<td>308</td>
<td>[62]</td>
</tr>
<tr>
<td>7</td>
<td>Pine bark wastes</td>
<td>0.03</td>
<td>30</td>
<td>55.6–111.2</td>
<td>303–333</td>
<td>[64]</td>
</tr>
</tbody>
</table>

The optimal ratio of (AS / FH-OUT) for removing both NOM and turbidity was (40% / 80%). The removal efficiency for NOM and turbidity at ratio (40% / 60%) was 81% and 93%. Increasing the dosage of MFH to (40% / 80%), significantly improved the removal efficiency to 98.0% for turbidity and 99.3% for NOM. Moreover, the MFH revealed a high removal efficiency at a higher initial NOM value which suggests the favorability for treatment of raw water contaminated with NOM.

In this study, many experiments have been carried out to investigate the performance of FHA to remove iron from aqueous solutions under various operational conditions. The removed iron by flaxseed husk ash was found to be depending on different types of FHA, Initial iron concentration, contact time, and dosage of the FHA. The initial pH values ranged between 7.0 to 7.8, Which did not cause any significant change in the pH after adding of the FHA200. It may be noted that the efficiency of de-iron removal of water solution is directly proportional to increase of adsorbent material, initial iron concentration and contact time. Maximum removal efficiency of 86-90 % was observed at adsorbent dosage 0.75 gm.l⁻¹. The FTIR spectra of FHA shows effective groups on the surface of FHA involved in iron removal from drinking water. Adsorption kinetics was used to analyze the equilibrium data. The experimental data of biosorption isotherms for the biosorbent yielded fitted well with the kinetics order, the pseudo- kinetics model of the first order and the pseudo-kinetics model of the second order. The results of the adsorption equilibrium by Langmuir and Freundlich model showed FHA’s ability to absorb iron ions at a high degree, hence, this was expressed in favorable. On the other hand, the correlation coefficient of the Langmuir and Freundlich models were 0.998 and 0.897, respectively, shows that the relationship between the absorbed iron ions and the FHA concentration in this study with the Langmuir model is better than that of the Freundlich model because the correlation coefficient in the
Langmuir model is greater than the correlation coefficient in the Freundlich model.

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[27] Water quality standards, Presidency of Meteorology and Environmental Protection (PME), Saudi Arabia

[28] GCC STANDARDIZATION ORGANIZATION (GSO), GSO/ FDS / 2012, Bottled drinking water


Title Arabic:
تحسين خصائص مياه الشرب باستخدام مادة مازة حيوية مبتكرة

Arabic Abstract:
في هذه البحث كشفنا عن مادة جديدة في عالم المخثرات والمواد المازة الطبيعية تحصل خصائص مياه الشرب وقد أثبتت فعاليتها مقارنة بالموارد الطبيعية المبتكرة. تم استخدام قشر بذور الكتان كمادة مازة حيوية وكمادة مازة لإزالة التكعك والمواد العضوية الطبيعية والدينار من المياه الجوفية لعده أربع مناطق الحوض شمال المملكة العربية السعودية. لإزالة التكعك والمواد العضوية الطبيعية (AS) تم تحميل قشر بذور الكتان المعالج (MFH) إلى قشر بذور الكتان المعالج (FH) إلى قشر بذور الكتان المعالج (FH) لإزالة المادة العضوية والزيت. يظهر توصيف (FH) قشر بذور الكتان المعالج (FH) سهولة نقل عالية (125.18 متر مكعب / جم) بقطر مسال بلغ 26.28 متر مكعب / جم. علاوة على ذلك أشار تحليل أطياف FTIR إلى وجود عديد كبير من المجموعات الأمينية ومجموعات وظيفية أخرى تحتوي على كفاءة إزالة التكعك والمواد العضوية الطبيعية. وكانت نسبة الخلط المثلى للإزالةFH = 99.3، وكانت فائدة الإزالة49.8٪ من المواد العضوية الطبيعية في المياه الغير مصبوغة. أيضا تم تخفيض نسبة الحديد في مياه الشرب عن طريق رماد قشر بذور الكتان (FHA) بعد إزالة المادة الصمغية والزيت. يُظهر توصيف (FHA) نسبة الخلط المثلى FH = 99.3، وكانت فائدة الإزالة49.8٪ من المواد العضوية الطبيعية في المياه الغير مصبوغة. أيضا تم تخفيض نسبة الحديد في مياه الشرب عن طريق رماد قشر بذور الكتان (FHA) بعد إزالة المادة الصمغية والزيت. يُظهر توصيف (FHA) نسبة الخلط المثلى FH = 99.3، وكانت فائدة الإزالة49.8٪ من المواد العضوية الطبيعية في المياه الغير مصبوغة. أيضا تم تخفيض نسبة الحديد في مياه الشرب عن طريق رماد قشر بذور الكتان (FHA) بعد إزالة المادة الصمغية والزيت. يُظهر توصيف (FHA) نسبة الخلط المثلى FH = 99.3.