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Hydrodeoxygenation of Anisole over Ni/ α -Al₂O₃ CatalystWen-wu Tang^{a,c}, Xing-hua Zhang^{b,c*}, Qi Zhang^{b,c}, Tie-jun Wang^{b,c}, Long-long Ma^{b,c*}*a. Department of Chemistry, University of Science and Technology of China, Hefei 230026, China**b. Key Laboratory of Renewable Energy, Chinese Academy of Sciences, Guangzhou 510640, China**c. Guangzhou Institute of Energy Conversion, Chinese Academy of Sciences, Guangzhou 510640, China*

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Ni-based catalysts supported on different supports (α -Al₂O₃, γ -Al₂O₃, SiO₂, TiO₂, and ZrO₂) were prepared by impregnation. Effects of supports on catalytic performance were tested using hydrodeoxygenation reaction (HDO) of anisole as model reaction. Ni/ α -Al₂O₃ was found to be the highest active catalyst for HDO of anisole. Under the optimal conditions, the anisole conversion is 93.25% and the hydrocarbon yield is 90.47%. Catalyst characterization using H₂-TPD method demonstrates that Ni/ α -Al₂O₃ catalyst possesses more amount of active metal Ni than those of other investigated catalysts, which can enhance the catalytic activity for hydrogenation. Furthermore, it is found that the Ni/ α -Al₂O₃ catalyst has excellent repeatability, and the carbon deposited on the surface of catalyst is negligible.

Key words: Anisole, Hydrodeoxygenation, Ni/ α -Al₂O₃, Hydrocarbons

I. INTRODUCTION

Lignin, a three-dimensional amorphous polymer consisting of methoxylated phenylpropane structures, is contained in plant biomass with high proportion [1–3]. It can be transformed into phenolic compounds via depolymerization [4, 5]. It has been reported that yield of 13.2% phenolic monomers was obtained from organosolv pine lignin via catalytic depolymerization in the presence of MgO and THF [6]. However, obtained phenolic products were complex mixtures with high oxygen content [7, 8]. The high oxygen content led to many undesirable properties, such as high viscosity, thermal instability, corrosiveness, poor heating value and immiscibility with hydrocarbon fuels [9, 10]. Hence, the removal of oxygen is required to obtain hydrocarbons as high-graded fuels.

Hydrodeoxygenation (HDO) is an effective method for the production of hydrocarbons from lignin-derived phenolic compounds [11–13]. Bifunctional catalyst comprised by active metal and solid acid exhibits excellent activity for HDO reaction. For example, guaiacol was completely converted with high selectivity for cyclohexane over Ni/SiO₂-ZrO₂ catalyst [10]. Normally, it is considered that the metal Ni acts as hydrogenation activity center while the support SiO₂-ZrO₂ provides acidic sites for the HDO reaction.

Support material is a crucial factor determining the catalytic performance. In the past years, γ -Al₂O₃

was widely used as catalyst support for HDO catalyst due to its cheap cost, excellent texture, and suitable acidity. For example, the HDO of guaiacol with conversion of 100% and yield of 88% of cyclohexane was obtained over the Pd-based catalyst supported on γ -Al₂O₃ [14]. Oxygen contained in phenolics was also efficiently removed by hydrodeoxygenation with the sulfided NiMo/ γ -Al₂O₃ and CoMo/ γ -Al₂O₃ catalysts [7, 15]. However, large amount of coke formed and deposited on the surface of the catalyst during the HDO reaction process [7, 15]. Worse, partial γ -Al₂O₃ can be transformed into boehmite under hydrothermal conditions, resulting in a decrease for catalytic activity [16]. To overcome these drawbacks, many support materials such as SiO₂ [17], TiO₂ [18], and ZrO₂ [15] were also explored in the past years, exhibiting impressive catalytic activity for HDO reaction.

α -Al₂O₃ is a type of mesoporous material with excellent hydrothermal stability. However, scarce work has been done to remove oxygen from phenolics with the HDO catalyst supported on α -Al₂O₃. In this work, HDO reaction of anisole with the Ni-based catalyst supported on α -Al₂O₃ was investigated to explore the effect of support on the catalytic properties. For comparison, the Ni-based catalysts supported on γ -Al₂O₃, SiO₂, TiO₂, and ZrO₂ were also investigated in the HDO reaction of anisole.

II. EXPERIMENTS

A. Catalyst preparation

SiO₂ was prepared by chemical precipitation using NH₄NO₃ and Na₂SiO₃ as materials. The precipitate was dried at 120 °C overnight and then calcinated at

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500 °C for 4 h. Similarly, ZrO₂ was prepared by chemical precipitation using NH₃·H₂O and ZrOCl₂ as material. TiO₂ was prepared by hydrolysis of TiCl₄. In addition, support materials α-Al₂O₃ and γ-Al₂O₃ were purchased from Aladdin Industrial Co. Ltd.

