Original Article





Kinetic study of the regeneration of spent caustic via the genetic algorithm method

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Abstract

Background: Spent caustic contains noxious components such as sulfide species and also high chemical oxygen demand content (COD). Oxidation of these materials to caustic and sulfate species is mostly the rate-controlling step within catalytic oxidation of spent caustic.

Methods: In this study, the kinetics of catalytic oxidation of spent caustic and the regeneration methodology of the sulfidic spent caustic were investigated. The kinetics of catalytic oxidation of spent caustic was studied in the presence of a heterogeneous catalyst. The developed mathematical model was verified via the batch bubble column reactor. The elementary and non-elementary models based on the genetic algorithm were used to obtain the rate coefficient and kinetic order.

Results: The experiments were carried out at various conditions. The results indicated that the error of objective function of the non-elementary and elementary models was 3.01% and 134.96%, respectively. **Conclusion:** According to the results, the non-elementary model had rational outcome compared to the elementary one. Also, non-elemental model is more concordance with experimental results. **Keywords:** Caustic, Kinetic, Regeneration, Catalysis

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Introduction

Chemical industries are among those fields which create various kinds of organic and inorganic waste in varying concentrations that must be treated. The contamination of these wastes is not only limited to water, air, soil and noise, but also has extended to the human's life (1,2). The petrochemical plants are among those industries that generate all kinds of wastewater. The aqueous caustic solutions are usually applied for the removal of impurities, like sulfur compounds and carbon dioxide in petrochemical plants such as the olefin unit (3). The effluent of caustic scrubbing as a waste solution is known as spent caustic. Spent caustic is a highly deadly and odorous waste. According to the US Environmental Protection Agency (EPA) reports, spent caustic is categorized as a hazardous and deleterious waste in the olefin unit (4). Table 1 shows some characteristics of spent caustic from the olefin unit in a petrochemical plant.

There are a couple of methods for treating spent caustic, such as incineration, disposal and oxidation methods, among which chemical oxidation is very practical and applicable (5). Some prominent methods are wet air oxidation (WAO), advanced oxidation process (AOP) and catalytic oxidation process (COP) that have been applied to treat spent caustic wastewater (6-14). WAO is carried out at high temperature and pressure depending on feed. Often, WAO is applied for pre-treating of spent caustic and it cannot lower the chemical oxygen demand (COD) of spent caustic to desired effluent (15). Therefore, it is necessary to use catalytic wet air oxidation (CWAO). Treatment of spent caustic wastewater was studied in catalytic reactor with graphene oxide (GO) and ruthenium/ GO catalysts (16). AOP can be defined as the acceleration of an oxidation reaction by generation of hydroxyl radicals. Also, AOP are defined as processes which treat wastewater steams with high COD and toxic contaminants (17). An AOP system involves a catalyst (photo-catalysis) and an oxidant such as H₂O₂, O₃/H₂O₂, fenton, and UV. A novel photocatalytic reactor was studied for the treatment of spent caustic wastewater and TiO₂-clinoptilolite was synthesized for the enhancement of its photocatalytic efficiency (18). COP is used to remove sulfide compounds from the spent caustic wastewater. In the previous study, spent caustic wastewater of an olefin unit was treated in

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Item	Value
Total sulfur	14000-21000 mg/L
COD	15000-30000 mg/L
pН	13.5-13.7
S ²⁻	670 mg/L
Phenols	300 mg/L
Specific gravity	1.1
NaOH	1-3 wt%

two reactor (the system can be represented via a series of bubble column reactor with mixed reactor) by the catalytic chemical oxidation method (19). This waste stream was finally regenerated and recycled to the unit. In the present study, the global reaction kinetics were modelled and optimized in a bubble column reactor via genetic algorithm (GA). Analysis of the kinetic models of catalytic reactions requires more accurate estimation of rate parameters and constants (20-22). The most commonly used methods in determining the optimal parameters model, are non-linear fitting algorithm like artificial neural networks and Marquardt algorithm. But, the structure of non-linear reaction kinetics treatment of spent caustic has more than one relative minimum value. So, it is difficult to achieve the optimal point by using these methods. One way to solve such kind of problems is the use of randomized algorithms. As a new method, GA has been introduced for optimization of non-linear models (23-25). Various applications of GA have been proposed in non-linear models to determine the optimal parameters. The flexibility of genetic algorithm is its major advantage which provides the objective function and constraints in farming (24). It can be seen that researchers have used GA to estimate the amounts of kinetic parameters of complex reactions. Finding a general kinetic model for hydrogenolysis of dibenzothiophene (DBT) based on Langmuir-Hinshelwood type (26), and estimating simultaneously the kinetic as well as energetic parameters of the complex reaction of the saponification of ethyl acetate (27) are examples of the use of genetic algorithm in these cases. In this study, a bench-scale pilot was employed for regenerating and treating spent caustic. The development of global reaction of spent caustic is the main purpose of this work. Another purpose of this study was to investigate the application of GA to choose a precise kinetic model and determine the optimal parameters.

