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Research Article

Design of a Synthetic Zinc Oxide Catalyst over Nano-Alumina for Sulfur Removal by Air in a Batch Reactor

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Abstract

Owing to the environmental regulations with respect to sulfur content and continuing challenges of finding a suitable catalyst of such impurity, a driving force for the development of more efficient technologies a deep research on new oxidative catalysts is considered an important issue in fuel quality improvement. Thus, the present study shows a novel percent of nano-catalyst with 18% zinc oxide (ZnO) of active component over nano-alumina that has not been reported in the public domain for sulfur removal from kerosene fuel by air (oxidative desulfurization (ODS) method). Where, such percent of the active component on the nano-alumina helps to add one or two atoms of oxygen to sulfur content in the kerosene. The nano-catalyst (ZnO/nano-alumina-particles composite) is prepared by precipitation of zinc oxide and loaded over nano-alumina in one-step. The activity of the prepared catalyst was tested utilizing ODS process of kerosene fuel by air in a batch reactor. A set of experiments were conducted with a wide range of operating conditions, where the reaction temperature was ranged from 150 to 190 °C, the reaction time from 30 to 50 min and the catalyst weight from 0.4 to 1 g. The experimental results showed that the chemical nature of zinc oxides showed higher conversion (70.52%) at reaction temperature of 190 °C, reaction time of 50 min, and 1 g catalyst weight used in the batch reactor. A kinetic model related to the sulfur removal from kerosene via ODS process in the batch reactor was also investigated in this study for the purpose of estimating the best kinetic parameters of the relevant reactions. The results showed that the prepared catalyst (ZnO over nanoalumina) can be applied confidently to reactor design, operation and control in addition to improve the fuel quality. Following the kinetic model of ODS process, a very well agreement between the experimental and predicted results is obtained. Copyright © 2019 BCREC Group. All rights reserved

Keywords: Nano-Catalyst; Oxidative Desulfurization; Zinc Oxide; Kerosene; Batch Reactor

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1. Introduction

Recently, the main goal of the current decade related to the environmental legislations across the world is to reduce the total sulfur content in feedstock of transportation fuels [1]. These regulations were reported that the maximum sulfur content should not be currently exceeded 0.005 wt%, and less than 0.0015 wt% or nil (freesulfur content) in the coming years [2-4]. The main source of air pollution is attributed to the generation of sulfur oxides by the combustion process leading to form acid rain. The oxidation desulfurization process used for removal all types of sulfur in fuel is expected to solve great environmental problems occurring wherever different petroleum fractions are used [5]. The

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catalytic hydrodesulfurization (HDS), which is commonly known as hydrotreating process requires modified catalyst, higher operating conditions (temperature, pressure, etc.) and toxic emissions may be generated due to the combustion of these fuels producing many gases [6].

A new sulfur removal technology without needing to use hydrogen (H_2) is the oxidation process (ODS) for catalytic decomposition of organosulfur compounds, which are classified as a non-hydrodesulphurization process (HDS) and adopted as ultra-deep desulfurization for many petroleum fractions. Organosulfur compounds in fuel will be converted to sulfone using a percent of oxygen or air directly rather than hydrogen and such process is considered an attractive issue in petroleum refining industries due to the availability of the reacting gas (air) and its low price in addition to safe operating conditions used. Where, the reaction temperature is ranged from 40-100 °C and the pressure is ranged from 1-2 bar and sometimes such process can be applied at room temperature and atmospheric pressure leading to reduce the operating cost compared with HDS process [7-11].

The ODS process is objected to oxidize sulfur compounds corresponding to sulfides or sulfones by adding electrophilic oxygen atom found in air giving a polar form and becoming as non-active compounds that can be removed by extraction or desorption process [12]. Ma et al. [13] studied the sulfur removal from different types of commercial jet fuel by oxidant molecular oxygen at moderate conditions using several sulfur models of jet fuel (BT, 2-MBT, 5-MBT and DBT dissolved in n-decane). They found that the sulfur content can be reduced by adsorption Fe(III) nitrate and Fe(III) bromide. Sundaraman et al. [4] have applied the ODS process of commercial jet fuel with sulfur content of 0.052 wt% and commercial diesel fuel with sulfur content at 0.0041 wt% using air as an oxidant for generating hydro-peroxides with CuO as a catalyst. The Authors have reported that the sulfur compounds in jet and diesel fuel are oxidized to sulfones with MoO₃/SiO₂ then the formed sulfones were adsorbed by beta zeolite to ultra-low sulfur jet and diesel fuel.

