PREPARATION OF V$_2$O$_5$ @MESOPOROUS SILICA SPHERE FOR NOVEL DUAL OXIDATIVE-ADSORPTIVE DESULFURIZATION

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Key Words: Oxidative-Adsorptive desulfurization; Mesoporous silica spheres; micelle-template; Vandia.

ABSTRACT:

V$_2$O$_5$ nanoparticles were supported on mesoporous silica spheres (V$_2$O$_5$/MSS) by impregnation technique. Where mesoporous silica sphere (MSS) was synthesized via micelle-template method. The structure and morphology were tested using XRD, FTIR, BET surface area, TEM, and DLS techniques. The results demonstrated that the samples were amorphous with a mesoporous structure, special interior spaces, and high specific surface area. The prepared sample was tested for duel oxidative/adsorptive performance of model diesel fuel. The desulfurization process was tested in the presence of H$_2$O$_2$ as an oxidizing agent at different reaction conditions (temperature, contact time, and initial concentration) using a batch system by either stirring or ultrasonic. The data indicate that V$_2$O$_5$/MSS shows good performance toward duel oxidative/adsorptive desulfurization with 100% removal efficiency under sonication within 1.5h but in stirring within 2h at 60°C using 600 ppm initial concentration.

1. INTRODUCTION

Mesoporous materials represent a range of porous materials with 2–50 nm diameters. They are commonly used in numerous fields, catalysis, biomedicine, and sensor. Because of their incredible properties and characteristics, including ultra-high surface areas, controllable pore sizes, and large volumes of pores (1). Various techniques such as Sol-Gel processing(2), Template assisted techniques(3), Microwave-assisted techniques(4), Chemical etching techniques can be used to synthesize mesoporous materials(5). Mesoporous silica are considered a good selection because of their amorphous character and the easy process of sphere formation (6–11).

Moreover, it is known that the earth's crust is rich in the element of vanadium (V), and the vanadium oxides have different forms of crystalline and

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multi-oxidation state (II – V). Vanadium oxides exhibit significant interactions with molecular or ionic species, higher catalytic efficiency and/or effective correlations between electron and electron (12). From the practical and economical points of view, V$_2$O$_5$ has a greater potential to be applied as catalyst active phase for oxidative desulfurization (ODS) process (13). Due to their large pore size with controlled distribution, which may be beneficial in allowing accessibility of large molecular size sulfones to the surface active sites, much attention was paid to development of mesoporous oxide-based materials (14). On the other hand, the worldwide attention has been drawn to the sulfur content of diesel fuel as after combustion, the sulfur compounds in the fuel are converted to sulfur oxides (SOx), which cause acid rain and air pollution, as well as catalyst toxicity. Governments worldwide have noticed these concerns and have imposed new strict environmental regulations on fuel sulfur levels to minimize SOx emissions (15,16). Several approaches have been investigated to remove those heterocyclic sulfur compounds (14,17–22). The traditional fuel sulfur removal method is called hydrodesulfurization (HDS), a well-known catalytic process in the refining industry. However, HDS is not efficient in removing heterocyclic sulfur compounds such as dibenzothiophene (DBT) and its derivatives. It is also demands strict operating conditions such as high temperatures, high pressure, and high consumption of hydrogen (23). Oxidative desulfurization (ODS) is a great alternative approach to hydrodesulfurization. This eliminates the use of costly hydrogen and can be performed under mild conditions with short time of reaction, greater selectivity and efficiency (24,25).

In the ODS process, these refractory aromatic organosulfur compounds (such as DBT and its derivatives) are quickly oxidized into their corresponding sulfoxides and highly polar sulfones, which can be removed from the fuel by adsorption or extraction (26). Since heterocyclic sulfur compounds are relatively large molecules, the pore diameter and surface area of the catalyst have to be taken as part of the catalyst design (13). Adsorption desulfurization technology is considered promising alternative method for deep desulfurization, owing to the potential benefits of low capital, the mild operating conditions of temperature and pressure, low environmental impact and greater reaction specificities, and availability of adsorbents with high adsorption capacity (24). The combination of oxidation and adsorption is better than direct adsorption to achieve a better desulfurization performance (27). However, there are few studies reported that consider insitu oxidative-adsorptive desulfurization as a more efficient than either oxidation or adsorption (28–30). In this case, the desulfurization efficiency is generally based on the catalyst/adsorbent system characteristics (31,32).

