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Treatment of oil and grease in wastewater using Electrocoagulation

معالجه الزيوت والشحوم في المخلفات السائلة باستخدام الترويب الكهربي

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الملخص

تم إجراء تجارب الترويب الكهربي بنظام الدفعة لتقييم أزاله الأكسجين الكيميائي الممتص والزيوت والشحوم من المخلفات السائلة باستخدام نو عين من الأقطاب الكهربية : الألومونيوم ، الحديد ومن ثم خليط من الألومونيوم والحديد . تم دراسة تأثير معايير التشغيل مثل شده التيار ، التركيز الابتدائي للأكسجين الكيميائي الممتص، ومدة التفاعل وتركيز كلوريد الصوديوم على كفاءة أزاله الأكسجين الكيميائي الممتص والزيوت والشحوم. وقد وجد من البحث أن زيادة شده التيار زاد من كفاءة أزاله الأكسجين الكيميائي الممتص. تركيز الأكسجين الكيميائي الممتص الابتدائي كان له تأثير قليل على نسب الإزالة. النتائج أظهرت أن أفضل أداء تم تحقيقه كان باستخدام أقطاب الألومونيوم عند تيار شدته 1.3 أمبير و 11.6 فولت مج/لتر و تركيز كلوريد الصوديوم 0,50 جم /لتر. 1500مع مدة تفاعل 15 دقيقه وتركيز أكسجين كيميائي ممتص ابتدائي تحت هذه الظروف، كفاءة أزاله الأكسجين الكيميائي الممتص وصلت إلى 95.07 % والتكلفة الكلية تعادل 0,84 متص يتم أزالته.

Abstract

Batch electrocoagulation (EC) experiments were carried out to evaluate the removal of COD and O&G from wastewater using two types of electrodes: aluminum, iron, and then mix of aluminum and iron. The effects of operating parameters such as current intensity initial COD concentration, contact time and NaCl concentration on COD and O&G removal efficiency had been investigated. It was found that increasing current intensity increased COD and O&G removal efficiency. Initial COD concentration had a little effect on removal efficiencies. Results showed that the best performance was obtained using aluminum electrode at a current intensity of 1.3 Ampere and 11.6 volt in 15 min contact time for 1500 mg/L initial COD concentration and 0.5 gm/L NaCl concentration. Under these conditions, COD removal efficiency reached 95.07% and total cost was 0 .84 $LE/m^3/gm$ COD removal.

Key words:

Electrocoagulation, Electrodes, Oil and grease and COD removal efficiency.

1. Introduction

O&G contained The in the wastewater aggregate and foul the sewer system and produce unpleasant odor (2) problematic .Environmentally wastewaters are produced in large volumes every day. O&G, are common pollutants found in wastes from a vast range of industries such as petroleum refineries, petrochemical, metal manufacture, machining and finishing, food processors, textile and carwash stations(3).

Main pollutant in oilfield wastewater is oil which may range between 100 and

1000 mg/l or still higher depending on the efficiency of emulsification and nature of raw oil (4). Oily wastewaters can be classification into three categories: freefloating oil, unstable oil/water emulsions, and highly stable (steady) oil/ water emulsions. Free floating oil can be readily removed by mechanical separation whilst unstable and stable oil-water (O/W) emulsions must be mechanically or chemically broken and detached gravitationally (3).

Carwash wastewaters can be harmful to humans and environment if released

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without treatment to surface water bodies as they contain a lot of pollutants such as detergents, oil, grease, solvent-based solutions, road grime, heavy metals, etc., that can be toxic to fish, organisms and the accumulation of these pollutant species in the aqueous bodies leads to undesirable effects on human life and on the environment (5).

The environmental requirements in the Egyptian law 93 for year 1962 and the modified law 44 for year 2000 recognize that water should be spent on sewage system does not exceed COD 1100 mg/l, oil-grease 100 mg/l and pH be in the range between 6 to 9.5.