Javadli *et al.* [14] utilized the oxidative desulfurization of the model feedstock (dibenzothiophene in bitumen) using air as an oxidant for generation sulfone or sulfoxide in bitumen. N-heptane have employed to prevent the bitumen hardening during autoxidation process based on H_2O used for washing step and they found that the sulfur removal is achieved by 46-47%.

Imtiaz et al. [15] have studied the ODS of sulfur for different feedstock models (thiophene, dibenzothiophene, 4-MDBT) using air/quaternary ammonium salt as oxidant. They found that an increase in temperature process, the rate of removal of sulfur increased rapidly. Nawaf et al. [16] employed the ODS of DBT in light gas oil (LGO) using several initial DBT contents, liquid hourly space velocities and different reactor temperatures in a trickle bed reactor (TBR) based on a prepared catalyst of manganese oxide (MnO_2/γ -Al₂O₃), and high sulfur removal was observed.

Nada et al. [17] have examined the ODS process of kerosene based on ZnO/AC composite prepared by thermal co-precipitation method in a batch system using air as oxidant. The highest conversion of sulfur is about 33% at optimum operating conditions. Thus, this work is aimed to prepare a new nano-catalyst (zinc oxide loaded on alumina nano-particles) for the purpose of testing the possibility of sulfur removal from kerosene utilizing the oxidation desulfurization process (ODS) in a batch system by air as an oxidant. Also, the kinetic model of the oxidative desulfurization process of such reactor is investigated here for further insight of the process.

2. Materials and Method

Kerosene feedstock with sulfur content of 2500 ppm (0.25 wt%) was supplied by the *AL-Dura Refinery Company-Iraq*. The physical properties of the kerosene used in this study are illustrated in Table 1. Nano-alumni particles were supplied by *Skyspring/USA*, zinc acetate C₄H₈O₄Zn.2H₂O with purity of 99% and deionized water were supplied by *Sinopharm Chemical Reagent*. The materials and the specifications of the catalyst used are shown in Table 2.

Various conditions are tested here for oxidtive desulfurization of kerosene as follow: (a) catalyst prepared (nano-catalyst = 18% ZnO/ nano- γ -Al₂O₃); (b). reaction time (30 min, 40 min, and 50 min); (c). reaction temperature (150 °C, 170 °C, and 190 °C); (d). weight of catalyst (0.4 g, 0.7 g, and 1 g).

2.1 Catalyst Preparation (18% ZnO/Nano- γ -Al₂O₃)

Zinc oxide loaded as active component on nano-alumina particles composite is prepared by a thermal co-precipitation method. The nano-alumina particles are dried in an oven for 1.5 h at 250 °C to remove the moisture out of the support pores of alumina. An amount of 8 g by weight of nano-alumina particles is used as the active component. Meanwhile, 4 g of zinc acetate salt was dissolved in 50 mL of deionized water and mixed for 2.5 h to obtain a good dissolving with the salt, where such quantity can

Table 1. Kerosene properties

Property	Value
Density at 15.6 °C	$0.82-0.86 \text{ g/cm}^3$
API	47.6
Flash point	54
Kinematic viscosity at 40 °C (cSt)	2.5 - 5.6
Water content% Vol.	0.05 max
Color	>+30
Sulfur Content	0.25% wt
Smoke Point	28
Ash content% wt	0.01 max
Pour point °C	-9 max Winter -3 max Summer
Distillation Data (°C)	
Initial Boiling point	163
10%	179
20%	184
30%	190
40%	194
50%	198
60%	203
70%	209
80%	215
90%	224
End Point	246

give 18 wt% of ZnO on Nano- γ -Al₂O₃ loaded. Then, zinc acetate solution prepared is slowly added on the Nano-alumina particles under continuous mixing and without heating. The prepared mixture was heated for 2 h in the oven at 125 °C for 5 °C/min with laminar air flow. After that, the mixture is firstly heated at 250 °C for 2 h and the temperature is then increased to 400 °C for 1 h to get a well drying of the mixture. Finally, the mixture is heated up to 560 °C for 2 h where the metal oxides are loaded on the supported nano-alumina followed gradually by a cooling process. The calcination procedure (as shown in Figure 1) has been conducted in Hadithah Diesel Power Plant-Al-Anbar-Iraq.