The parameters of this reaction are rate constants and kinetic order of concentrations. The kinetics of the reaction was surveyed as a global reaction in a bubble column reactor. This function objective is prediction error of the model and the experimental concentration. The kinetic steps are shown in Figure 1.

Materials and Methods

Experimental procedure

The waste solution was provided by a petrochemical plant



Figure 1. The flowchart of the overall kinetics.

(Olefin unit) in Iran. The catalytic chemical oxidation was conducted to regenerate and treat spent caustic in a bubble column reactor, with capacity of 500 ml and 40 μ m pore size (Figure 2). IVKAZ catalyst of the JSC company was employed to regenerate spent caustic in a bubble column reactor. The reactor was operated at ambient pressure and 90°C. In previous work, the treatment of spent caustic was accomplished in two reactors by the full factorial design (19). It has been found that the optimal operating temperature and air flow for treatment and regeneration of spent caustic were 85 to 95°C and 10 L/min, respectively, while temperature of 90°C and air flow of 10 L/min were chosen for this kinetic study.

All the demanded materials purchased from Merck Chemicals were used for data analysis procedure. The mercaptide and hydrosulfite sodium were measured using a digital pH meter (Metrohm). These compounds were determined according to ASTM-UOP 209. Thiosulfate was measured using a spectrophotometer. The standard thiosulfate titrant and standard iodine titrant, HCl and indicator were used for the titration of S²⁻ and alkalinity, respectively. The alkalinity, S²⁻ and O₂ of the spent caustic were analyzed by standard methods for the examination of water and wastewater (28). All solutions were prepared using deionized water.



Figure 2. The Schematic of bench scale pilot.

Kinetic modelling

Spent caustic contains a wide variety of compounds, each one has its own specific kinetic characteristics. In the majority of the reactions (1 to 5), thiosulfate was produced as an intermediate compound. Several previous studies have shown that thiosulfate converts to sulfate in an alkaline environment at 76-85°C. The reaction rates 1 to 5 was considered as both elementary and non-elementary reactions. The GA method was used for each of the reactions. Table 2 indicated the reaction rate equations. The reaction rate equations of a, b, c, d, and f belong to caustic, thiosulfate sodium, oxygen, hydrosulfite sodium, mercaptide, and sodium sulfide, respectively.

As air flow enters during the process, the reactor is a semi batch. It is assumed ideal for studying the reaction kinetic (29,30). The bubble column reactor was considered as a batch reactor. For this purpose, the oxygen injection was computed every 5 seconds and the total amount has been added in 20, 25, and 30 minutes.

$$In - Out + r_i * V = d(C_i * V)/dt$$
(6)

Where r_i is the rate of reaction for i material, V is the volume of the batch reactor and t is time.

Oxygen is the only component which enters the reactor and other ones have no entery or exit flow. The solution volume (V) is not constant due to high evaporation especially at high temperatures. To solve this problem, the reaction time is divided to shorter intervals (some seconds) and the output of each interval assumes as the input of the next interval. This technique removes the input and output terms from equation 6 as mentioned below:

1- At the beginning of each interval, it is assumed that the total amount of oxygen which was supposed to enter the reactor gradually, is injected at the beginning.

2- As the reaction time is divided to shorter intervals, the volume variations at these intervals will be small enough to be neglected. It is worth noting that the shorter intervals to be considered, the error of this method will be less. In this way, the equation 7 is rearranged as follows: $r_i = d(C_i)/dt$ (7)

tudies random manner, depending on the range of defined in an changes. This vector consists of seventeen variables and five variables for non-elementary reactions and elementary reactions, respectively. An example of these vectors is $V_i = (k_1 k_2 k_3 k_4 k_5 X_6 X_7 X_8 X_9 X_{10} X_{11} X_{12} X_{13} X_{14} X_{15} X_{15} X_{16} X_{17})$ for non-elementary reaction and $V_i = (k'_1 k'_2 k'_3 k'_4 k'_5)$ for elementary reaction. The range of variables of unique the vectors were between $(10^{-3} - 10^{+1})$ and population size was 500.