Although there were a lot of techniques available, including a variety of filters, gravity separation, air flotation, biological process, membrane bioreactor, carbon adsorption, chemical coagulation, electrocoagulation, electro flotation etc., for separation of oil—water emulsions, some of the researchers stated that there is still a lack of efficient universal technique in treating oily wastewaters (4, 6).

There are two kinds of coagulation: conventional coagulation and electrocoagulation. Conventional (chemical) coagulation refers to addition of chemicals such as Alum [Al2 (SO₄)₃ . 18H₂O] to an aqueous solution to join small dispersed particles into bigger agglomerates which can be removed by some other method such as sedimentation, floatation, filtration air or (3,7).Conventional coagulation involves number of drawbacks such as the high amount of require coagulant, corrosion problems with reducing pH and problems with produced sludge (3).

The rule of electrocoagulation (EC) has been popular knowledge for over 120 years with electrochemical methods first being used for water and wastewater treatment in 1887. In 1906 Dietrich invented the first electric water purifier which used aluminum electrodes whilst J.T. Harries received a patent in 1909 for wastewater treatment by electrolysis with

sacrificial aluminum and iron anodes (3). The electrocoagulation of drinking water was first implemented on a large scale in the US in 1946(8).

The EC unit is environmentally friendly so that it does not produce corrosion or any pollutants. This technique has some merits when compared to conventional methods such as simple equipment, easy to apply, less retention time required and less sludge production (5, 9). Furthermore, Electrocoagulation is efficient in removing suspended solids as well as O&G. Many investigators found that it removes metals, colloidal solids, particles containing arsenic, dyes, paper mills, breaking oil emulsions in water, phosphate, boron and bacteria, viruses and cysts $(9 \rightarrow 11)$.

The nature of the electrode material is a main issue in electrochemical treatment and the appropriate selection of electrode material is very essential. The most popular electrode materials used for electrocoagulation technique are aluminum and iron because they are cheap, readily available and proven effective since their dissolution in aerated media produce trivalent species (12).

In the literature it was found that the iron electrode was more effective than aluminum (2, 13). On the other hand, Rupesh et al. said that oil removal efficiency was 90% at 4.72 pH within 30 treatment time for concentration of oil and 94.44% of oil removed in 30 min at 4 mg/l of salinity using Aluminum electrode (4). In the second place, Fouad treated oil-water emulsions, In his work the initial O&G concentrations investigated were 200, 300 and 400 ppm, pH values used were 3, 5, 7, 9 and 11(11). Maha et al., treated oil tanning effluent and proved that iron (Fe) and aluminum (Al) electrodes were considered as sacrificial electrodes in different combinations. They found that under best operating conditions such as 20 mA/cm^2 current density. 5% concentration, 1.0 g/l NaCl concentration

and Fe/Fe electrodes, % COD removal, energy consumption and operating cost were 89.65%, 1.279 kW h/m3 and 6.28 US \$/m3 ,respectively (15) . Bensadok et al., (2011) focused on study the effect of NaCl concentration between 0.5 and 2.0 g/L on the removal efficiency and they proved that the greatest removal efficiency was obtained with the use of both cathode and anode made of aluminum (Al–Al system) (12).

This paper is primarily aimed at development the technology of O&G removal by electrocoagulation. In the present study, the effect of numerous operating conditions such as electrode material (iron or aluminum), initial COD concentration, current intensity, NaCl concentration, and contact time on the removal efficiency of COD have explored and discussed to identify the optimum operational conditions and low cost.

2. Theory of Electrocoagulation

As shown in Fig. 1, aluminum or iron is usually used as electrodes and upon the application of a direct current their cations are produced by dissolution of sacrificial anodes. metal The produced are hydrolyzed in the electrochemical cell to create metal hydroxide ions according to anodic and solution reactions and the solubility of the metal hydroxide complexes formed depends on pH and ionic power. Insoluble flocs are produced at pH range between 6.0 and 7.0. Positive metal species react with negatively charged particles in the water to form destabilized colloids and then flocs. The in situ generation of coagulants means that electrocoagulation processes do not require the addition of any chemicals. The gases generated at the cathode during the electrolysis of water metal dissolution according cathodic reaction permit the resulting flocs to float and it may be removed by any skimming technique (4, 5, 7, 9, 16, 17).