2.2 Experimental Procedure for ODS Process

The oxidative desulfurization experiments were carried out into batch reactor (250 mL) using kerosene as a feedstock and air as oxidant for oxidative desulfurization reactions. In all experiments, 100 mL of kerosene has charged for each run. The air flow rate in ODS reactions was 0-3 L/h with keeping constant pressure at 0.5 bar. The prepared catalyst was charged to the batch reactor containing the liquid feedstock (kerosene), where the ODS reactions are carried out. Several experiments were carried out under moderate operating conditions (reaction temperatures (150 °C, 170 °C, and 190 °C), reaction time (30 min, 40 min, and 50 min) and catalyst weight (0.4 g, 0.7 g, and 1)g) with keeping constant stirring rate used for all experiments at 500 rpm for the purpose of obtaining a good mixing and avoiding segregation of both fluid and catalyst [18]. The evaporated gases during the oxidation reactions are

Table 2. Materials and specifications of the catalyst used

	Materials
Chemical Properties of Zinc Ac	<i>etate</i>
Chemical Structure	$\mathrm{C_{4}H_{8}O_{4}Zn.2H_{2}O}$
Molecular Weight	219.51
Purity	≥99.0%
PH (50 g/L, 25 °C)	6.0-7.0
Typical Impurities	Cl<0.001% , SO2 ≤0.002% , Fe<0.003%, As<0.00002% , Pb<0.002%
Specifications of Alumina Nan	o Particles
Appearance	White
Empirical formula	$\gamma ext{-}\mathrm{Al}_2\mathrm{O}_3$
Purity	$\geq 99.9\%$ (trace metal basis)
APS	20 nm
Typical Impurities	Si = 20 ppm, Na = 10 ppm, K = 10 ppm, Ca = 20 ppm Fe = 20 ppm



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condensed through the condensation supplied to the process. Product sulfur concentrations were analyzed via X-ray diffraction instrument based on ASTM D7039 method. The product sulfur content of kerosene was determined in *AL-Dura Power Station Laboratories/Ministry* of *Electricity-Iraq*. Process flow diagram of the batch reactor system is illustrated in the Figure 2.

2.3 Characterization of Nano-Catalyst Prepared

The amount of the active component (metal oxide) used for preparing the nano-catalyst from zinc oxide via composite zinc acetate was tested by BET surface area, pore volume and pore size. Determination of the prepared nano-catalyst particles has been carried out at *The Petroleum Research and Development Center/Ministry of Oil-Iraq*. The data obtained with respect to surface area of the nano-catalyst is shown in the Table 3 and illustrated in Figure 3. Operation conditions of the process analysis are as follow: analysis adsorptive of N₂; analysis bath temperature of 77.441 K; warm free space of 28.8645 cm³; and equilibration interval of 10 s.

Langmuir surface area as an isotherm for gases adsorbed on the solid particles has also been estimated, which is an empirical isotherm derived from a proposed kinetic mechanism. It is based on the surface of the adsorbent as uniform, where all the adsorption sites are equals and the adsorbed molecules should not be intersected with each other. At maximum adsorp-



Figure 3. Isotherm data of the prepared nanocatalyst

tion, a monolayer is formed and the molecules of the adsorbate are not deposited on other sites, but the molecules will be adsorbed on the free surface of the adsorbent only. The obtained isotherm tabular data of the prepared nano-catalyst is reported in the Table 4.

2.4 Mathematical Modelling of ODS Reactions

2.4.1 Model based on kinetics of oxidation desulfurization

Kinetics model related to the oxidative desulfurization process is studied here based on oxygen as oxidant and $ZnO/nano-\gamma-Al_2O_3$ catalyst including the pseudo first and second order kinetics, external diffusion model and intra particle diffusion model, which are expressed within the kinetic rate equation to find the best kinetic model through the comparison among correlation coefficients.