According to the previous statement, our aim to prepare ordered mesoporous materials with narrow pore size distribution for removal of sulfur compounds from model diesel fuel. Firstly, the mesoporous silica spheres (MSS) will synthesized via micelle-template method then will impregnated with
V$_2$O$_5$ using a wet impregnation technique to deposit V$_2$O$_5$ on mesoporous silica spheres. The prepared catalyst will be characterized by XRD, FTIR, BET, TEM and DLS techniques. The efficiency of the prepared sample as a catalyst-adsorbent system will be examined for oxidative-adsorptive desulfurization of dibenzothiophene (DBT), as a model diesel oil of sulfur-containing compound, in presence of H$_2$O$_2$ as oxidant. To achieve the best selectivity and activity, different factors were studied such as contact time, reaction temperature, and initial concentration of DBT. Additionally, the effect of ultra-sonication versus stirring will be also studied.

2. EXPERIMENTAL:

2.1. Materials:
Tetraethoxysilane (TEOS, 99.9%, Sigma Aldrich), Aqueous ammonia solution (25–28 wt.%, Merck), Ammonium metavanadate (NH$_4$VO$_3$, 99.8%, Sigma Aldrich), Cetyltrimethylammonium bromide (CTAB, Sigma Aldrich), Ethanol (EtOH (99 %, Merck Chemicals), Oxalic acid (99.5%, Merck), Dibenzo-thiophene (99 %, Merck), Hydrogen peroxide (aqueous solution, 30%), Hexane and Deionized water.

2.1.1. Synthesis of mesoporous silica spheres (MSS):
Mesoporous silica sphere (MSS) was prepared via the hydrolysis and condensation of TEOS in an aqueous basic-ethanol medium. Typically, 0.16g of CTAB was dissolved in 100 mL of mixed aqueous solution of ethanol and water with continuous stirring. Then 1.0 mL of TEOS and 1.0 mL of ammonia were added sequentially. The resulting mixture was further stirred for 6 h at 30 °C until it gives a white suspension. After reaction, the suspension was centrifuged and washed with water and ethanol by two cycles to remove any residual organics and ammonia. The final particles were collected after drying over-night at 80 °C then calcined at 500 °C for 5 h in static air (heating rate 2 °C min$^{-1}$).

2.1.1. Synthesis of V$_2$O$_5$/mesoporous silica spheres:
To prepare V$_2$O$_5$/MSS, the dried sample of mesoporous silica sphere (MSS) was impregnated with an aqueous solution of NH$_4$VO$_3$ that dissolved in oxalic acid solution and stirred at ambient temperature for 10 h. The obtained sample was first dried for 24 h at 120 °C and then calcined for 5 h at 550 °C in static air (heating rate 2 °C min$^{-1}$).

2.3. Characterization of the prepared sample:
X-ray diffractometer (XRD) (XPert PRO, Analytical, Netherlands) tested the crystal structure of the prepared samples using CuKp radiation at 40 kV, 2θ = 0.02 ° phase size and 0.4 sec step time of scanning. N$_2$ adsorption-desorption isotherms (BET) estimated at liquid N$_2$ (-196 ° C) by using a NOVA2000 gas sorption analyzer (Quanta Chrome Corporation) device was examined the texture properties. Fourier-transform infrared
spectroscopy (FT-IR) spectrum was reported between 500 and 4000 cm\(^{-1}\) with Perkin Elmer FTIR spectrometer (model one FT-IR spectrometer, USA). High-resolution transmission electron microscopy (HRTEM) was conducted by using a JEOL 2100F TEM at 200 kV of the accelerating voltage. To prepare TEM sample, A dilute colloidal sample mixture was ultra-sonicated in ethanol for 30 minutes and a drop of this solution was placed on a carbon-coated Cu TEM grid. The average particle size distribution of prepared samples was obtained by Dynamic light scattering (DLS). Analysis was carried out using measurements at 25 °C using a Zetasizer NanoZS (Malvern Instruments Ltd., UK).

2.4. Activity test:
The dual oxidation-adsorption efficiency of the V\(_2\)O\(_5\)/MSS was examined for DBT removal from liquid fuels. In a standard experiment, a condenser, a thermometer, a magnetic bar and a hot plate were fitted with a 25 ml three-necked flask. In this flask, 0.02 g V\(_2\)O\(_5\) / MSS and 0.4 ml H\(_2\)O\(_2\) are applied to a definite concentration of 10 ml of DBT solution (in hexane as a solvent).