The oxidation - reduction reactions involved in the electrochemical cell are as follow:

Anodic (oxidation) reactions:

$$Al_{(s)} \to Al_{(aq)}^{+3} + 3 e^{-}$$

 $Fe_{(s)} \to Fe_{(aq)}^{+2} + 2 e^{-}$
 $Fe_{(aq)}^{+2} + 2 OH_{(aq)}^{-} \to Fe(OH)_{2(s)}$

Cathodic (reduction) reactions:

$$2 H_2O + 2 e^- \rightarrow H_2(g) + 2 OH_{(ag)}^-$$

In the solution:

$$AL_{(aq)}^{+3} + 3 H_2O \rightarrow Al(OH)_{3(s)} + 3 H_{(aq)}^+$$

 $Fe_{(aq)}^{+3} + H_2O \rightarrow Fe(OH)_{(aq)}^{+2} + 2 H_{(aq)}^+$

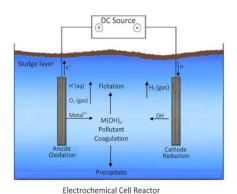


Fig. 1

When the concentration of NaCl salt in solution increased, conductivity of the solution and the current intensity increased. The higher ionic power would generally cause an increase in current intensity at the same cell voltage (8). It was that the addition of significantly boosted the oil removal but further increase in NaCl decreased the oil removal (4). The presence of NaCl reduced the size of gas bubbles, especially hydrogen Gas. Since the buoyancy of smaller bubbles was lower than bigger bubbles, they rise slowly to the surface with high opportunities for collision with oil drops. This leads to a progress in the oil removal process (4).

When chlorides were presented in the solutions the products from anodic discharge of chlorides were Cl_2 and OCl^- . The OCl^- itself is a strong oxidant, which capable of oxidizing organic molecules present in wastewater (9). It decreased passivation effect and raised the current efficiency. The following reactions (1, 2, 3) explain the formation of hypochlorite (15).

When we add NaCl there are three equations:

$$2 Cl^{-} - 2e \rightarrow Cl_{2}$$
 (1)
 $Cl_{2} + H_{2}O \rightarrow HOCl + Cl^{-} + H^{+}$ (2)

$$HOCl \leftrightarrow OCl^- + H^+$$
 (3)

The quantity of electrode material dissolved or consumed during the electrocoagulation process depended heavily on the current intensity as explained by the Faraday's law as follow (5,7,8).

$$m = \frac{M I t}{n F vol}$$

where : m is the mass of dissolved metal (gm/L); M is the molecular weight (g/mol) (M_{Al} =26.98 g/mol; M_{Fe} = 56 g/mol); I is the current intensity (Ampere); t is the contact time (second); n is the number of electrons involved in the oxidation reduction reaction (n_{Al} =3 and n_{Fe} =2); and F is the Faraday's constant (96,485 C/mol)and vol is the sample volume (liter).

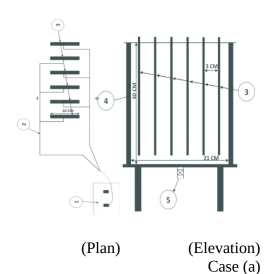
3. Materials and methods 3.1. Experimental set-up

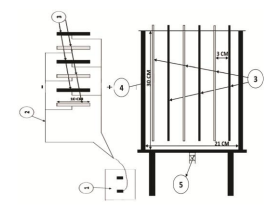
Experiments were implemented in a batch electrochemical rectangular glass cell shown in Fig. 2 that had the following dimensions: 21 cm long, 15 cm wide, and 30 cm height. The total volume of 6 liters of wastewater was treated in the electrochemical cell with 20 cm wetted depth and 10 cm free board. Six parallel plate electrodes made of rectangular metal sheets with dimensions of 30 cm height, 10 cm wide, and 1 mm thickness were