Catalysts Ni/α-Al₂O₃, Ni/γ-Al₂O₃, Ni/SiO₂, Ni/TiO₂ and Ni/ZrO₂ with 10 wt% Ni loadings were prepared by wet impregnation using Ni(NO₃)₂·6H₂O as nickel precursor. Firstly, supporting material dipped into the nickel nitrate solution. Then the solution was heated with continuous stir until water was evaporated to dryness. The obtained residues were dried at 120 °C overnight and calcinated at 500 °C for 4 h. The prepared catalyst was crushed into powder with sizes of 100–200 mesh.

The prepared catalyst Ni/α-Al₂O₃ was reduced at 700 °C while the other catalysts were reduced at 550 °C for 5 h in H₂ atmosphere before using.

B. Catalyst characterization

Brunauer-Emmett-Teller (BET) surface area (S_{BET}), average pore diameter, and pore volume of the catalyst were measured by N₂ isothermal (−196 °C) physisorption through Autosorb-iQ-2 (Qudrasorb SI, Quantachrome Instruments). The catalyst was degassed for 12 h at 250 °C under vacuum condition before N₂ adsorption.

H₂-temperature programmed reduction (H₂-TPR), NH₃-temperature programmed desorption (NH₃-TPD), and H₂-temperature programmed desorption (H₂-TPD) measurements were carried out on an automatic chemical adsorption instrument (CBP-1, Quantachrome Instruments) equipped with a thermal conductivity detector. Dispersity of Ni over different supports was calculated with the following formula:

$$D = \frac{\text{Mole of adsorbed hydrogen}}{\text{Total moles of Ni in the catalyst}} \quad (1)$$

X-ray diffraction (XRD) analysis of catalyst was carried out on an equipment (PANalytical, Netherlands) with Cu Kα ($\lambda=0.154$ nm) radiation. X-ray photoelectron spectroscopy (XPS) analysis of the catalyst was performed on the instrument of Thermo Scientific ESCALAB 250XI using Al Kα_{1,2} radiation as spectra excitation. Thermogravimetry (TG) analysis of the used catalyst was carried out under a flow of air (50 mL/min) on a thermal analyzer (TGA Q50, US). The heating rate is 15 K/min. Transmission electron microscope (TEM) profiles were obtained on a JEM-2100F (JEOL, Japan) instrument equipped with an EDX spectrometer.

C. Catalytic activity test

HDO process of anisole was carried out in a 50 mL stainless steel autoclave equipped with electric mechanical agitator. For each run, catalyst (0.5 g), solvent

n-octane (21.5 mL) and anisole were loaded into the autoclave. 5.0 MPa H₂ was pressured into the reactor after displacing the air. The reactor was heated to a desired reaction temperature while the reagents were stirred at a rate of 800 r/min. Liquid product was collected for subsequent analysis when the reaction was completed.

D. Products analysis

Liquid products obtained from the HDO of anisole were analyzed by gas chromatography (Shimadzu GC-2010 with a FID detector and a DB-5 column) and GC-MS (Agilent 7890A-5975C with DB-FFAP capillary column). The carrier gas was He (99.995% purity), and the oven temperature program increased from 50 °C (holding for 1 min) to 260 °C (holding for 10 min) at a rate of 10 °C/min.

Anisole conversion (C) and hydrocarbon yield (Y_h) were determined with the following equation:

$$C = \frac{\text{Mole}(\text{anisole})_{\text{initial}} - \text{Mole}(\text{anisole})_{\text{end}}}{\text{Mole}(\text{anisole})_{\text{initial}}} \quad (2)$$

$$Y_h = \frac{\text{Mole of hydrocarbons}}{\text{Initial mole of anisole}} \quad (3)$$

III. RESULTS AND DISCUSSION

A. Catalyst characterization

The parameters of the catalysts are shown in Table I and the structures are in Fig.S1 (supplementary materials). It was found that all tested catalysts were mesoporous materials. Their average pore diameter is in the range of 7–29 nm, and the most probable pore diameter is in the range of 3–60 nm. Among the investigated catalysts, the average pore diameter and the most probable pore diameter of Ni/α-Al₂O₃ catalyst were the smallest. They were 7.3 and 3.9 nm, respectively. In addition, TEM image of the Ni/α-Al₂O₃ catalyst was presented in Fig.S2 (supplementary material). It can be clearly seen that the metal Ni was well dispersed on the surface of α-Al₂O₃. The size of Ni particle is about 30–40 nm.

