



Cite this: *RSC Adv.*, 2019, 9, 21175

Selective hydrodeoxygenation of *p*-cresol as a model for coal tar distillate on Ni–M/SiO₂ (M = Ce, Co, Sn, Fe) bimetallic catalysts

Liuyi Pan,^{ab} Yulong He,^a Menglong Niu,^c Yong Dan^a and Wenhong Li^{*a}

Ni–M/SiO₂ with different binary metals M (M = Ce, Co, Sn, Fe) prepared by an incipient impregnation method was used in the hydrodeoxygenation (HDO) of low-temperature coal tar distillate, which is rich in phenolic compounds. *p*-Cresol, as a model compound of the distillate, was used to evaluate the activity and selectivity of BTX products on the series of reduced Ni–M/SiO₂ catalysts in a fixed bed reactor. The properties of the catalysts were characterized by N₂ adsorption–desorption, ICP-AES, XRD, H₂-TPR, and XPS. Benzene and toluene as the direct deoxygenation (DDO) products and cyclohexane and methylcyclohexane as the hydrogenolysis (HYD) products were detected to evaluate the selectivity of the path in the deoxygenation process. In this series of catalysts, the order of reactivity was Ni–Ce > Ni–Sn > Ni–Co > Ni–Fe > monometallic Ni. Meanwhile, the addition of Ce and Co loaded in the Lewis acid sites of the catalyst affected the electron distribution of nickel atom and its atomic arrangement on the surface of the carrier. Compared to monometallic Ni, the DDO path become dominant on Ni–Ce and Ni–Co and the selectivity for BTX products increased from 58.8% to 77.4% and 71.1%, respectively. The binary metal Sn, unlike the former two metals, formed a Ni₃Sn crystal form with Ni, which resulted in significant enhancement of the HYD path while obviously increasing the reactivity.

Received 13th April 2019

Accepted 31st May 2019

DOI: 10.1039/c9ra02791b

rsc.li/rsc-advances

1 Introduction

The production of fuels such as gasoline and diesel by the hydrogenation of coal tar has been frequently reported and has been industrialized.^{1–3} In low-temperature coal tar, more than 90% of the distillate below 230 °C consists of phenolic compounds,^{4,5} mainly monocyclic phenols, which account for more than 60% of the total phenols.⁶ In the process of hydro-treating, these compounds often lead to the formation of large amounts of naphthenes and water; thus, it is not only difficult to obtain high-octane fuel, but the catalyst life is adversely affected.⁷ Moreover, if the intent is to use the hydrogenation product downstream to produce BTX, a dehydrogenation reforming process is also necessary.⁸ Therefore, the selective hydrodeoxygenation of monocyclic phenolic compounds enriched in coal tar can improve the quality of fuel oil and can also benefit the production of BTX products.^{9,10}

Hydrodeoxygenation of monocyclic phenolic compounds usually involves two basic reaction pathways. One involves breaking C–O bonds to obtain aromatics, called direct

deoxygenation (DDO); the other proceeds *via* prehydrogenation of the aromatic rings, leading to alicyclic hydrogenation dehydrogenation (HYD).^{11,12} The traditional catalyst used in hydro-treating coal tar is generally a sulfided catalyst which contains γ -Al₂O₃ as a carrier and is loaded with active metals such as Ni, W, Co, and Mo.^{13–15} The sulfurized CoMo/ γ -Al₂O₃ catalysts used in studies on phenol hydrodeoxygenation possess both high reactivity and high selectivity but are deactivated easily.^{16,17} Senol¹⁸ studied HDO experiments of phenol on sulfided CoMo/Al₂O₃ and NiMo/Al₂O₃ catalysts. The results showed that the main hydrodeoxygenation product of phenol on the sulfided Ni–Mo catalyst was alicyclics, while the product on Co–Mo was mainly benzene; also, the catalyst deactivation was mainly due to coking and loss of sulfur atoms. Viljava¹⁹ observed obvious coke deactivation of CoMo/Al₂O₃ catalyst during the HDO of guaiacol, especially at high reaction temperatures.