Mole balance in the batch reactor for desulfurization:

$$Mole in = Mole out = 0 (in the batch reactor)$$
(2)

Substitution of Eq. (2) into Eq. (1) results in:

$$Accumulation = -(Dissapparance by reaction)$$
(3)

$$Accumulation = (-r_{sulfur})(V_{Batch})$$
(4)

Dissapparance by reaction =
$$\frac{dN_{sulfur}}{dt}$$
 (5)

From Equations (3), (4), and (5), Eq. (6) becomes:

$$-\frac{dN_{sulfur}}{dt} = \left(-r_{sulfur}\right) V_{Batch}$$
(6)

At constant density,

$$-\frac{d\left(\frac{N_{sulfur}}{V_{Batch}}\right)}{dt} = \left(-r_{sulfur}\right)$$
(7)

$$-\frac{d C_{sulfur}}{dt} = \left(-r_{sulfur}\right) \tag{8}$$

The chemical reaction rate equation is stated as follows:

For Pseudo first order =
$$(-r_A) = KC_{sulfur}$$
 (9)

$$-\int_{C_{sulfur(0)}}^{C_{sulfur}} \frac{dC_{sulfur}}{C_{sulfur}} = \int_{0}^{t} K dt$$
(10)

$$\ln \frac{C_{sulfur(0)}}{C_{sulfur(f)}} = Kt \tag{11}$$

For Pseudo second order =
$$(-r_A) = KC_{sulfur}^2$$
 (12)

Thus,

$$-\int_{C_{sulfur(0)}}^{C_{sulfur(f)}} \frac{dC_{sulfur}}{C_{sulfur}^2} = \int_0^t K dt$$
(13)

$$\frac{1}{C_{sulfur(f)}} = \frac{1}{C_{sulfur(0)}} + Kt$$
(14)

2.4.2 Pore diffusion in the prepared catalyst $(18\% ZnO/nano-\gamma-Al_2O_3)$

Studying the effect of air and sulfur mass transfer in kerosene for oxidative desulfurization process inside pore catalyst prepared (ZnO/nano- γ -Al₂O₃) is calculated through the Thiele modulus (M_T) as presented in the following equation [19]:

$$M_T = L \left(\frac{K(n+1)C_{sulfur}^{(n-1)}}{2D_{Eff}} \right)^{\left(\frac{1}{2}\right)}$$
(15)

For second order kinetic reaction:

$$-r_{sulfur} = KC_{sulfur}^2 \tag{16}$$

Hence:

$$M_T = L \left(\frac{3KC_{sulfur}}{2D_{Eff}} \right)^{\left(\frac{1}{2}\right)}$$
(17)

Effective diffusivity (D_{Eff}) can be determined as follows:

$$D_{Eff} = \frac{D_{AB} \in_P}{\tau_P} \tag{18}$$

In this work, the ZnO/nano- γ -Al₂O₃ catalyst have been assumed to be as sphere particles. So, the porosity ϵ_P can be estimated depending on the experiments of the total pore volume, while, the tortuosity was found to be 1.978 [20,21]. The molecular diffusion coefficient (D_{AB}) of the air in kerosene is estimated to be 0.0495 cm²/s at 67 °C and equals to 0.5789 cm²/s at 25 °C [22]. The diffusivity function with temperature can be stated as:

$$\frac{D_{AB}}{D_{AB0}} = \left(\frac{T, K}{T_0, K}\right)^{1.5} \tag{19}$$

3. Results and Discussion

3.1 Catalyst Characterizations

3.1.1 BET surface area (ASTM D5604)

The surface area of the prepared nanocatalyst 18% ZnO/nano-y-Al₂O₃ composite was measured by BET method. The values of the surfaces area, pore size and pore volume are listed in Table 5. The surface area of nano catalyst decreases after loading metal oxide on the catalyst supported approximately by 18% (weight percent) from zinc oxide, but such surface area of 18% ZnO/nano-y-Al₂O₃ is still high. It can be observed based on the results obtained before and after preparation process that the surface area can close to that of the original source. Occasionally, the surface area and pore volume increases or be the same as the original source in case of loading ratio less or equals to 10 wt% due to a good dispersion of the metals oxides on the supported area [18].

Among the physical and chemical properties of the prepared nano catalyst, the specific surface area of 18% ZnO/nano-y-Al₂O₃ is an important factor affecting the catalytic activity. As the specific surface area of the catalyst preparation increases, the number of effective active site increased leading to increase the oxidation process of the reaction rate. The BET specific surface areas of the prepared catalyst and the Langmuir surface areas are shown in the Tables (3) and (4), respectively. The specific surface area is significantly dependent on the calcination temperature used during the catalyst preparation. The optimal specific surface area is found by increasing the calcination temperature up to 580 °C, but an increase in such calcination temperature above 600 °C a decrease in the surface area is observed. Such behavior can be attributed to decrease in the pore volume by increasing the calcination temperature owing to the melting of metals oxide upon the surface of the catalyst at high calcination temperature [24].