used. Three electrodes of them were connected parallel as anodes and the other connected as cathodes. electrodes connected in a monopolar mode separated by a space of 3 cm and an immersed height of 16 cm (because if the spacing between electrodes <10 mm it impeded movement of liquid hindering removal efficiency) (18). The electrode gap was kept constant in all experiments. The immersed area of one $160cm^{2}$. The electrode was electrodes were dropped to the wastewater sample and connected to digital multi meter, KEW SNAP model-2007, for measurement the current and the potential between the electrodes. The D.C. power supply output had three different current conditions: 1.3 A, 1.6 A and 1.9 A with the volts of 11.6 V, 14 V and 16.4V, respectively.

3.2. Synthetic oil—water emulsions:

Emulsions were set by adding dosages of dirty vehicles motor oil (Mobil Oil) to 1 liter of tap water and violently mixed for 3 min. The mixture showed a uniform white color.





(Plan) (Elevation) Case (b)

Fig. 2.A schematic diagram: (1) DC power supply, (2) Anode and cathode connections, (3) six electrodes of aluminum or iron (case a) or three electrodes of aluminum and three electrodes of iron (case b), (4) electrocoagulation cell, and (5) drainage valve.

After preparation the emulsion, it was left for 30 min to observe its stability. The emulsion was prepared by initial oil and grease concentrations of 400 mg/l, 550 mg/l and 700 mg/l that gave initial COD 1100 mg/l, 1280 mg/l and 1500 mg/l respectively. Therefore COD value was taken as a measure of O&G. The initial conductivity ranged from 499 µs to 546µs. In order to study the effect of conductivity, NaCl was added to the Emulsion in the last group experiments. The concentrations in that experiments were 0.50, 1.00, 1.50 mg/l. The initial pH ranged from 5.9 to 8.2.

3.3. Experimental method

First EC-cell was filled with the synthetic —water emulsion. Electrodes were submerged and then the current was passed by the regulated DC power supply. The reaction was timed, beginning when the D.C. power supply was switched on. The electrodes polarity was changed at time intervals of 15 min. Samples of 5 ml of oil—water emulsion were withdrawn from the depth of 5 cm below the free surface of oil—water emulsion at regular time intervals of 15 min. The effect of the electrochemical treatment was determined by measuring COD at the regular time

intervals of 15 min. After each run the electrodes were cleaned and rinsed with HCl (10%concentration) to remove the oxides formed at the anode surface and then dried.

Anodic dissolution was determined by measuring weight of sacrificial anode before and after experiments.

3.4. Analytical measurement

The experimental parameters measured were COD, O&G concentration, conductivity, TDS and pH. Analysis was carried out by the standard method for the examination of water and wastewater (22nd edition, 2012) (1).

The COD was measured by the closed reflux, colorimetric method and O&G was measured by hexane extractable method according to standard methods (1). The removal efficiency was determined as $(C_0 - C)/C_0$.

In order to accomplish the aims of this study ten sets of runs were planned. Each set of them contained experiments with different calculated O&G concentration (400,550. mg/lit.). The first three sets were conducted with Al electrodes (Al/Al) under the previously mentioned output current conditions of D.C. power supply. The second and third three sets similarly were conducted with (Fe/Fe) and with combination of (Al / Fe) electrodes. The last set of experiments was conducted with emulsion samples of calculated COD concentration of 1500 mg/lit and NaCl concentration of 0.50, 1.00, 1.50 mg/lit. With Al electrodes and with current of 1.3 A and 11.6V.

4. Result and discussion

4.1. Effect of electrode material & current density

4.1.1. Aluminum electrodes (Al/Al system)

Results of the first 3 sets of runs with Al electrodes were depicted in figures (3 -5). From it can be noticed that after 60 min, for initial COD concentration of 1500

mg/l the percentage of COD removal was 52.87%, at 11.6 volt and 1.3 ampere, 84.87%, at 14 volt and 1.6 ampere and was 95.73 % at 16.4 volt and 1.9 ampere. As shown in figures (3 - 5), the COD removal efficiencies after 60 min. were 47.63 – 52.87 % at current of 1.3 A, 82.36-84.87 % at 1.6 A and 90.73-95.73 % at 1.9 A.