> The objective function is defined as the minimum error model results and the experimental results. The error of objective function is shown in equation 8.

> In this study, the following steps were performed to

determine the kinetic parameters. At first, members of the

primary population including constants of reactions and

kinetic order of reaction were generated in a completely

$$\operatorname{Error} = \frac{1}{n} \sum_{i=1}^{n} \frac{\left| C_i^{\exp} - C_i^{cal} \right|}{C_i^{\exp}} * 100$$
(8)

After generating the initial populations via the GA, the value of the objective function of each member is calculated. By using the objective function values based on the roulette wheel method, members are ranked. Obviously, any member with a better objective function will be considered more valuable. GA operators are reproduction, crossover and mutation (31,32).

Reproduction: By this operator, 50% of the members of the current generation based on the roulette wheel are transferred to the new generation as follows:

I) Based on elitism: The best members of the current generation (5%) are transferred to the new generation directly (and not randomly),

II) 45% of the members are randomly assigned to the new generation (30).

Crossover: To produce new generation by the crossover operation, both one- and two-point crossovers are employed. After selecting two vectors by the roulette wheel, a random number is generated to decompose the chromosome (i.e. the vector) into several pseudo parents. Then, these parents are composed together to

	Non-elementary		Elementary
1-	$R_a = -k_1 C_a^{X_6} C_b^{X_7} C_c^{X_8} - k_2 C_a^{X_9} C_c^{X_{10}} C_d^{X_{11}} + 2k_4 C_e^{X_{14}} C_c^{X_{15}} +$	1-	$R_{a} = -k_{1}C_{a}C_{c}C_{b}^{0.5} - k_{2}C_{a}C_{c}^{2}C_{d} + 2k_{4}C_{e}^{2}C_{c}^{0.5} +$
	$2k_5 C_c^{X_{16}} C_f^{X_{17}}$		$2k_5^{\prime}C_cC_f$
2-	$R_b = -0.5k_1 C_a^{X_6} C_b^{X_7} C_c^{X_8} + 0.5k_3 C_d^{X_{12}} C_c^{X_{13}} C_d^{X_{11}} +$	2-	$R_b = -0.5k_1C_aC_b^{0.5}C_c + 0.5k_3C_c^2C_d + k_5C_cC_f$
	$k_5 C_f^{X_{17}} C_c^{X_{16}}$		
3-	$R_{c} = -k_{1} C_{a}^{X_{6}} C_{b}^{X_{7}} C_{c}^{X_{8}} - 2k_{2} C_{a}^{X_{9}} C_{c}^{X_{10}} C_{d}^{X_{11}} - 2k_{3} C_{d}^{X_{12}} C_{c}^{13} -$	3-	$R_{c} = -k_{1}C_{a}C_{b}^{0.5}C_{c} - 2k_{2}C_{a}C_{c}^{2}C_{d} - 0.5k_{3}C_{d}C_{c}^{2} - 0.5$
	$0.5k_4 C_e^{X_{14}} C_c^{X_{15}} - 2k_5 C_c^{X_{16}} C_f^{X_{17}}$		$0.5k_{4}C_{e}^{2}C_{c}^{0.5}-2k_{5}C_{c}C_{f}$
4-	$R_{d} = -k_{2} C_{a}^{X_{9}} C_{c}^{X_{10}} C_{d}^{X_{11}} - k_{3} C_{d}^{X_{12}} C_{c}^{13}$	4-	$R_d = -k_2 C_a C_c^2 C_d - k_2 C_d C_c^2$
5-	$R_e = -2k_4 C_e^{X_{14}} C_c^{X_{15}}$	5-	$R_e = -2k_4 C_e^2 C_c^{0.5}$
6-	$R_f = -2k_5 C_c^{X_{16}} C_f^{X_{17}}$	6-	$R_f = -2k \int_{S} C_c C_f$

Table 2. The equation rate of the reactions (1 to 5)

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produce offspring. In this study, the rate of crossover was considered to be 50%.