Supported noble metal catalysts are also common in HDO.^{17,20,21} Gutierrez²² studied the HDO of *p*-cresol on zirconia-supported mono- and bimetallic noble metal (Rh, Pd, Pt) catalysts. They found that the noble metal catalysts simultaneously showed higher catalytic activity and lower carbon deposition compared with the conventional sulfided CoMo catalyst. As new hydroprocessing catalysts, the application of transition metal phosphides in HDO has attracted great attention in recent years. Especially, metal–nickel phosphide catalysts show superior reactivity and sulfur resistance.^{23–26} Zhao²⁷ prepared a series of metal phosphide catalysts, including Ni₂P/SiO₂, Fe₂P/SiO₂,

^aSchool of Chemical Engineering, Northwest University, Xi'an, Shaanxi 710069, P. R. China. E-mail: liwenhong@mwu.edu.cn

^bCollege of Chemistry & Chemical Engineering, Baoji University of Arts and Sciences, Baoji 721013, Shaanxi, China

^cCollege of Chemistry & Chemical Engineering, Xi'an Shiyou University, Xi'an, Shaanxi Province, 710065, P. R. China



MoP/SiO₂, Co₂P/SiO₂ and WP/SiO₂, for hydrodeoxygenation of guaiacol. The results showed that Ni₂P/SiO₂ has the highest reactivity, and the products were mainly phenol and benzene. In contrast, commercial CoMoS/Al₂O₃ showed little activity and was deactivated quickly. Metal-reduced catalysts are widely used in HDO processes, especially those loaded with nickel.^{28,29} In general, a small amount of a binary metal, such as La,³⁰ Mo,³¹ Cu,^{32,33} or Fe,³³ is considered to improve stability and reactivity.

Various types of catalysts have been studied in the hydrodeoxygenation reactions of phenol, guaiacol, anisole *et al.* as bio-oil models. However, phenolic compounds in coal tar have different species and distributions than those in bio-oil; these compounds mainly consist of methylphenol, dimethylphenol and higher phenols (such as naphthol and indophenol). Few studies have been reported on the selective hydrodeoxygenation of these phenolic compounds to obtain BTX products *via* the DDO pathway. Especially, research about the effects of different Ni–M binary metals on the hydrodeoxygenation pathway of alkyl phenols is rare. In the present study, a series of Ni–M/SiO₂ bimetallic catalysts (M = Ce, Co, Sn, Fe) were prepared by an incipient-wetness impregnation method. *p*-Cresol, a monocyclic phenolic compound with high content in the fraction of coal tar, was selected as a model compound to evaluate the deoxidation and transalkylation during the hydrogenation process. The fixed bed hydrogenation reaction was carried out in the temperature range of 330 °C to 390 °C to evaluate the activity and product selectivity of the catalysts, and the catalysts were also characterized by BET, XRD, H₂-TPR, NH₃-TPD and XPS.

2 Experimental

2.1 Materials and chemicals

p-Cresol (≥98.0%), Ni(NO₃)₂·6H₂O (≥98.0%), and SnC₂O₄ (≥98.5%) were obtained from Aladdin Chemical Reagent Co. Ltd., while Ce(NO₃)₃·6H₂O (≥99.0%), Fe(NO₃)₃·9H₂O (≥98.5%), and Co(NO₃)₂·6H₂O (≥98.5%) were supplied by Sinopharm Chemical Reagent Co. Ltd. SiO₂ was purchased from a local manufactory.

2.2 Preparation of catalysts

Five catalysts, Ni/SiO₂, Ni–Ce/SiO₂, Ni–Co/SiO₂, Ni–Sn/SiO₂, and Ni–Fe/SiO₂, were prepared through an incipient wetness impregnation method in which the Ni loading was kept constant at 20 wt% (calculated by carrier quality) and the Ni/M molar ratio was maintained at 10. First, 15 g SiO₂ were dried at 120 °C for 12 h. Second, 14.86 g Ni(NO₃)₂·6H₂O with different bimetallic nitrates (2.21 g Ce(NO₃)₃, 1.48 g Co(NO₃)₂·6H₂O, 1.05 g SnC₂O₄, or 1.23 g Fe(NO₃)₃·9H₂O; in addition, SnC₂O₄ should be hydrolyzed first with hydrochloric acid) were dissolved in 15 mL deionized water (the volume of dissolved water was calculated by measuring the water absorption of the carrier) in a beaker and treated by ultrasonic dissolution for 20 min. Then, the metal salt solution was slowly added dropwise to the carrier with magnetic stirring for 12 h at 45 °C. Third, the precursors were removed from the immersion solution by drying at 120 °C for 12 h and finally calcined at 550 °C for 4 h.

The catalyst precursors were subjected to reduction treatment before being used in the reaction. The reduction conditions are as follows: H₂/N₂ (10% H₂) mixed gas with a flow rate of 200 mL min^{−1}; increase temperature from 30 °C to 350 °C at 5 °C min^{−1}, maintain for 60 min at 350 °C, then increase again from 350 °C to 550 °C at 2 °C min^{−1} and maintain for another 4 h.