4.1.2. Iron electrodes (Fe/Fe system)

Results of the second 3 sets of runs with Fe electrodes were depicted in figures (6-8). From it can be noticed that after 60 min, for initial COD concentration of 1280 mg/l the percentage COD removal was 83.13%, at 11.6 volt and 1.3 ampere, 97.5%, at 14 volt and 1.6 ampere and was 98.6 % at 16.4 volt and 1.9 ampere ,after 45 min. As shown in figures (6-8), the COD removal efficiencies after 60 min. was 80.87 – 84.1 % at current of 1.3 A, 96.1 - 97.9 % at 1.6 A and after 45 min. was 97.9 - 98.6 % at 1.9 A.

4.1.3. Mix of aluminum and iron electrodes (Al/Fe system)

Results of the third 3 sets of runs with Al & Fe electrodes were depicted in figures (9 -11). From it can be noticed that after 60 min, for initial COD concentration of 1100 mg/l the percentage COD removal was 96.63%, at 11.6 volt and 1.3 ampere, 97.18%, at 14 volt and 1.6 ampere and was 98.36 % at 16.4 volt and 1.9 ampere . As shown in figures (9), the COD removal efficiency after 60 min. was 95.47 – 96.63 % at current of 1.3 A, after 45 min. was 93-94.45 % at 1.6 A and after 30 min. was 94.6- 95 % at 1.9A as shown in figure (10, 11).

As a result, iron and mix of aluminum and iron electrodes were better than aluminum electrodes. The percentage of COD was more than 90% after less time and at less current density. It is commonly noted that pH increases during an electrochemical process because the process leads to the formation of metal hydroxide according to previous equations (19).

pH increasing was in the range of 0.0 to 0.5, 3.0 to 4.5 and 0.8 to 2.7 at using aluminum electrodes, iron electrodes and mix of aluminum and iron electrodes, respectively.

At using aluminum electrodes and for emulsion of initial COD concentration of 1500 mg/l After 60 min, The COD removal efficiency increased from 52.87 % to 84.87 % by increasing the current voltage from 11.6 volt to 14 volt. When the applied current voltage was increased from 14 volt to 16.4 volt, the COD removal efficiency increased from 84.87 % to 95.73 % as seen in figure $(3\rightarrow 5)$. Furthermore at using iron electrodes, The COD removal efficiency increased from 80.87 % to 96.1% by increasing the current voltage from 11.6 volt to 14 volt. When the applied current voltage was increased from 14 volt to 16.4 volt, there was not noticeable change in COD removal efficiency as seen in figures $(6\rightarrow 8)$. There was not noticeable change in COD removal efficiency when using mix of iron and aluminum electrodes when current voltage increased from 11.6 volt to 16.4 volt as seen in figures $(9\rightarrow11)$.

It can be showed from figs (6, 9) that in the low range of current intensities, as the quantity of dissolved metal was small, the solid particles of metal hydroxide formed have of this fact, small dimension and did not permit to an efficient adsorption of the emulsions. These small particles of metal hydroxide, which remain in suspension, lead to no increase in COD removal efficiency (12).

Based on Faraday's law, increasing applied voltage (or current intensity) resulted in an increasing amount of metal hydroxide flocs for the removal of colloidal particles (17).

It was also recognized that the rate of bubble-generation increased and the bubble size decreased with increasing current intensity; both of these facts were beneficial in terms of high pollutant removal efficiency by H_2 flotation (14, 16).

The difference of conductivity between the beginning and the end of experiments increased with increasing current intensity.

Initial COD concentration had a low effect on COD removal efficiency. The COD removal efficiencies at using aluminum electrodes were less than that at using iron electrodes and mix of aluminum and iron electrodes.

4.2. Effect of contact time

The time range from 0 to 60 min was studied to show how time affected the removal efficiencies. As the time of electrolysis increased comparable changes in the removal efficiency of COD was observed.