Different parameters that influence the efficiency of DBT removal have been investigated, such as the absence of adsorbent or an oxidizing agent, contact time (up to 300 minutes), reaction temperature (30-80°C) and initial DBT concentration (200-1000 mg L\(^{-1}\)). Also, the effect of stirring versus ultra-sonication on the removal efficiency was investigated. GC-FPD has been used to detect the concentration of residual DBT. The efficiency of DBT removal (%E) and the adsorption capacity q\(_e\) (mg g\(^{-1}\)) were determined as follows:

\[
\% E = \frac{C_o - C_e}{C_o} \times 100
\]

(1)

\[
q_e = \frac{(C_o - C_e) \times V}{m}
\]

(2)

Where C\(_o\) is the initial DBT concentration of its solution and C\(_e\) are remaining concentration of DBT after the adsorption (mg L\(^{-1}\)). V is the DBT volume in the solution, and m is the mass of the V\(_2\)O\(_5\)/MSS (g).

3. RESULTS AND DISCUSSION

3-1 Characterization of the Prepared Samples:
XRD pattern of the prepared V\(_2\)O\(_5\)/MSS after thermal treatment at 550°C in air is shown in Fig. 1. showed a broad diffraction peak in the region of 2\(\Theta\) 20-38° indicating that the silica samples are amorphous structure matrix (1,33). Despite the amount of V\(_2\)O\(_5\), some low-intensity diffraction peaks related to the V\(_2\)O\(_5\) supported on the amorphous silica pattern are observed (34).
Fig. 1. XRD pattern of the prepared V$_2$O$_5$/MSS

Fig. 2. showed the FT-IR spectrum of the obtained samples at different preparation steps, dried core-shell solid silica spheres (SSCS), Mesoporous silica sphere (MSS) and V$_2$O$_5$/MSS. A distinctive Si-OH stretching vibration was displayed at about 3423 cm$^{-1}$. The structural differences between obtained (SSCS) and (MSS) were observed through the transverse optical mode of Si—O—Si asymmetric stretching vibration band exhibits a distinct red shift from 1101 to 1086 cm$^{-1}$ of (MSS). The redshift of the Si—O—Si band suggests a more open structure in (MSS), which demonstrates the higher degree of condensation of silicate species (35). Also, The peaks at 2925 cm$^{-1}$ and 2854 cm$^{-1}$ were attributed to asymmetrical stretching vibration of -CH$_3$ and -CH$_2$-, which caused by organic matter inside the dried core-shell silica sample (SSCS) (36). The strong absorption peak at 1082 cm$^{-1}$ was belonged to Si-O-Si asymmetric stretching(35). The peaks at 801 cm$^{-1}$ and 464 cm$^{-1}$ were assigned to symmetric stretching vibration of Si-O, and the peak at 964 cm$^{-1}$ was due to bending vibration absorption of Si-OH. Peak around 1635 cm$^{-1}$ was the bending vibration peak of H-O-H in water (36). The structural differences between obtained (MSS) and V$_2$O$_5$/MSS were observed where V$_2$O$_5$/MSS spectrum shows that the transverse. The peaks at 939, 804 and 469 cm$^{-1}$ belong to stretching vibration of terminal oxygen bonds of Si-O-V vibrations (37), the vibration of doubly coordinated oxygen (bridge oxygen) bonds, and the asymmetric and symmetric stretching vibration of triply coordinated oxygen (chain oxygen) bonds, respectively (38).
**Fig. 2.** FT-IR spectrum of the obtained samples at different preparation steps, Dried core shell solid silica spheres (SSCS), Mesoporous silica sphere (MSS), V$_2$O$_5$/MSS

HR-TEM images show the shape of the as-prepared samples. Typical show that as-prepared of dried core-shell solid silica spheres (SSCS) and Mesoporous Silica Sphere (MSS) after calcination are of spherical morphology (Fig. 3a, b), and SSCS exhibit a smooth surface while MSS show rough pore-like channels indicating the presence of micelle aggregates (35). The average radius of the spheres was not affected by calcination processes (PSD ~ 100 nm), as confirmed by DLS data (Fig. 3c).