Table 5. Specification	of nano-supported and	l nano-catalyst prepared
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Specification	Nano- γ -Al ₂ O ₃	18% ZnO/nano- γ -Al $_2O_3$	
Surface area	180-200 m²/g	$159.2676 \text{ m}^2/\text{g}$	
Langmuir surface area	$245.5 \text{ m}^2/\text{g}$	223.6012 m²/g	
Pore volume	-	$0.845870 \ { m cm^3/g}$	
Pore size	-	212.4398\AA	
Catalyst nano particle	-	20 nm	



Figure 4. Chemical reactions occur in pore size distribution



3.1.2 Pore size distribution (ASTM D4641) and pore volume (ASTM ISO 9277)

Catalyst pore size distribution (Å) structure estimates how easily the reactants can access the interior surface of the catalyst that has been determined by N_2 desorption measurement at *The Petroleum Research and Develop*-



Figure 7. Effect of reaction time on sulfur removal using the prepared nano-catalyst (a) at 1 g and temperatures (150-190 °C), (b) at 0.7 g and temperature (150-190 °C), (c) at 0.4 g and temperature (150-190 °C)

ment Center/Ministry of oil-Iraq by using the testing device. The pore structure parameters include pore sizes and their distribution, pore volume and surface area. Various classifications of pore sizes can be found in literature, but the International Union of Pure and Applied Chemistry (IUPAC) has classified the pores for different sizes by the following manner: micropores > $(20 \ ^{\circ}A)$; mesopores (20-500)°A); and macropores > (500 °A) in diameter. In general, hydrotreating catalysts are prepared by precipitation methods and the catalyst pore structure is mainly determined by the catalyst support. The pore size distribution and the total pore volume of the prepared catalyst 18% ZnO/nano-y-Al₂O₃ has determined to be 235.7984 °A and 0.845870 cm³/g, respectively. The catalyst pore structure is also a significant issue for oxidation process of kerosene liquid due to diffusional limitations. Figure 4 shows the chemical reactions in pore size distribution.

3.1.3. Metal dispersion (ASTM D3908)

Figure 5 illustrates the elemental image distribution of the active component on the supported site for 18% ZnO/nano- γ -Al₂O₃. From the results presented in this Figure, there is a good surface distribution of metal oxides (zinc) over alumina Nano particles surface giving a clear indication that the distribution of zinc oxide showed very well correspondence for oxidation desulfurization reactions. These results ensure that the precipitation is an efficient method to prepare such type of the catalyst with a good active metal distribution.

3.2. Effect of Operating Conditions on Sulfur Removal

In this work, the effect of the operating conditions on the oxidation process in the batch reactor was studied. All sulfur compounds were converted to sulfone then followed by sulfur conversion process as shown in Figure 6. The effect of different reaction time, reaction temperature, and several catalyst weights on sulfur removal are described.

3.2.1. Reaction time

An increase in the mixing time (reaction time) from 30-50 min of the oxidation process, the removal of sulfur content was increased as shown in in Figure 7a, b, and c. The maximum conversion based on the ODS reactions was found to be at 50 min reaction time and 190 °C (reaction temperature) with 70.52% of the sulfur removed using 1 g of the prepared catalyst

(18% ZnO/Nano- γ -Al₂O₃) as shown in the Figure 7a. Increasing the reaction time leading to increase the contact time between the sulfur compounds found in the feedstock and the oxygen dissolved as well as the surface area of the prepared Nano-catalyst giving high residence time for the purpose of converting sulfur to sulfone or sulfoxide [25]. However, at long time, the observed adsorption capacities in such oxidation reactions of the trend sulfur is probably



Figure 8. Effect of reaction temperature on sulfur removal using the prepared nano-catalyst (a) at 1 g and reaction times (30-50 min), (b) at 0.7 g and reaction times (30-50 min), (c) at 0.4 g and reaction times (30-50 min)

reflected upon the accessible adsorption surface due to the presence of micro-meso porosity as revealed in the textural properties of the prepared catalyst (18% ZnO/nano- γ -Al₂O₃).