XRD profiles of different catalysts are shown in Fig.1. The weak and broad characteristic peaks centered at 2θ of 37.3°, 43.3°, and 62.9° were assigned to NiO. When XRD characteristic peak of the samples showed a weak and broad peak of NiO species, it actually indicated that NiO particles were very small and widely dispersed on the catalyst support [19]. Notably, the catalyst supported on α-Al₂O₃ showed the smallest NiO crystallite size among the tested catalysts, as shown in Table I. The reason might be that NiO well dispersed on the support, suppressing its aggregation.

H₂-TPR profiles of catalyst are shown in Fig.2(a). The peaks positioned at lower reduction temperature

TABLE I Parameters of the catalysts.

Catalyst	$S_{\text{BET}}/(\text{m}^2/\text{g})$	$V_{\text{total}}/(\text{cm}^3/\text{g})$	D_p/nm	D_m/nm	NiO size/nm
Ni/ α -Al ₂ O ₃	172.9	0.316	7.3	3.9	11.4
Ni/ γ -Al ₂ O ₃	133.3	0.662	19.9	17.1	30.2
Ni/SiO ₂	213.1	1.131	21.2	17.2	17.2
Ni/TiO ₂	13.1	0.094	28.6	59.0	21.5
Ni/ZrO ₂	34.2	0.153	17.9	17.0	17.8

Note: D_p is average pore diameter. D_m is the most probable pore diameter. NiO size is the size of NiO crystallite and calculated by Scherrer equation, the Bragg angle of 2θ was $\sim 43.3^\circ$.

TABLE II Ni dispersion and H₂ uptakes of different catalysts.

Catalysts	H ₂ uptaking/ $(\mu\text{mol}/\text{g})$	Ni dispersion/%
Ni/ α -Al ₂ O ₃	976.33	13.50
Ni/ γ -Al ₂ O ₃	459.72	6.36
Ni/SiO ₂	566.95	7.84
Ni/TiO ₂	201.86	2.79
Ni/ZrO ₂	498.57	6.90

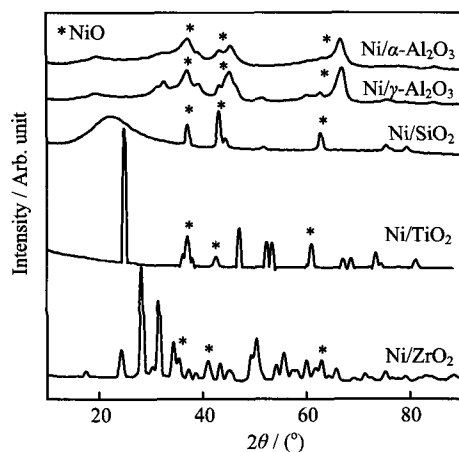
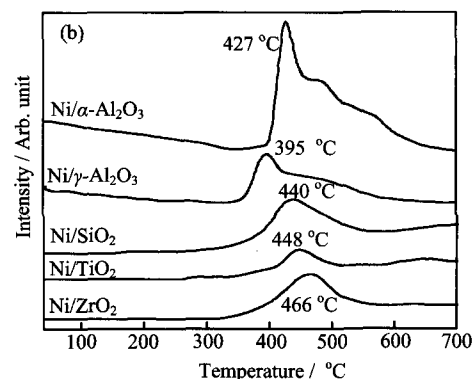
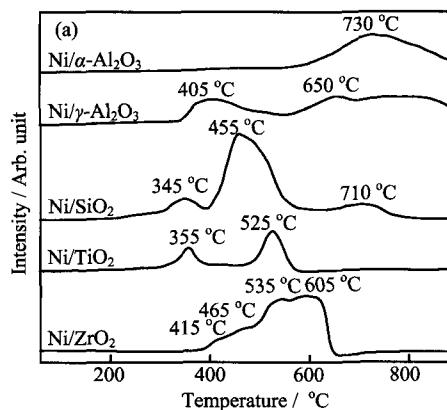


FIG. 1 XRD profiles of different catalysts.

(usually below 400 °C) were ascribed to the superficial NiO, which can be easily reduced to Ni⁰. The peaks at higher temperature were associated with the reduction of bulk NiO, which weakly interacted with support material. It is noteworthy that the reduction peak of Ni/ α -Al₂O₃ was centered at 730 °C, suggesting a stronger interaction between NiO and the support [20].

H₂-TPD experiments of different catalysts were also carried out, and the results are shown in Fig.2(b). H₂ desorption peaks stepped from about 320 °C for all investigated catalysts. The quantitative determination based on the H₂-TPD profiles demonstrated that the H₂ uptakes of Ni/ α -Al₂O₃ (976.33 $\mu\text{mol}/\text{g}$) were far more than that of the Ni-based catalysts supported on γ -Al₂O₃, SiO₂, ZrO₂, and TiO₂ (as shown in Table II).

FIG. 2 (a) H₂-TPR and (b) H₂-TPD profiles of different catalyst samples.