2.3 Characterization of catalysts

N₂ adsorption–desorption isotherms of the catalysts were measured by a Quantachrome SI instrument at −196 °C. Before these measurements, the catalyst samples were degassed under vacuum at 350 °C for 3 h. The Brunauer–Emmett–Teller (BET) equation was used to calculate the specific surface area (*S*_{BET}). The average pore diameter was calculated using the desorption branch of the isotherm according to the BJH method.

Inductively coupled plasma-atomic emission spectroscopy (ICP-AES) measurements were carried out on an Agilent 725 instrument. Prior to these measurements, the samples were first dissolved in mixed acid with concentrated sulfuric acid and concentrated nitric acid in a certain proportion.

CO pulse chemisorption studies were carried out in an AutoChem II 2920 instrument (Micromeritics). 80 mg sample was first reduced in a H₂ flow (50 mL min^{−1}) at 550 °C for 1 h, then cooled with a He flow to 25 °C. After the TCD signal was stable, CO (50 μL) was pulsed in until the signal no longer changed; subsequently, the CO uptake was calculated.

Powder X-ray diffraction (XRD) was performed on a Bruker D8 Avance instrument with Cu–Kα radiation (λ = 1.5409 Å). The X-ray tube was operated at 40 kV and 40 mA. The XRD patterns of the samples were recorded in the range of 10° ≤ 2θ ≤ 80° with a scanning step size of 5°.

H₂ temperature-programmed reduction (H₂-TPR) was measured on a self-built test instrument containing a U-tube reactor, a heating jacket and a thermal conductivity detector (TCD). The catalyst sample (80 mg) was maintained in the reactor, then dried and degassed under N₂ flow (50 mL min^{−1}) at 200 °C for 2 h. Reduction was conducted at a heating rate of 10 °C min^{−1} from 200 °C to 800 °C in a 10 vol% H₂/N₂ flow (50 mL min^{−1}).

Temperature-programmed desorption of ammonia (NH₃-TPD) was carried out on the AutoChem II 2920 (Micromeritics) to evaluate the total acidities of the catalysts. The catalysts were reduced under flowing H₂ (50 mL min^{−1}) at 450 °C for 2 h. After cooling and cleaning with He (50 mL min^{−1}) and adsorption of NH₃ at 100 °C, the NH₃-TPD was performed between 100 °C and 800 °C with a heating rate of 10 °C min^{−1} using a He flow.

X-ray photoelectron spectroscopy (XPS) was carried out using a ThermoFisher ESCALAB 250Xi with Al Kα radiation (1486.6 eV). The binding energies were referenced to C 1s at 284.8 eV.

2.4 HDO tests

HDO reactions were performed on a fixed-bed reactor (ID 17 mm, length 1200 mm) with separate gas and liquid feed systems and a thermocouple to monitor the reaction temperature. Before the hydrogenation reaction, 30 mL catalyst was placed in



the reactor and reduced at 550 °C for 2 h in a H₂ flow (100 mL min⁻¹); then, the reactor was cooled to reaction temperature in H₂ atmosphere. The liquid products were analyzed by an Agilent gas chromatograph (9790 A) equipped with an HP-INNOWax polar column (30 m × 0.32 mm × 0.5 mm) and a flame ionization detector.

During the experiment, the main hydrogenation products were quantitatively analyzed by the external standard method; products with very low contents, such as biphenyls and olefin, were not calculated.

The conversion of *p*-cresol and the product selectivity were defined as follows:

$$\text{Conversion}[\%] = \frac{N(\text{p-cresol})_{\text{in}} - N(\text{p-cresol})_{\text{out}}}{N(\text{p-cresol})_{\text{in}}} \times 100$$

$$\text{Selectivity}[\%] = \frac{N_i}{N(\text{p-cresol})_{\text{in}} - N(\text{p-cresol})_{\text{out}}} \times 100$$

BTX products are the target products for this research. The main BTX products obtained by hydrogenation of *p*-cresol are benzene and cyclohexane. The content of xylene in the product is extremely low; thus, it was neglected when calculating selectivity_{BTX}.