**Fig . 3.** HRTEM images of obtained samples at different preparation steps, (a) TEM image of as-prepared dried core-shell solid silica spheres (SSCS). (b) TEM of Mesoporous Silica Sphere (MSS) after calcination (c) Particle size distribution of Silica Sphere at different preparation steps and (V$_2$O$_5$/MSS) after calcination
Textural properties of the prepared V$_2$O$_5$/MSS samples are illustrated in (Fig. 4) Nitrogen adsorption-desorption shows type IV of isotherms, which are typical of mesoporous materials. The hysteresis loop of these isotherms can be classified as type H$_3$, characterized by a triangular shape (39). Since the absorption and desorption branches appear at a relatively low pressure of about 0.4, materials with this kind of loop can be identified as materials with relatively uniform channel like-pores (40). Likewise, the pore diameter distribution determined with BJH method showed a broad range of pore diameters, mostly below 4 nm for samples synthesized in the case of spherical, solid silica particles, specific surface areas as high as $403.501\, \text{m}^2/\text{g}$ correspond to diameters of 3.33 nm and total pore volume $0.5436\, \text{cm}^3/\text{g}$ which indicates the high porosity of the prepared sample (1,41).

Fig. 4. N$_2$ adsorption-desorption isotherms (a), the corresponding pore size distribution plots obtained by BJH method (b)

3.2. Oxidative/Adsorptive Desulfurization Process

The duel oxidative/adsorptive desulfurization of DBT was tested by using the prepared catalyst V$_2$O$_5$/MSS. The desulfurization process was carried by using either stirring system or ultrasonicator as batch reaction systems in presence of an oxidizing agent as H$_2$O$_2$. The model diesel fuel was prepared by dissolving DBT in n-hexane. Moreover, the duel oxidation-adsorption experiments were performed by changing temperature (30-70°C), initial concentration of DBT (200-1000 ppm) and the time from 10 to 300 min. To prove the coupling of catalytic oxidation and the adsorption process of prepared catalyst, the experiment was alternately carried out in the absence and presence of the V$_2$O$_5$/MSS or H$_2$O. First of all, the desulfurization process was carried on 600 ppm of DBT for 60 min under stirring system at 60°C, in the absence and presence of 0.02 g V$_2$O$_5$/MSS and/or 0.4 ml of H$_2$O$_2$. The removal efficiency (%E) was detected and presented in (Table. 1). From the obtained result, It is obvious that Using V$_2$O$_5$/MSS without the oxidizing agent showed low removal efficiency (%E) which prove the low adsorptive ability of V$_2$O$_5$/MSS toward DBT. However, using of H$_2$O$_2$ in absence of V$_2$O$_5$/MSS results in poor removal
which indicates the slow rate of DBT oxidation. Using both V₂O₅/MSS and H₂O₂ enhances significantly the removal (٪E) which indicates the duel oxidative adsorption property of the prepared catalyst. 

**Table 1. Removal (٪E) of DBT in presence and absence of V₂O₅/MSS**

| Experiment no. | V₂O₅/MSS (g) | H₂O₂ (ml) | (٪E)  
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.02</td>
<td>-</td>
<td>39.34</td>
</tr>
<tr>
<td>2</td>
<td>-</td>
<td>0.4</td>
<td>27.61</td>
</tr>
<tr>
<td>3</td>
<td>0.02</td>
<td>0.4</td>
<td>71.5</td>
</tr>
</tbody>
</table>

This data indicates the oxidative ability of the prepared catalyst V₂O₅/MSS of DBT by H₂O₂ producing sulfones and/or sulfoxides forms (scheme 1). In consequence, The adsorption capacity increases due to the greater adsorptive affinity toward sulfoxides and sulfones (polar O=S=O) than nonpolar thiophenic compounds (42,43).

**Scheme. 1. Catalytic Oxidative desulfurization of DBT to DBT sulfone**

3.2.2. Effect of contact time

In order to find a suitable equilibrium contact time, this test was carried out at time intervals ranging from 10 to 300 min. the removal (٪E) of model diesel of DBT using oxidative/adsorptive system as a function of interval contact time at 60 °C shown by Fig. 5 comparing both stirring and sonication techniques. At either stirring or sonication, the removal (٪E) increases by increasing the reaction time till equilibrium that’s due to the availability of vacant active sites on V₂O₅/MSS surface. then the oxidative desulfurization process to sulphone slows down due to occupying the active sites. Hence, In the case of stirring, 120 min was selected for further experiments. Comparing these results with the previous work indicates that, V₂O₅/MSS duel system achieved to equilibrium faster than those based only on adsorption (44,45), but the desulfurization process attained equilibrium 90 min in the case of sonication. This can be recognized by the specific effect of ultrasound waves, where micro-streaming bubbles are produced by the acoustic cavitation of the ultrasonic technique, which improves mass transfer in the reaction mixture (46,47).
Fig. 5. Effect of contact time (% E) under stirring or sonication on 0.02 g of adsorbent, 10 ml of 600 ppm DBT solution for 300 min