In addition, Ni dispersion degree of Ni/ α -Al₂O₃ (13.5%) was also higher than that of other tested catalysts. This conclusion is in good agreement with that of XRD analysis.

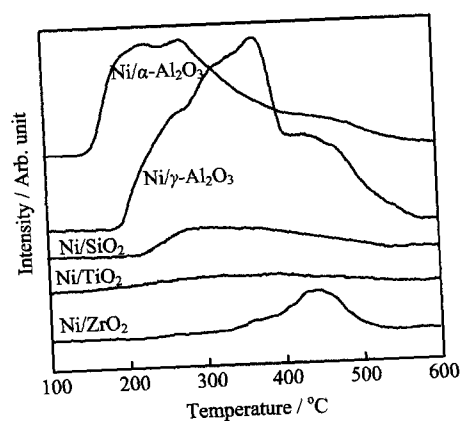
NH₃-TPD experiment was carried out to investigate the acidity of the catalyst. It is widely accepted that peaks at a temperature range of 100–400 °C are assigned to desorption of NH₃ on weak acidic sites and that peaks at a temperature range of 400–600 °C are assigned to the desorption of NH₃ on strong acidic sites. As shown in Fig.3, a NH₃ desorption band was observed in the pattern of Ni/ α -Al₂O₃, suggesting the presence of acidic sites. Normally, α -Al₂O₃ is an in-

TABLE III Anisole conversions and product yields over different catalysts.

Catalyst	Anisole conv./%	Yield/%				
		Hydrocarbons	Methoxy-cyclohexane	Cyclohexanol	Phenol	<i>o</i> -Cresol
Ni/ α -Al ₂ O ₃	93.25	90.47	0.81	0.63	0.92	0.31
Ni/ γ -Al ₂ O ₃	99.62	73.83	0.53	0	20.83	1.27
Ni/SiO ₂	99.40	71.98	0	0.16	17.27	0
Ni/TiO ₂	77.95	56.13	0	4.97	15.39	0
Ni/ZrO ₂	93.52	62.02	0.32	0	25.91	2.88
α -Al ₂ O ₃	10.33	1.47	0	0.16	6.65	0.97

TABLE IV Different hydrocarbons yields (%) over different catalysts.

Catalyst	Methylpentane	Cyclohexane	Cyclohexene	Methylcyclohexane	Toluene	Phenylcyclohexane
Ni/ α -Al ₂ O ₃	0.15	90.02	0.17	0	0	0.13
Ni/ γ -Al ₂ O ₃	0.69	50.83	4.13	2.11	15.91	0.16
Ni/SiO ₂	0.94	66.24	2.34	0	2.46	0
Ni/TiO ₂	0.24	54.83	0.35	0.58	0	0.13
Ni/ZrO ₂	0.79	57.37	1.60	0.34	1.74	0.18
α -Al ₂ O ₃	0	0.67	0.24	0	0.56	0

FIG. 3 NH₃-TPD profiles of the different catalyst samples.

ertial material. Detected acidic sites were deemed to be created by the introduction of Ni²⁺ cation [20]. It has been reported that trace Ni could be incorporated into α -Al₂O₃ through ion exchange [20]. A broad NH₃ desorption band was also observed as expected in the pattern of Ni/ γ -Al₂O₃ catalyst. Compared with that of Ni/ α -Al₂O₃, NH₃ desorption peak of Ni/ γ -Al₂O₃ moved to higher temperature, suggesting stronger acidity. With regard to catalysts of Ni/SiO₂ and Ni/ZrO₂, relatively small peaks of NH₃ desorption were observed, suggesting a small amount of acidic sites. Notably, no NH₃ desorption peak was observed in the case of Ni/TiO₂.

B. Catalyst testing

Catalytic performances of catalysts and pure support α -Al₂O₃ were evaluated in HDO reaction process with

anisole as substrate. As shown in Table III, the highest yield of hydrocarbons (90.47%) was obtained over Ni/ α -Al₂O₃ catalyst. Cyclohexane was the main component, and its yield was as high as 90.02% (Table IV). This result indicates that the catalytic performance of α -Al₂O₃ was obviously superior to other catalysts under the same reaction conditions. This can be explained as: H₂ uptake of Ni/ α -Al₂O₃ was obviously higher than that of other catalysts as discussed in H₂-TPD analysis suggesting more active nickel, thereby causing a better catalytic activity for hydrogenation. In addition, a large number of acidic sites were observed on the surface of Ni/ α -Al₂O₃. These acidic sites could co-work well together with active metal sites, and facilitate the HDO reaction [21]. However, the yield of hydrocarbons was rather low (1.47%) when pure support α -Al₂O₃ was used as catalyst, suggesting a negligible catalytic activity for the HDO reaction.