COD removal efficiency increased with increasing contact time. According to the Faraday's law, the amount of iron or aluminum released to the EC system using Fe or Al electrodes was affected by the residence time which leads to an increase in Fe or Al ions freed to the system.

At using aluminum electrodes COD removal efficiency increased fast at first 45 min and increased slowly at last 15 min as shown in figures $(3\rightarrow 5)$. Likewise, at using iron electrodes and mix of aluminum and iron electrodes, it is shown in figures $(6\rightarrow 11)$ That COD removal efficiency increased fastly at first 30 min and increased slowly at last 30 min.

4.3 Sludge production

The effluent with aluminum electrodes was found very clear and stable, while the effluent with iron electrodes appeared brown first, and then turned black and turbid. About two thirds of the sludge floated on the top and came out from a sludge outlet, while the other third was generated after sedimentation (2).

Volume of sludge from each patch increased with increasing initial concentration and current intensity. Minimum and maximum sludge volume per patch were $63cm^3$ and $567cm^3$ for aluminum electrodes, $220.5 cm^3$ and $472.5cm^3$ for iron electrodes, $315cm^3$ and

 $504cm^3$ for mix of aluminum and iron electrodes.

4.4. Effect of salinity (NaCl)

At using emulsion of initial COD concentration of 1500 mg/l and NaCl of 0.50 mg/l in (Al/Al) system, COD removal efficiency reached to 95.07% after 15 min, at 11.6 volt and 1.3 Ampere, as shown in figure (12).

Increasing NaCL dosage to 1 gm/l and 1.5 gm/l did not effect on COD removal efficiency. Initial conductivity was 1446, 2550 and 3630 in case of 0.5 mg/l, 1.0 mg/l and 1.5 mg/l.

5. Total cost

One of the most important parameters that greatly affect the application of any technique of wastewater treatment is the cost.

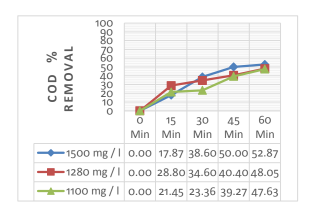


Fig. 3 COD removal with Al electrodes at 11.6 volt and 1.3 ampere.

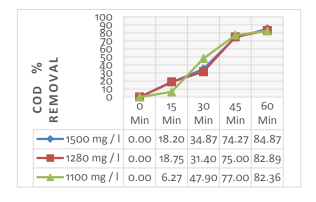


Fig. 4 COD removal with Al electrodes at 14 volt and 1.6 ampere.

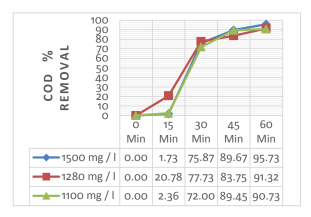


Fig. 5 COD removal with Al electrodes at 16.4 volt and 1.9 ampere.

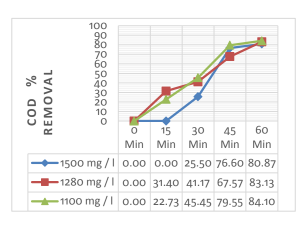


Fig. 6 COD removal with Fe electrodes at 11.6 volt and 1.3 ampere.

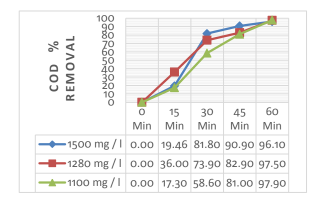


Fig. 7 COD removal with Fe electrodes at 14 volt and 1.6 ampere.

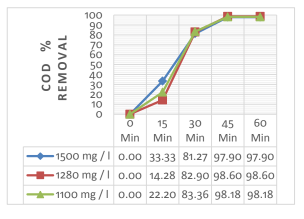


Fig. 8 COD removal with Fe electrodes at 16.4 volt and 1.9 ampere.