Adsorption kinetics study

Kinetics of the experimental results have been studied by applying the well-known pseudo-first-order and pseudo-second-order models (48,49). Their expressions of mathematics are given in Eqs. (3–5).

The pseudo-first-order model is written as follows:

$$\ln (q_e - q_t) = \ln q_e - k_1 t$$

(3)

The pseudo second order equation is stated as follows:

$$\frac{t}{q_t} = \frac{1}{k_2 q_e^2} + \frac{t}{q_e}$$

(4)

$$R_s = k_2 q_e^2$$

(5)

where $q_e$ and $q_t$ are the adsorbed amounts at equilibrium (mg g⁻¹) and time $t$ respectively while $k_1$ (min⁻¹) is the rate constant of pseudo first-order adsorption that calculated from linear plot of Log ($q_e - q_t$) versus time. $k_2$ (g.mg⁻¹.min⁻¹) is the rate constant of pseudo second order adsorption which can calculated by Plotting a straight line from $t/q_t$ verses $t$. the value of the correlation coefficient ($R^2$) was used to choose the best-fit model. Table 2 and (Fig. 6) showed a better fitting to pseudo–first–order model with $R^2 \approx 1$ under sonication or stirring. Also, it shows the comparable theoretical values $q_e$ with to the experimental values.

Table. 2. Detected kinetics parameters for duel oxidative adsorption of DBT (600 mg/l); 14 min at 60°C

<table>
<thead>
<tr>
<th>Conditions</th>
<th>experimental $q_e$ (mg/g)</th>
<th>pseudo-first order</th>
<th>pseudo-second order</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>$q_e$ (mg/g)</td>
<td>$K_1 \times 10^3$ (min⁻¹)</td>
</tr>
<tr>
<td>Stirring</td>
<td>299.268</td>
<td>317.770</td>
<td>0.008</td>
</tr>
<tr>
<td>Sonication</td>
<td>299.585</td>
<td>309.318</td>
<td>0.019</td>
</tr>
</tbody>
</table>
3.2.3. Effect of initial concentration

To find the maximum capacity of V₂O₅/MSS as adsorbent, five samples of DBT solution with different concentrations of 200, 400, 600, 800, and 1000 ppm were used. This parameter was carried by using 0.02g of V₂O₅/MSS duel system for 120 min at 60°C under stirring. As shown fig. 7 the amounts adsorbed (mg g⁻¹) increases with increasing of initial concentration that’s due to improve the mass transfer of adsorbate species from the bulk solution to the absorbent surface (44,45,50). Additionally, the sample did not achieve saturation up to initial concentration of 1000 mg L⁻¹.

![Graph showing adsorption capacity as function of initial concentration.](image)

**Fig. 7.** Adsorption capacity as function of DBT initial concentration at 60°C under stirring on 0.02 g of adsorbent, 10 ml of 600 ppm DBT solution for 120 min

3.4. Adsorption isotherm

Models of adsorption isotherms were also used to study the Adsorption process at equilibrium as a function of initial DBT concentration. Two models Langmuir and Freundlich. The Langmuir isotherm (51) can be expressed mathematically as the following Eq (6).
\[
\frac{C_e}{q_e} = \frac{1}{bQ_e} + \frac{C_e}{Q_e}
\]  

(6)

Where \(C_e\) is the equilibrium concentration of DBT in solution (mg/L), \(Q_e\) is the maximum theoretical capacity at the monolayer (mg/g) and \(b\) is the constant of Langmuir (l/mg).