The conversion of anisole was as high as 99.62% over the catalyst Ni/ γ -Al₂O₃. The major reason is the strong acidity of catalyst favors activation of the C—O bond, resulting in the cleavage of C_{Ar}O—CH₃ bond [2]. It was reported that the phenolic compounds could also be converted over the support of γ -Al₂O₃ alone [2]. However, compared with that of Ni/ α -Al₂O₃ catalyst, the yield of hydrocarbon products was relatively low (only 73.83%). One of the reasons was ascribed to inefficient hydrogenation activity. For similar reasons, the hydrocarbon yields over Ni/SiO₂ and Ni/ZrO₂ were also relatively low.

It should be noted that the yield of toluene (Table IV) was 15.91% over the catalyst Ni/ γ -Al₂O₃. This may also relate to the acidity of the catalyst. Normal anisole molecule can be adsorbed on the acidic sites due to its well accessible basic oxygen electronic

blet, resulting in the activation of ArO–CH₃ bond. The activated ArO–CH₃ bond cleaved heterolytically, and the positively charged methyl group can be transferred to aromatic ring, leading to the formation of methylated products [24]. Therefore, it is plausible that the catalyst Ni/ γ -Al₂O₃ favors the transmethylation due to its strong acidity, yielding higher methylated products during the HDO process of anisole. Similarly, methylated products were also observed over catalysts of Ni/SiO₂ and Ni/ZrO₂, although it was minor.

In addition, anisole conversion and hydrocarbon yield were relatively poor over the catalyst Ni/TiO₂. There are two possible reasons accounting for this. Firstly, H₂ uptake of Ni/TiO₂ is far less than that of other catalysts as discussed in H₂-TPD experiments, suggesting less active nickel. Moreover, the acidity of Ni/TiO₂ is almost negligible as discussed in NH₃-TPD experiments. Two handicaps resulted in an inferior catalytic activity for HDO reaction. Secondly, the BET surface area, Ni dispersion and porous structure of Ni/TiO₂ are clearly inferior to those of other investigated catalysts, which are also unfavorable for the improvement of catalytic activity.

Effects of reaction temperature, time and catalyst dosage on HDO of anisole were investigated carefully using Ni/ α -Al₂O₃ as a representative catalyst. As shown in Fig.4(a), the anisole conversion gradually increased with the increased temperature. The yield of hydrocarbons also increased gradually from 38.90% to 90.47% when the reaction temperature increased from 220 °C to 300 °C. However, yield of hydrocarbons swiftly decreased as the reaction temperature further increased from 300 °C to 340 °C. In addition, bicyclic compound (phenylcyclohexane) noticeably increased in the products at 340 °C, suggesting that the intermolecular polymerization took place at a higher temperature. Detailed product distribution was presented in Table S1 (supplementary materials).

Reaction time significantly influenced the HDO reaction of anisole. As shown in Fig.4(b) and Table S2 (supplementary material), the conversion of anisole and the yield of hydrocarbons gradually increased with reaction time over Ni/ α -Al₂O₃. At the beginning of the reaction, the yield of phenol is considerably high, and gradually reduced with reaction time being lengthened. According to this result, it can be speculated that anisole was converted to phenol via demethylation firstly, followed by hydrogenation of aromatic ring to form cyclohexanol and then cyclohexane. The detected cyclohexanol and cyclohexene were other evidences for this presumption. Interestingly, methoxycyclohexane was detected in the products obtained over Ni/ α -Al₂O₃, suggesting that hydrogenation of the aromatic ring of anisole could occur prior to the demethylation. Generally, methoxycyclohexane was only mentioned as a reaction intermediate or product over noble metal catalyst due to its efficient hydrogenation activity [25].

The effect of catalyst dosage on the HDO of anisole

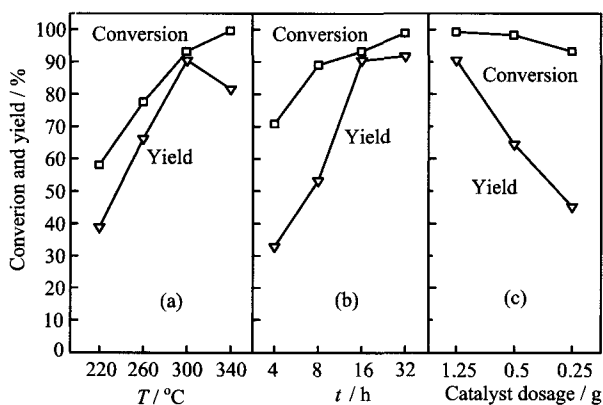


FIG. 4 Anisole conversion and hydrocarbon yield varied with (a) temperature, (b) time, and (c) catalyst dosage over Ni/ α -Al₂O₃ catalyst. Reaction conditions: (a) catalyst of 0.5 g, anisole of 2.5 g, octane of 21.5 mL, P_{H_2} =5.0 MPa, t =16 h; (b) catalyst of 0.5 g, anisole of 2.5 g, octane of 21.5 mL, P_{H_2} =5.0 MPa, T =300 °C; (c) anisole of 2.5 g, octane of 21.5 mL, P_{H_2} =5.0 MPa, T =300 °C, t =16 h.