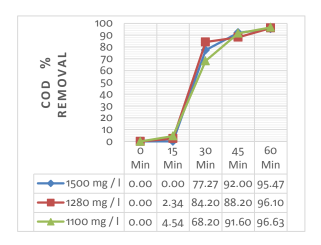


Fig. 9 COD removal with Al & Fe electrodes at 11.6 volt and 1.3 ampere

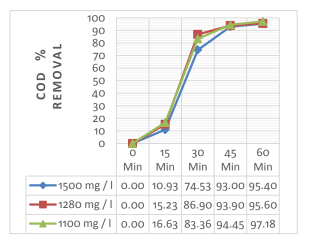


Fig. 10 COD removal with Al & Fe electrodes at 14 volt and 1.6 ampere.

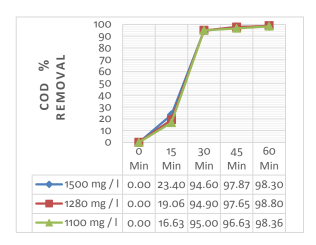


Fig. 11 COD removal with Al & Fe electrodes at 16.4 volt and 1.9 ampere.1.3 ampere.

The electrical energy consumption increased with increasing current. Since the current is a main variable in controlling the performance of the electrocoagulation, it is preferable to decrease cell voltage rather than decrease current to minimize the energy consumption (4). Operating time is the significant performance the electrocoagulation parameter in process as higher operating time results in higher energy requirement (15). Electrical energy Consumption was calculated using the following equation:

$$E = \frac{\text{VIt}}{(\text{cod}_{i\text{-}}\text{cod}_{f})\text{vol}}$$

Where E is the energy consumption (kwh/g COD); V is the current voltage (volt); I is the current intensity (ampere); t is the contact time (hour); COD_i, COD_f is initial and final chemical oxygen demand (mg/l) respectively; and vol is the sample volume (liter).

Total Cost (LE/g COD removal/ m^3) =E (kwh/g COD) ×Price (LE/kwh) + m (kg/ m^3) ×metal Price (LE/kg) + salt price (LE/ m^3)

Where the price of kwh of electricity equals 0.25 LE; price of kg aluminum equals 31.75 LE; Price of kg Iron equals 6.85 LE and price of NaCL; for 0.5 gm/l equals 0.16 LE; for 1.0 gm/l equals 0.32 LE; for 1.5 gm/l equals 0.48 LE.

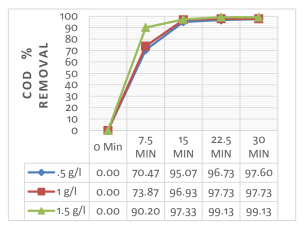


Fig. 12 Effect of NaCl dosage on COD removal with Al electrodes at 11.6 volt and 1.3 ampere for 1500 mg/l initial COD concentration.

Total costs of cubic meter for each gm COD removal after different interval times were calculated, the low cost results were at 11.6 volt and 1.3 Ampere, for initial COD concentration of 1500 mg/l, adding .5 gm/l NaCl using aluminum electrodes. Low cost was .51 LE/g COD removal/ m^3 , after 7.5 min with removal efficiency of 70.74% and it was 0.84 LE/g COD removal/ m^3 , after 15 min with removal efficiency of 95.07%.

Conclusion

- 1) The results of this study showed that electrocoagulation could be applied in the treatment of oily wastewater.
- 2) At using (Al / Al) system the COD removal efficiency after 60 min. was 90.73 95.73 % at current intensity of 1.9 A. Under the same current intensity, it was 97.9 98.6 % after 45 min. using (Fe / Fe system) and 98.3 98.8 % after 60 min. using (Al / Fe) system.
- 3) The current intensity is an important operating factor influencing the performance of electrocoagulation process and initial COD concentration has a little effect.
- 4) Results showed that the best performance was obtained using aluminum electrode at a current intensity of 1.3 A and 11.6 volt in 15 min contact time for 1500 mg/L initial

COD concentration and 0.5 mg/L NaCL concentration. Under these conditions, COD removal efficiency reached 95.07% and total cost was 0.84 $LE/m^3/gm$ COD removal.

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