3.2.2. Reaction temperature

Increasing the reaction temperature through oxidation desulfurization process will contribute to an increase the air dissolved in



Figure 9. Effect catalyst weight on sulfur removal using the prepared nano-catalyst (a) at 50 min and temperatures (150-190 °C), (b) at 40 min and temperatures (150-190 °C), (c) at 30 min and temperatures (150-190) °C)

the feedstock (kerosene) causing an increase in the chemical reaction rate within pore site of the prepared catalyst. Also, at moderate reaction temperature, an important physical properties such as diffusivity (D_{AB}) and Henry's constant (H) was increased with decreasing the kerosene viscosity and surface tension at the same time [26,27]. The impact of the reaction temperature on sulfur removal is illustrated in Figure 8a, b, and c at different reaction times and catalyst weight of 1 g, 0.7 g, and 0.4 g, respectively. Such behavior with respect to sulfur removal can be explained by the molecule of sulfur compound adsorbed upon the surface of the prepared catalyst (18% ZnO/nano- γ -Al₂O₃)

Table 6. Kinetics models results

Order	K (s ⁻¹ .Con ¹⁻ⁿ)	Weight of catalyst, g
	Temperature = 150 °C	
n=1	0.0246	1
n=2	1.646×10^{-5}	1
	Temperature = 170 °C	
n=1	0.0288	1
n=2	2.1012×10^{-5}	1
	Temperature = 190 °C	
n=1	0.0312	1
n=2	2.448×10^{-5}	1
	Temperature = 150 °C	
n=1	0.0143	0.7
n=2	7.974×10^{-6}	0.7
	Temperature = 170 °C	
n=1	0.0187	0.7
n=2	1.1079×10^{-5}	0.7
	Temperature = 190 °C	
n=1	0.0229	0.7
n=2	1.48808×10^{-5}	0.7
	Temperature = 150 °C	
n=1	0.0092	0.4
n=2	4.444×10^{-6}	0.4
	Temperature = 170 °C	
n=1	0.01397	0.4
n=2	7.4483×10^{-6}	0.4
	Temperature = 190 °C	
n=1	0.01612	0.4
n=2	9.033×10^{-6}	0.4

acting as adsorbent for sulfur molecules. Where, zinc oxide has showed high adsorption efficiency for different sulfur compounds owing to its high surface area based on the metal oxides.

3.2.3 Catalyst weight

The influence of the catalyst weight on sulfur removal during oxidative desulfurization reactions of kerosene fuel utilizing the prepared nano-catalyst at different reaction times and temperatures is presented in Figure 9a, b and c. Increasing the weight of the prepared nano-catalyst from 0.4 to 1 g leads to reduce the sulfur content in the kerosene products leading to increase the conversion process from 53.8% to 70.52% at 50 min and 190 °C. These results were obtained due to the high surface area available whenever the nano-catalyst weight was increased giving high sulfur removal. Where, such surface area of the prepared nano-catalyst plays a significant role in providing more surface area of the reaction to occur. As well as, such behavior can increase the vacuum size of the pore sites leading to increase the chemical reaction rate and as a result, the sulfur molecules interaction with oxygen will increase.

3.3. Kinetics Model for Kerosene ODS Reactions

The reaction time can effect on the sulfur content in kerosene liquid fuel at different temperature. The reaction rate constants based on pseudo first and second order kinetics model have been estimated at different reaction times and temperatures. Table 6 lists the comparison between such two kinetic models (first and second order) used in the oxidation desulfurization process with several weights of the nanocatalyst (0.4 g, 0.7 g, and 1 g) for the purpose of giving further insight of the process. Based on the results presented in Table 6, it has been found that the ODS reactions of kerosene fuel oil employing the prepared nano-catalyst is followed by the pseudo second order kinetic giving a good fit than pseudo first order kinetics.

3.3.1. Activation energy of ODS process

Depending on Arrhenius correlation, a plot of $(\ln K)$ against (1/T) will give a straight line with slope equals to (-E/R) and the activation energy (E) is then linearly evaluated as shown in Figure 10a, b, and c at 0.4 g, 0.7 g, and 1 g, respectively. The generated activation energies of the relevant reactions are introduced in Ta-

ble 7 and such values are close to that obtained by Nawaf *et al.* [16,20]. It is clearly observed based on the results reported in this Table, the low amount of the prepared nano-catalyst gives higher activation energy for such reactions studied here indicating that the new nanocatalyst prepared in this work can be utilized confidently to reactor design, operation and control.





3.3.2. Thiele modulus

Tables 8 shows the results of Thiele modulus obtained here with respect to ODS reactions of kerosene using the prepared nanocatalyst at different reaction temperature and different catalyst weight. From the results of reported in this Table, it has been shown that all values of Thiele modulus (MT) for all products are less than 0.4 giving a clear indication that the effect of internal mass transfer on the overall oxidation rate can be neglected. Such results could be attributed to the small size of the particle making the pore diffusion resistance very small.