was also investigated in detail. As shown in Fig.4(c), the anisole conversion and the hydrocarbons yield gradually increased with the increased catalyst dosage in the investigated range. Generally, higher catalyst dosage implies more catalytic active sites, thereby resulting in better catalytic performance [26]. Therefore, it is plausible that this process is of high catalyst dosage dependence.

The repeatability of Ni/ α -Al₂O₃ was also tested. The catalyst was reused directly without any retexture (such as calcination and reduction). As shown in Fig.5(a), anisole was efficiently converted with high hydrocarbons yield when the catalyst Ni/ α -Al₂O₃ was repeatedly used. This result implied that the catalyst Ni/ α -Al₂O₃ had a stable catalytic activity for the HDO of anisole. To examine the extent of coke formed during the HDO process, TG analysis of reused catalyst was carried out and the result was exhibited in Fig.S3 (supplementary materials). Only a slight weight loss was observed even if the catalyst was repeatedly reused for five times. This result suggests that the catalyst Ni/ α -Al₂O₃ possesses excellent resistance to coking. However, as shown in the XPS patterns of catalysts (Fig.5(b)), the peak of Ni⁰ positioned at 852.54 eV became small when the catalyst was repeatedly used for five times. Conversely, the peak of Ni²⁺ positioned at about 856 eV intensified obviously. This result suggests that active metal Ni⁰ would be oxidized to Ni²⁺ during the HDO process of anisole.

IV. CONCLUSION

Effects of support materials α -Al₂O₃, γ -Al₂O₃, SiO₂, TiO₂, and ZrO₂ on catalytic performance of Ni-based catalysts were studied for the HDO of anisole to hydro-

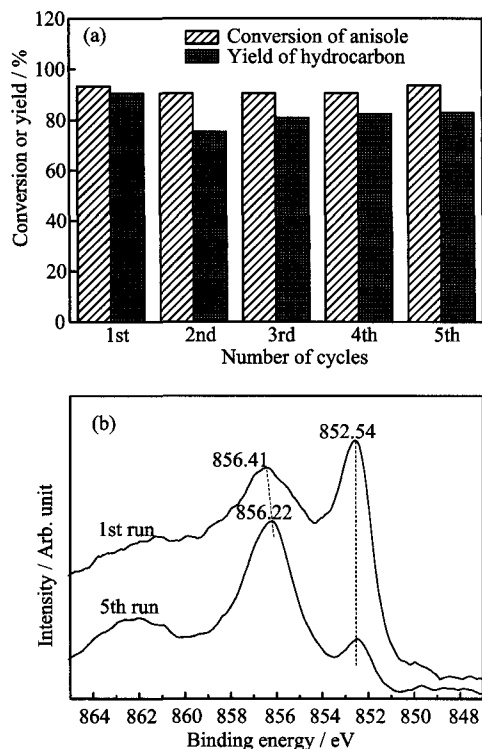


FIG. 5 (a) Catalyst repeatability test and (b) surface chemical state of catalysts. Reaction conditions: catalyst Ni/ α -Al₂O₃ of 0.5 g, anisole of 2.5 g, octane of 21.5 mL, P_{H_2} =5.0 MPa, T =300 °C, and t =16 h.

carbons. The Ni/ α -Al₂O₃ catalyst exhibited the most H₂ uptakes and the highest catalytic activity among these investigated catalysts, revealing that the higher hydrogenation activity can promote the HDO of anisole, obtaining higher hydrocarbon yield. Furthermore, the Ni/ α -Al₂O₃ catalyst exhibited excellent repeatability for the HDO of anisole. However, the result of XPS analysis of the used catalyst demonstrated that active metal Ni⁰ was oxidized to Ni²⁺ during the HDO process when the catalyst was repeatedly used for five times.

Supplementary materials: Table S1 and S2 show the conversion of anisole and product distribution varied with temperature and time, respectively. Figure S1 shows the pore size distribution of the different catalysts. Figure S2 shows the TEM image of Ni/ α -Al₂O₃ catalyst. Figure S3 shows the TG curves of Ni/ α -Al₂O₃ used in the first and the fifth run.