The BTX selectivity was defined as follows:

$$\text{Selectivity}_{\text{BTX}} = \frac{N(\text{benzene}) + N(\text{toluene})}{N(\text{p-cresol})_{\text{in}} - N(\text{p-cresol})_{\text{out}}}$$

3 Results and discussion

3.1 Characterization of the catalysts

3.1.1 BET. The BET surface areas, pore volumes and pore sizes of Ni-M/SiO₂ and the carrier measured by N₂ physisorption are listed in Table 1. Compared with the SiO₂ carrier, the surface area of Ni/SiO₂ slightly decreased after loading the monometallic nickel. After the addition of the binary metal M, the surface area and pore volume of each Ni-M catalyst further decreased, which was especially obvious for Ni-Co and Ni-Fe; this probably occurred because the surface and the pores of SiO₂ were blocked by the loaded metal elements.^{34,35} In this series of catalysts, the loadings of nickel and binary metals were

similar in molar composition and basically consistent with theoretical values. This indicates that the preparation process of the catalysts was stable throughout the study.

3.1.2 XRD. The XRD pattern of each catalyst is shown in Fig. 1. A broad peak with 2θ between 15° and 30° was attributed to the characteristic diffraction peak of amorphous silica. In agreement with previous observations,^{36,37} the obvious peaks at 2θ = 44.5° (111), 51.8° (200), and 76.4° (220) were attributed to metallic nickel (JCPDS 87-0712), indicating that the main active sites of the catalysts are all reduced metal nickel. New diffraction peaks at 2θ of 28.5° and 56.3°, indexed to the (111) and (311) diffraction planes of CeO₂ crystal phase (JCPDS 81-0792), were observed in the pattern for Ni-Ce/SiO₂. However, this reveals that Ce added as a binary metal exhibited a partially oxidized state; that is, Ce was not completely reduced to the metallic state at this reduction temperature. In contrast to Ni-Ce, in the spectrum of Ni-Co, metallic cobalt diffraction peaks appeared at 2θ = 42.3° and 44.8° (JCPDS 05-0727); this indicates that the surface of the catalyst contained dispersed cobalt that was reduced to a metallic state. Consistent with the literature,³⁸ new diffraction peaks that are different from the metallic and oxidized states of nickel and tin were observed at 2θ = 33.8°, 39.3° and 42.5° in the spectrum of Ni-Sn; this exhibits the formation of a new Ni₃Sn alloy phase. A slight shift to a lower angle compared to Ni-M suggests that the addition of Sn may cause lattice distortion after entering the lattice. Surprisingly, the diffraction peaks of iron were not observed, whether in the metallic or oxidized state, and only a slight shift to a lower angle was observed in the diffraction peak of nickel. Nie *et al.*³³ concluded that the shift of the nickel diffraction peak in the XRD pattern indicated the formation of Ni-Fe solid solution after the addition of the binary metal Fe.

The average particle diameters of the Ni-M/SiO₂ catalysts, as shown in Table 1, were calculated by Scherrer's formula through selecting the diffraction peak at 2θ = 51.8°; meanwhile, the dispersions of the active metal were calculated from the relationship between the particle diameter *d* and the degree of dispersion *D* (*d* ≈ 0.9/*D*). Simultaneously, CO chemisorption was used to characterize the particle sizes and dispersions, which are listed in Table 1. The particle sizes of the catalysts, measured by both methods, are distributed in the range of 10 to 18 nm; this indicates good consistency.³⁷ After the addition of the binary metal, compared with Ni/SiO₂, the particle size of Ni-

Table 1 Physical properties and CO uptake of the Ni-M/SiO₂ (M = Ce, Co, Sn and Fe) catalysts

	Ni ^a wt%	M ^a wt%	BET surface area ^b (m ² g ⁻¹)	Pore volume	Pore size	Particle size ^c nm	Dispersion ^d (%)	CO uptake mmol g ⁻¹	Particle size ^e nm
SiO ₂			438	0.85	6.88				
Ni	15.7	—	402	0.77	6.28	19.8	4.5	0.12	18.3
Ni-Ce	14.8	3.75	384	0.70	6.12	11.4	7.9	0.17	10.5
Ni-Co	14.6	1.48	371	0.69	6.01	12.2	7.4	0.16	11.5
Ni-Sn	14.3	3.10	389	0.72	6.04	14.6	5.5	0.18	9.9
Ni-Fe	15.0	1.42	360	0.71	6.14	13.7	6.6	0.13	15.1

^a Determined by ICP-AES. ^b BET surface areas, pore volumes, and pore sizes determined by N₂ physisorption. ^c Particle size calculated by the Scherrer formula. ^d Dispersion (*D*) calculated with particle size (*d*) by the equation *D* ≈ 0.9/*d*. ^e Calculated by CO pulse adsorption.