The Freundlich isotherm (52) can be expressed mathematically as the following Eq (7).

\[
\log q_e = \log k_f + \frac{1}{n} \log C_e
\]

(7)

where \(q_e\) (mg/g) is the DBT amount adsorbed at equilibrium, \(C_e\) (mg/L) is the remaining concentration of DBT in model oil, \(K_F\) the Freundlich constant \([\text{mg/g (mg/l)}^n]\) represents adsorption capacity and \(n\) is adsorption favorability constant which they are calculated from slope and intercepts of linear plot of \(\ln q_e\) against \(\ln C_e\). The value of \(1/n\) is being closer to 0, which implies that the adsorbent surface is more heterogeneous and promotes the multilayer adsorption process.

Table. 3. and Fig. (8) showed the calculated values of models. The model of Langmuir isotherm has a good fittings of the experimental data with \(R^2 = 0.909\) which suggests a monolayer adsorption process on the homogeneous surface of the adsorbent.

### Table. 3. Obtained data from fitting experimental results with the three models:

<table>
<thead>
<tr>
<th>Model</th>
<th>(b) (L mg(^{-1}))</th>
<th>(q_{max}) (mg g(^{-1}))</th>
<th>(R^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Langmuir</td>
<td></td>
<td></td>
<td>0.909470584</td>
</tr>
<tr>
<td>Freundlich</td>
<td></td>
<td></td>
<td>0.07895289</td>
</tr>
</tbody>
</table>

Table. 3. Obtained data from fitting experimental results with the three models:

![Graph](image_url)

**Fig. 8.** The dual oxidative adsorption models for DBT removal: (a) Langmuir isotherm, and (b) Freundlich isotherm.
3.5. Adsorption mechanism

It is known that the adsorption mechanism follows a complex pathway and is followed by both the surface and the diffusion of the pore but in different areas. The intra-particle diffusion model was evaluated according to “Weber and Morris equation” to investigate the diffusion process (53,54).

The initial rate of intra-particle diffusion in this model can be expressed mathematically as the following equation:

\[ q_t = k_i t^{1/2} + C \]  

where \( k_i \) is the constant of the intra-particle diffusion rate (mg/g.min\(^{0.5}\)), and \( C \) (mg/g) is a constant attributed to the boundary layer effect and obtained from the intercept value by plotting \( q_t \) versus \( t^{0.5} \). In this equation, the \( C \) value give a picture about the rate-determining step where, \( C = 0 \), intraparticle diffusion is the only rate-determining step and \( C \neq 0 \), the process was controlled by a variety of mechanism.

As shown in (Fig. 9), the multilinear relationship with \( C > 0 \), indicate that, more than one step occurred during the desulfurization process over the prepared catalyst adsorbent (V_2O_5@MSS) system under both stirring and ultrasonication. This implies that adsorption takes place in three stages: the early stage is the rapid adsorption of the surface, followed by the slower stage of intra-particle diffusion and finally the equilibrium stage (46,47).

![Intraparticle diffusion model](image)

**Fig. 9. Intra-particle diffusion plot for adsorption of 600 mg/l of as initial concentration of DBT at 60°C for 120 min**
3.6. Thermodynamics studies

The key factor for both catalytic and adsorptive desulfurization reactions is temperature (55). The influence of temperature was set on the DBT solution desulfurization process using V$_2$O$_5$ / MSS as a dual catalyst-adsorbent system has been investigated. The adsorption process was performed at various temperatures under stirring (303, 313, 323, 333 and 343 K). (Fig. 10) displays the relation between adsorption capacity ($q_e$) and temperature. It shows that a gradual increase in removal (percent E) was obtained by increasing the temperature applied from 303 to 333 K and improving the dual oxidative / adsorption process. However, a further rise in the temperatures of reaction leads to a reduction in the potential of adsorption. This has been shown by the decomposition of H$_2$O$_2$ into water, which reduces sulfone production (43). These results indicate that the process of desulfurization using V$_2$O$_5$ / MSS occurs adequately at temperatures equal to or even lower than 333 K (56).

![Graph](image)

**Fig. 10.** The adsorption capacity as a function of applied temperature using V$_2$O$_5$/MSS under stirring; 600 ppm of DBT as initial concentration at 60°C for 120 min.