V. ACKNOWLEDGMENTS

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BiFe_{1-x}Nb_xO₃的结构、磁性和光学性质..... 578

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摘要: 用溶胶凝胶法制备了Nb掺杂多铁BiFe_{1-x}Nb_xO₃粉晶样品(0≤x≤0.05), 研究Nb掺杂对样品的结构、磁学和光学性质的影响. 根据XRD图谱和Rietveld精修的结果可知, 所有的样品仍保持R3c相, 但晶格常数a, c, 晶胞体积V和Fe—O—Fe键角发生变化. 适当的Nb掺杂使得样品晶粒尺寸减小, 导致剩余磁化强度的增强, 使得BiFe_{1-x}Nb_xO₃样品的禁带窄化.
关键词: 铁酸铋, 晶体结构, 磁学性质, 带隙

光沉积Ru和RuO₂的锐钛矿TiO₂纳米片的光解水产氧..... 585
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摘要: 采用水热法以HF作为结构调控剂合成了主要暴露(001)面的锐钛矿TiO₂纳米片, 通过光沉积方法分别合成了负载Ru和RuO₂物种的光催化剂. 利用X射线衍射、透射电镜和氢气程序升温还原等分析表征了催化剂的结构性质. 通过光解水产氧反应来研究催化剂的催化性能, 详细考察了Ru含量、负载方式以及氧化和还原处理等因素的影响. 光解水产氧速率的差异证明了Ru物种在不同晶面的电荷-空穴分离效应. 与负载单一助催化剂的Ru/TiO₂和RuO₂/TiO₂样品相比, 活性最优的0.5%Ru-1.0%RuO₂/TiO₂样品由于负载了双助催化剂, 其催化活性得到更大的提高, 证实了在锐钛矿TiO₂上的晶面电荷-空穴分离效应.
关键词: 锐钛矿TiO₂纳米片, 光解水产氧, 晶面, Ru助催化剂, 电荷分离

三甲基镓在Pd(111)表面吸附解离及表面预吸附H和O的影响. 591
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摘要: 利用X-射线光电子能谱(XPS)和程序升温脱附谱(TPD)研究了三甲基镓在Pd(111)表面的吸附和解离行为, 并考察了表面预吸附H和O的影响. 结果表明, 在吸附温度为140 K时, 三甲基镓在Pd(111)上主要为解离吸附, 此时表面物种为Ga(CH₃)_x (x=1, 2, 3)和CH_x物种. 加热将导致Ga的甲基化合物中的Ga—C键发生分步断裂, 在不同温度下产生CH₄和H₂从表面脱附. 同时, XPS结果证实了在275~325 K的温度区间内存在Ga甲基化合物的分子脱附. 退火至更高温度, 表面只观察到积碳和金属Ga物种, 这二者随着温度的继续升高逐渐向体相扩散. 在Pd(111)表面预吸附O和H对上述吸附和解离行为存在显著的影响. 当表面预吸附H时, 脱附产物CH₄和H₂的脱附主要位于315 K, 可归属为一甲基镓的解离脱附. 当表面预吸附O时, 只在258 K观察到CH₄和H₂的脱附峰, 可能来自于Pd-O-Ga(CH₃)₂吸附结构的解离.
关键词: 三甲基镓, Pd(111), 吸附, 解离, XPS, TPD

三元异质结构Ag-Bi₂MoO₆/BiPO₄光催化剂高效降解苯酚红 600
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摘要: 多组分复合体系有利于电荷的有效分离, 减少电子空穴对的复合几率. 通过低温液相法首次合成Ag-Bi₂MoO₆/BiPO₄三元异质结构光催化剂. 利用XRD、SEM、EDX及XPS等技术对样品进行了表征. 结果表明, Ag纳米粒子光照积累在Bi₂MoO₆/BiPO₄的表面, 通过表面等离子共振增加对可见光的吸收, 同时作为电子受体促进了光生电子的转移. Ag、BiPO₄和Bi₂MoO₆形成三元异质结构有效地抑制了光生电子空穴对的复合. Ag-Bi₂MoO₆/BiPO₄表现出优异的光催化性能, 其光催化活性较BiPO₄、Bi₂MoO₆和Bi₂MoO₆/BiPO₄样品有较大提高. 并且对Ag-Bi₂MoO₆/BiPO₄三元异质结构的光催化机制进行了讨论. 光催化过程中反应活性物种捕获实验结果表明h⁺和O₂⁻是主要的活性基团.
关键词: 异质结构, 钼酸铋, 磷酸铋, 银, 光催化, 可见光

含硫磷酸钙纳米颗粒的制备及对铅离子的高效选择性去除..... 607
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摘要: 用一种简易共沉淀法制备了非晶含硫磷酸钙(SCP)材料, 实现硫原子原位引入磷酸钙纳米颗粒中, 并研究了其对Pb(II)的吸附特性和机理. 与羟基磷灰石相比, SCP对Pb(II)的去除性能显著增强, 在10 min内能快速将20 ppm的Pb(II)溶液降低至饮用水标准