It is noteworthy that in this operator, it is possible to choose more members with a better objective function. Mutation: New generation of the present operation is used to all members after producing. The definition of the mutation and its rate plays an important role in the convergence of the algorithm. As a general fact, inappropriate definition of this operator leads to premature convergence of the GA. Therefore, mutation should be defined in such a manner that the GA has sufficient diversity during its evolutionary procedure. The mutation rate is based on the following equation:

 $Mu=Mu_{min}+(Mu_{max}-Mu_{min})$. exp $(-10^*(C_w-C_b)/C_w)$ (9) Where Mu is the mutation rate, calculated in each generation. Mu_{min} and Mu_{max} are the minimum and maximum mutation rates allowed and their values are 10% and 100%, respectively, in the equation 9. C_w is the value of the cost of the worst individual and C_b is the value of the cost of the best one (28). The flowchart of the applied GA method is shown in Figure 3.

Results

The concentrations of components in the bubble column reactor were obtained. As mentioned before, the temperature and flow rate of reaction were constant and only the reaction time changed. The experimental amount of each component is indicated in Figure 4-9. The value of spent caustic is illustrated for elementary and non-elementary models in Figure 4. At an earlier time (20 minutes), the elementary model can better predict the experimental results, also non-elementary model can be seen to be better with time.

The obtained results of thiosulfate sodium are demonstrated in Figure 5. As shown in Figure 5, the thiosulfate amount will be reduced by the increase of reaction time. The oxidation process of thiosulfate to sulfate is a heterogeneous gas-liquid reaction.

Figure 6 compares oxygen amount botained by experimental and GA methods. The real amount of





oxygen has been computed based on the stoichiometric calculations in a 500 mL bubble column reactor. As shown in Figure 6, the results of the non-elementary model is near to the actual values.

According to the amount of NaHS shown in Figure 7, the elementary model is not a desirable model to predict

experimentally-obtained results of NaHS. The hydrosulfite sodium is consumed in two different reactions, so the prediction of this component is difficult by the elementary model.

The oxidation of NaSR is illustrated in Figure 8. The nonelementary model has really low error compared to the













Figure 8. The results of NaSR obtained from experimental data and kinetic model.

elementary one. The results illustrate that non-elementary model is more concordance with experimental results (error of non-elementary model was lower than 10%). The change of amount of Na_2S is indicated in Figure 9. The catalytic oxidation with increase of the reaction time, led to higher conversion of sulfides to thiosulfate, therefore, the amount of sulfide species reduced. The increase of reaction time helped approach the value of

non-elementary model to the experimental amount and resulted in a low error at the end of the reaction time. The kinetic coefficients obtained from elementary and non-elementary models are shown in Table 3. The kinetic coefficient of the reaction 4 had the highest value among

other reactions. This value made the reaction rate much faster than the other reactions. Table 4 indicates the order of concentration in the nonelementary model. These values were obtained by the GA

Discussion

method.

Different chemical oxidation reactions took place in bubble column reactor. The main oxidation reactions of spent caustic are as follows (33):

$$NaOH + \frac{1}{2} Na_2 S_2 O_3 + O_2 Na_2 SO_4 + \frac{1}{2} H_2 O$$
(1)

$$NaSH + 2O_2 + NaOH Na_2SO_4 + H_2O$$
(2)

$$NaSH + 2O_{2} \frac{1}{2} Na_{2}S_{2}O_{3} + \frac{1}{2} H_{2}O$$
(3)
2RSNa + ¹/₂O + H O RSSR + 2NaOH (4)

$$Na_{0}S + O_{0} + \frac{1}{2}H_{0}O NaOH + \frac{1}{2}Na_{0}S_{0},$$
(5)

One of the aims of this study was to investigate the regeneration of spent caustic, therefore, the amount of caustic was expected to increase. The present claim was confirmed by the experimental data. The reaction time variables had a significant effect on the regeneration process of spent caustic. It is mostly proffered to increase reaction time of the whole process. The error of the elementary model was less than the non-elementary one, but the concentration of spent caustic was decreased. It is obvious that the elementary model couldn't properly predict the components of spent caustic in Figure 4.

So, the experiments should be executed in a regime where diffusional resistance in gas and liquid phases could be ignored to find out the true kinetics (34). The kinetics of catalytic oxidation was carried out in the presence of a cobalt phthalocyanine (IVKAZ) catalyst. As shown in Figure 5, the thiosulfate content was significantly reduced, so that the experimental data as well as the results of elementary and non-elementary models' showed the same trend regarding to the studied kinetics. Figure 5 showed





Table 3. Kinetic rate coefficient value for the reactions (1-5)

Kinetic rate Elementary coefficient model		Non-elementary model		
k ₁ & k' ₁	k'_=0.0001	k ₁ =0.0636		
k ₂ & k' ₂	k'2=0.5658	k ₂ =0.0313		
k ₃ & k' ₃	k' ₃ =0.4999	k ₃ =0.0005		
k ₄ & k' ₄	k' ₄ =0.5000	k ₄ =2.0000		
k ₅ & k' ₅	k' ₅ =0.1000	k ₅ =0.3000		