The thermodynamic adsorption parameters as change of free Gibbs energy ($\Delta G$), change of enthalpy ($\Delta H$) (kJ mol$^{-1}$) and change of entropy ($\Delta S$) (J mol$^{-1}$ K$^{-1}$) were determined by using the results of experimental adsorption isotherms up to 333 K. The thermodynamic adsorption parameters ($\Delta H$, $\Delta G$) and ($\Delta S$) were calculated by using the following equations (9-11):
\[ K_d = \frac{q_e}{C_e} \]  

(9)

\[ \Delta G = -RT \ln K_d \]  

(10)

\[ \ln K_d = \frac{\Delta S}{R} - \frac{\Delta H}{RT} \]  

(11)

where \( R \) (8.314 X10^{-3} \text{ kJ K}^{-1} \text{ mol}^{-1}) is the gas constant and \( T \) is the applied kelvin temperature (K). From (Eq. 13), \( \Delta G \) was demonstrated while from slope and intercept from the plot of \( \ln K_d \) versus \( 1/T \) of (Eq. 14), (\( \Delta H \)) and (\( \Delta S \)) were measured. The values of \( \Delta H \) and \( \Delta S \) are presented in (Table. 4) (\( \Delta S \)) and (\( \Delta H \)) showed positive values that suggest that the adsorption mechanism of endothermic catalytic oxidation contributes to a growing disorder. (\( \Delta S \)) and (\( \Delta H \)) show positive values and hence the negative sign of (\( \Delta G \)) indicates the spontaneity of adsorption process (54,57,58).

It showed positive values

**Table. 4. Thermodynamic parameters for duel oxidation adsorption process of DBT (600 mgg^{-1}) for 120 min under stirring.**

<table>
<thead>
<tr>
<th>( T ) (K)</th>
<th>( \ln K_d )</th>
<th>( \Delta G ) (kJ mol^{-1})</th>
<th>( \Delta H ) (kJ mol^{-1})</th>
<th>( \Delta S ) (J mol^{-1} K^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>303</td>
<td>0.332362832</td>
<td>-0.837269169</td>
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</tr>
<tr>
<td>313</td>
<td>0.778222795</td>
<td>-2.02515517</td>
<td>37.57273881</td>
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<tr>
<td>323</td>
<td>1.807364104</td>
<td>-4.853535328</td>
<td></td>
<td></td>
</tr>
<tr>
<td>333</td>
<td>5.347294101</td>
<td>-14.80431525</td>
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</tr>
</tbody>
</table>

**CONCLUSION**

In conclusion, our study successfully synthesized mesoporous silica spheres via micelle-template and \( V_2O_5 \) nano-particles supported on it (\( V_2O_5/MSS \)) through impregnation technique. This novel catalyst proved its effect as a novel duel oxidative/adsorptive system of model diesel fuel. The Desulfurization process is performed in the presence a suitable oxidizing agent (\( H_2O_2 \)) at different conditions of temperature, contact time and initial concentration under either stirring or ultrasonication. The novel catalyst shows an effective removal about 100 % under sonication within 1.5h but in stirring within 2h. The pseudo-first - order model could explain adsorption kinetics (\( V_2O_5 / MSS \)) of the studied adsorbents. The experimental data matches well with the Langmuir model of the isotherm which indicates a monolayer adsorption process on the homogeneous surface of the adsorbent. The determined thermodynamic parameters suggested that the endothermic adsorption process was favorable for (\( V_2O_5 / MSS \)) adsorption.

**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.
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Microwave-induced catalytic oxidative desulfurization of gasoil fraction over phosphotungstic acid-based magnetic silica (Ni@SiO2\PWA) nanocatalyst. Catalysis Communications.;136:105926.


59. تحضير خامس أكسد الفانديم النانوني المحمل على كريات السيكا المسامية (V₃O₅/MSS) لإزالة الكبريت بطريقة الأكسدة-الامتزازية الجديدة