下. 由Langmuir吸附等温线模型计算可知, SCP对Pb(II)的最大饱和和吸附量高达1720.57 mg/g, 这个数值远远超过以往所报道的绝大部分吸附剂材料. 在竞争离子Ni(II), Co(II), Zn(II)和Cd(II)共存的条件下, SCP还表现出对Pb(II)的选择性去除. 研究表明, SCP对Pb(II)超高的去除效率和优异的亲和力归因于其可通过溶解沉淀和离子交换反应在其表面形成棒状的羟基磷酸铅晶体, 以及形成沉淀物硫化铅. SCP以其对Pb(II)快速、高效和优异选择性成为在实际铅污染治理中的理想材料.
关键词: 含硫磷酸钙, 铅离子, 选择性去除

Ni/α-Al₂O₃催化剂作用下苯甲醛的加氢脱氧..... 617
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摘要: 通过化学沉淀法制备了以α-Al₂O₃、γ-Al₂O₃、SiO₂、TiO₂和ZrO₂为载体的五种镍基催化剂, 以苯甲醛为模型化合物对催化剂进行活性评价, 考察载体对加氢脱氧催化反应的影响. 实验表明, Ni/α-Al₂O₃对苯甲醛加氢脱氧反应的催化活性最高. 在优化的工况下, 苯甲醛的转化率达到93.25%, 碳氢化合物收率达到90.47%. H₂-TPD测试表明五种镍基催化剂中, Ni/α-Al₂O₃表面拥有更多的活性金属位, 具有更高的加氢催化活性. Ni/α-Al₂O₃催化剂具有优异的可重复使用性能, 反应后催化剂表面的积碳量几乎可以忽略.
关键词: 苯甲醛, 加氢脱氧, Ni/α-Al₂O₃, 碳氢化合物

基于分布参数系统和单粒子模型的锂离子电池工作状态实时监控模型..... 623

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摘要: 提出了一种基于分布参数系统偏微分方程描述和单粒子模型的锂离子电池工作状态实时监控建模方法, 能够实时跟踪锂离子电池阳极的锂离子密度和残差变化, 并通过仿真实验验证了本文模型对锂离子电池工作状态故障报警的有效性和精确性.
关键词: 锂离子电池, 分布参数系统, 单粒子模型, 工作状态监控

基于支持向量机、支持向量回归和分子对接的CYP450 1A2抑制剂的发现研究..... 629

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摘要: 支持向量机, 支持向量回归和分子对接的计算方法已广泛应用于化合物的药理活性计算. 为了提高计算的准确性和可靠性, 拟以细胞色素P450酶1A2为研究载体, 运用建立的联合SVM-SVR-Docking计算模型预测潜在的CYP1A2抑制剂. 其中, 建立的最优SVM定性模型训练集, 内部测试集和外部测试集的准确率分别为99.432%, 97.727%和91.667%. 最优SVR定量模型训练集和测试集的R²和MSE分别为0.763, 0.013和0.753, 0.056. 实验表明两个模型具有较高的准确性和可靠性. 通过对SVM和SVR模型结果的分析, 发现连接性指数、分子构成描述符和官能团数目等分子描述符可能与CYP1A2抑制剂的辨识和活性预测密切相关. 随后利用分子对接技术分析化合物与CYP1A2的结合构象及相互作用的稳定性. 形成氢键相互作用的关键氨基酸包括THR124, ASP320; 形成疏水相互作用的关键氨基酸包括ALA317和GLY316. 所获得模型可用于天然产物化学成分中CYP1A2潜在抑制剂的活性计算及其介导的药物-药物相互作用预测提供理论指导, 也为合理联合用药提供一定参考. 共获得20个对CYP1A2具有潜在抑制活性的化合物. 部分结果与文献结果相互印证, 进一步说明了模型的准确性和联合计算策略的可靠性.
关键词: 支持向量机, 支持向量回归, 分子对接, CYP1A2抑制剂

生物油制备生物质氢气和生物质燃料..... 635

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摘要: 研究了一种以生物质裂解油为原料制备氢气和生物燃料的催化转化过程. 该过程包括生物油催化裂解制氢气和生物合成气, 合成气的调变, 烯烃聚合和费脱合成耦合制备生物燃料. 在优化反应条件下, 氢气产率达到120.9 g H₂/(kg bio-oil), 烯烃聚合-费脱耦合反应形成的生物燃料产率达到526.1 g/(kg bio-syngas). 基于产物分析和催化剂特性表征, 探讨了生物燃料合成过程中的反应路径和化学反应过程.
关键词: 生物油, 生物质氢气, 烯烃聚合, 费脱合成, 生物燃料