Table 4. The order of concentration in non-elementary model

Variables of non-elementary model							
Item	X ₆	X ₇	X ₈	X ₉	X ₁₀	X ₁₁	
Value	4.994	4.993	4.996	8.029	4.574	5.195	
Item	X ₁₂	X ₁₃	X ₁₄	X ₁₅	X ₁₆	X ₁₇	
Value	10.00	0.0010	3.825	2.416	5.524	3.814	

that the error of non-elementary model was acceptable and could predict well. According to the Figure 5, thiosulfate had the highest concentration at the early times of the reaction. Due to the main oxidation reactions (1), decrease the amount of thiosulfate with time. Specially, by increasing the amount of oxygen and time, the sulfate content also increases (35,36).

Oxygen participates in all five reactions, so it has more terms in the reaction equation. The results of elementary model for oxygen (despite the relatively low error compared to the other compounds), show that these models has not been able to accurately estimate the amount of oxygen in the reactor (Figure 6). The effects of oxygen concentration on the removal of sulfur compounds is directly related with evaluation of kinetics of reaction (16).

NaHS (Figure 7) was accelerated to thiosulfate in the reaction 3 and estimation was rather complex and unsatisfactory, because thiosulfate is an intermediate and unstable component. The kinetics of NaHS is predicted by non-elementary model slightly. The kinetic of the overall reaction tends to the consumption of sulfide. The unstable sulfur compound like hydrosulfide, is converted to thiosulfate and sulfate via increasing oxygen (35).

Mercaptide (NaSR) is considered rather stable under normal ambient temperature and pressure. The chemical oxidation of the mercaptide components to disulfides is actually slow (37). So, an appropriate catalyst is needed to increase the rate of conversion to disulfide. The disulfides are usually insoluble in the alkaline solution and can be separated easily.

According to Figure 9, the increase of reaction time helps approach the value of non-elementary model to the experimental amount and results in a low error at the end of the reaction time. Chemical oxidation of sulfide in alkaline media is usually generated thiosulfate (38).

As shown in Table 3, the orders of the non-elementary model are quite different from the elementary model. The non-elementary model seems to display really better results compared with the elementary one. The overall error of the non-elementary model was less than the elementary one. The comparison of others shows that the order of concentration of the reaction (4) was reasonable. One of the reasons that spent caustic is regenerated in these conditions is due to the value obtained in the reaction (4). According to the results obtained from the third reaction and Table 3-4 (kinetic coefficient and order of concentration), the effect of this reaction can be considered insignificant. The temperature and catalyst concentration are parameters which are prominent in the kinetic model. In this case, temperature and catalyst concentration are fixed. It is very interesting that kinetic coefficient of reactions (2) and (3) in the elementary model was higher than the non-elementary one. These reactions are related to NaHS components. This finding can be attributed to the order of oxygen in reactions (2) and (3). However, the regeneration of spent caustic was the main purpose. The value of oxygen order was lower than the elementary model order.

Conclusion

The conclusions drawn from this study can be resumed as follows:

In order to regenerate and treat the spent caustic, experiments were conducted via a catalytic air oxidation process in a bubble column reactor. A mathematical model was developed for regeneration of spent caustic in the batch system and it was investigated by the experiment. The temperature, catalyst concentration, pH, and air flow are constant and reaction time from operating conditions is unstable.

It was found that heterogeneous cobalt phthalocyanine is a suitable catalyst for chemical oxidation of spent caustic under highly alkaline conditions.

The GA was used to predict the rate coefficient and kinetic order of component concentrations. The elementary and non-elementary models are considered for the whole of reactions. Based on the GA method, the nonelementary model has reasonable results comparison to the elementary one.

Fundamentally, when the materials are produced or consumed in different reactions the estimation is difficult via the elementary model. The errors of the objective function of elementary and non-elementary models was 134.96% and 3.01%, respectively. It was also revealed that the non-elementary model is able to compute true values with the lowest error.

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Ethical issues

Ethical issues (plagiarism, informed consent, misconduct, data fabrication and/or falsification, double publication and/or submission redundancy) have been strictly observed by the authors.

Competing interests

The authors declare that they have no competing interest.

Authors' contributions

All authors contributed equally in all aspects of this research like data collection, analysis, and interpretation.

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