3.4. Simulation Results

For further insight of the ODS process, simulation results employing the kinetic model proposed in this study based on the nanocatalyst prepared are required. The comparison results between experimental and model predicted data for ODS of kerosene utilizing the prepared nano-catalyst over a wide range of the operating conditions investigated in this work, are shown in the Figure 11a, b, c, and d. As clearly seen from these Figure, process simulation results have found to agree well with the experiments conducted in a wide range of the studied operating conditions (different catalyst weights, several reaction times and temperatures). Such agreement indicated that the model can be applied confidently for other operations. Thus, the nil content of sulfur compound based on the optimal design and operation of such process is a worthy subject for future study.

4. Conclusions

Sulfur removal by air of kerosene fuel in a batch reactor based on the new nano-catalyst (18% ZnO/nano- γ -Al₂O₃) prepared in this study can successfully be achieved. Oxidative desulphurization process (ODS) appears to be technically and economically available for processing ultra-low sulfur fuel. It can be considered as an alternative process in comparison with

Table 7. Activation energies of the preparednano-catalysts

Weight of Catalyst, g	Activation energy, kJ/mole
0.4	28.9402
0.7	25.3095
1	16.1449



Figure 11. Comparison between experimental and predicted results at different reaction temperatures with (a) at 1 g and 30 min, (b) at 1 g and 40 min, (c) at 0.7 g and 40 min, (d) at 0.4 g and 50 min

Table 8. Effect of operation conditions on Thiele modulus

Temperature, K	$K(\mathrm{sec})^{\cdot 1}\mathrm{Con}^{1\cdot \mathrm{n}}$	$D_{AB}~({ m cm^{2}/ m sec})$	$D_{\it Eff}~({ m cm^2/sec})$	M_T
	Wei	ght of Catalyst, 1 g	Ţ	
423.15	2.7433×10^{-7}	0.05789	0.0166	4.2856×10^{-10}
443.15	3.5020×10^{-7}	0.10490	0.0301	9.9871×10^{-10}
463.15	4.0800×10^{-7}	0.11208	0.0322	9.9520×10^{-10}
	Weig	ght of Catalyst, 0.7	g	
423.15	1.3290×10^{-7}	0.05789	0.0166	1.0817×10^{-9}
443.15	1.8465×10^{-7}	0.10490	0.0301	8.7667×10-10
463.15	2.4801×10^{-7}	0.11208	0.0322	9.0604×10-10
	Weig	ght of Catalyst, 0.4	g	
423.15	7.4066×10^{-8}	0.05789	0.0166	6.1609×10-10
443.15	1.2413×10^{-7}	0.10490	0.0301	7.8764×10^{-10}
463.15	1.5055×10^{-7}	0.11208	0.0322	8.0329×10^{-10}

the conventional hydrodesulfurization process (HDS). Notably, ODS does not require hydrogen (only air) as presented by HDS process in addition to its capital cost and operating cost that would significantly be lower than HDS process (further studies will be investigated in future).

The oxidative desulfurization process using the new catalyst showed that the chemical reactions can be carried out under moderate reaction temperature with very short reaction time. Also, the air can be used as an efficient oxidant based on the new nano-catalyst prepared in this study compared with other costly oxidants, mainly H_2O_2 , ozone and TBHP (reported in the literatures for ODS processes). Second order kinetics model have been obtained for ODS reaction and it can be used to represent the system. The effect of internal mass transfer can be neglected.

Nomenclatures

- r_{sulfur} : Sulfur rate of reaction
- V_{Batch} : Reactor volume, cm³
- C_{sulfur} : Sulfur concentration, ppm
- t : Reaction time, min
- K : Kinetic rate constant, (time)⁻¹(conc.)¹⁻ⁿ
- M_T : Thiele modulus
- *n* : Order of reaction
- $D_{E/f}$: Effective Diffusivity Coefficient, cm³/cm solid. sec
- D_{AB} : Molecular Diffusion Coefficient, cm³/sec
- E : Activation energy, kJ/mol
- L : Characteristic length of the catalyst, cm

Greek Letters

- τ_P : Tortuosity of the nano-catalyst pores
- ϵ_P : Porosity of the nano-catalyst particles
- γ : Gamma

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