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تم تحضير خامس أكسد الفانديم النانوني المحمل على كريات السيكا المسامية (V₃O₅/MSS) باستخدام تقنية الشرب impregnation. حيث تم تحضير كريات السيكا المسامية باستخدام FTIR و XRD، و أظهرت النتائج أن العينات كانت غير متبمورة ومسامية الشكل، وذات مساحات داخلية خاصة، وكذلك مساحة سطح مميزة عالية. تم اختبار العينة المحضرة للآداء الأكسيدي / الامتزازي oxidative/adsorptive بواسطة توضيح وقود الديزل. تم اختبار عملية إزالة الكبريت في نموذج H₂O₂ مع عامل مؤكسد تحت ظروف تفاعل مختلفة (درجة الحرارة ووقت التلامس والتركيز الأولي) باستخدام نظام المفاعل الثابت عن طريق القليل أو في وجود الموجات فوق الصوتية. تشير البيانات إلى أن V₃O₅/MSS يظهر أداءً جيدًا تجاه إزالة الكبريت في ظروف العناية بالإضافة إلى أن العينات كانت غير متبمورة ومسامية الشكل، وذات مساحات داخلية خاصة، وكذلك مساحة سطح مميزة عالية. تم اختبار العينة المحضرة للآداء الأكسيدي / الامتزازي oxidative/adsorptive بواسطة توضيح وقود الديزل. تم اختبار عملية إزالة الكبريت في نموذج H₂O₂ مع عامل مؤكسد تحت ظروف تفاعل مختلفة (درجة الحرارة ووقت التلامس والتركيز الأولي) باستخدام نظام المفاعل الثابت عن طريق القليل أو في وجود الموجات فوق الصوتية. تشير البيانات إلى أن V₃O₅/MSS يظهر أداءً جيدًا تجاه إزالة الكبريت في ظروف العناية بالإضافة إلى أن العينات كانت غير متبمورة ومسامية الشكل، وذات مساحات داخلية خاصة، وكذلك مساحة سطح مميزة عالية. تم اختبار العينة المحضرة للآداء الأكسيدي / الامتزازي oxidative/adsorptive بواسطة توضيح وقود الديزل. تم اختبار عملية إزالة الكبريت في نموذج H₂O₂ مع عامل مؤكسد تحت ظروف تفاعل مختلفة (درجة الحرارة ووقت التلامس والتركيز الأولي) باستخدام نظام المفاعل الثابت عن طريق القليل أو في وجود الموجات فوق الصوتية. تشير البيانات إلى أن V₃O₅/MSS يظهر أداءً جيدًا تجاه إزالة الكبريت في ظروف العناية بالإضافة إلى أن العينات كانت غير متبمورة ومسامية الشكل، وذات مساحات داخلية خاصة، وكذلك مساحة سطح مميزة عالية. تم اختبار العينة المحضرة للآداء الأكسيدي / الامتزازي oxidative/adsorptive بواسطة توضيح وقود الديزل. تم اختبار عملية إزالة الكبريت في نموذج H₂O₂ مع عامل مؤكسد تحت ظروف تفاعل مختلفة (درجة الحرارة ووقت التلامس والتركيز الأولي) باستخدام نظام المفاعل الثابت عن طريق القليل أو في وجود الموجات فوق الصوتية. تشير البيانات إلى أن V₃O₅/MSS يظهر أداءً جيدًا تجاه إزالة الكبريت في ظروف العناية بالإضافة إلى أن العينات كانت غير متبمورة ومسامية الشكل، وذات مساحات داخلية خاصة، وكذلك مساحة سطح مميزة عالية. تم اختبار العينة المحضرة للآداء الأكسيدي / الامتزازي oxidative/adsorptive بواسطة توضيح وقود الديزل. تم اختبار عملية إزالة الكبريت في نموذج H₂O₂ مع عامل مؤكسد تحت ظروف تفاعل مختلفة (درجة الحرارة ووقت التلامس والتركيز الأولي) باستخدام نظام المفاعل الثابت عن طريق القليل أو في وجود الموجات فوق الصوتية. تشير البيانات إلى أن V₃O₅/MSS يظهر أداءً جيدًا تجاه إزالة الكبريت في ظروف العناية بالإضافة إلى أن العينات كانت غير متبمورة ومسامية الشكل، وذات مساحات داخلية خاصة، وكذلك مساحة سطح مميزة عالية. تم اختبار العينة المحضرة للآداء الأكسيدي / الامتزازي oxidative/adsorptive بواسطة توضيح وقود الديزل. تم اختبار عملية إزالة الكبريت في نموذج H₂O₂ مع عامل مؤكسد تحت ظروف تفاعل مختلفة (درجة الحرارة ووقت التلامس والتركيز الأولي) باستخدام نظام المفاعل الثابت عن طريق القليل أو في وجود الموجات فوق الصوتية. تشير البيانات إلى أن V₃O₅/MSS يظهر أداءً جيدًا تجاه إزالة الكبريت في ظروف العناية بالإضافة إلى أن العينات كانت غير متبمورة ومسامية الشكل، وذات مساحات داخلية خاصة، وكذلك مساحة سطح مي...