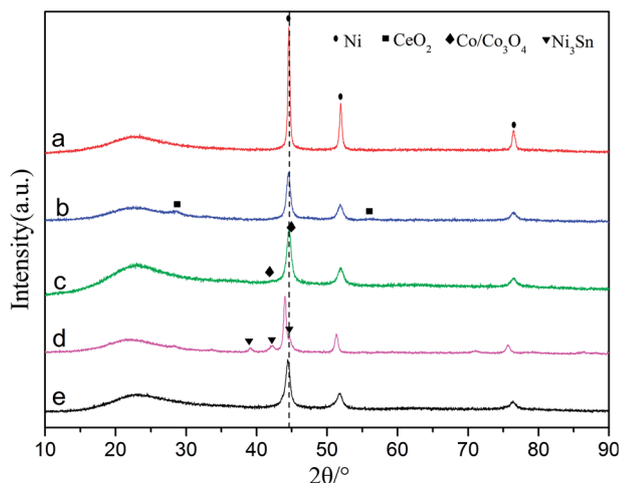


Fig. 1 X-ray diffraction patterns for (a) Ni/SiO₂, (b) Ni-Ce/SiO₂, (c) Ni-Co/SiO₂, (d) Ni-Sn/SiO₂ and (e) Ni-Fe/SiO₂.

M decreased, indicating that the dispersion of the active metal improved.²⁷ Based on the premise that each active site adsorbs a CO molecule, the order of dispersion was Ni-Ce > Ni-Co > Ni-Fe > Ni-Sn > Ni, suggesting that the addition of Ce results in higher reactivity due to its effects on the dispersibility.

3.1.3 H₂-TPD. The reduction properties of the catalysts were characterized by temperature programmed reduction experiments with H₂ (H₂-TPR), and the TPR profiles of the monometallic Ni/SiO₂ and the bimetallic Ni-M/SiO₂ catalysts are compared in Fig. 2. As in previous literature,^{38,39} an obvious peak observed in the profile of Ni/SiO₂ at around 390 °C is attributed to the reduction of NiO to metallic nickel, while a smaller shoulder peak appears on the right side due to the formation of poorly reducible nickel silicates. The temperature of the main reduction peak of each binary metal catalyst, compared to mono Ni/SiO₂, changed as follows: Ni-Ce (388 °C), Ni-Co (395 °C), Ni-Sn (450 °C), and Ni-Fe (425 °C). As can be seen from the pattern of Ni-Ce/SiO₂, the introduction of Ce caused a slight decrease in the reduction temperature of NiO,

indicating improvement of the reductivity. Zhang *et al.*⁴⁰ attributed this lower reduction temperature to the formation of Ni-Ce-O species and the lattice distortion of ceria. Moreover, a broad peak ranging from 500 °C to 600 °C was attributed to the hydrogen consumption peak of Ce⁴⁺ to Ce³⁺; this suggests that metal Ce has a higher reduction temperature and cannot be completely reduced to metallic Ce at 550 °C.⁴¹ This result is consistent with the XRD analysis. An observed difference is that in the profile of Ni-Co, a shoulder peak appeared at 450 °C and a broad peak appeared at 550 °C; these were attributed to the reduction of Co⁴⁺ to Co²⁺ and the reduction of Co²⁺ to metallic Co, respectively.⁴² Obviously, in the profile of Ni-Sn/SiO₂, the peak temperature range from 450 °C to 600 °C is higher than the others; this proves the formation of Ni₃Sn crystal phase and the presence of strong interactions between the Ni-Sn biphasic alloy and the carrier.⁴³ In addition, after the addition of the binary metal Fe to Ni/SiO₂, the TPR spectrum contained no peak above 500 °C; only the main peak was observed, and it moved to a higher temperature. Sitthisa *et al.*⁴⁴ reported that under a consistent load of Ni, hydrogen consumption increased as the loading of Fe increased in H₂-TPR; however, no peak related to the reduction of iron oxide to α-Fe was observed, but a shift to higher temperature was observed. This indicates the reduction of iron oxide and close interactions between Ni and Fe.

3.1.4 NH₃-TPD. The NH₃-TPD profile of each catalyst, shown in Fig. 3, shows an obvious peak of weak acid sites (at about 200 °C) and a weak peak of medium-strength acid sites (at about 450 °C). Nickel cations can usually bring Lewis acid sites to a carrier.^{24,45} The addition of a binary metal may have two effects on the acidity of the monometallic nickel catalyst.

On the one hand, the acidity of the Ni-M catalyst increased in comparison with that of Ni/SiO₂ with the addition of the binary metal, as can be seen from the acid quantitative results in Table 2.

This indicates that the reduced or partially oxidized binary metal M increased the number of new acid sites, especially cerium ions and cobalt ions, which were not completely

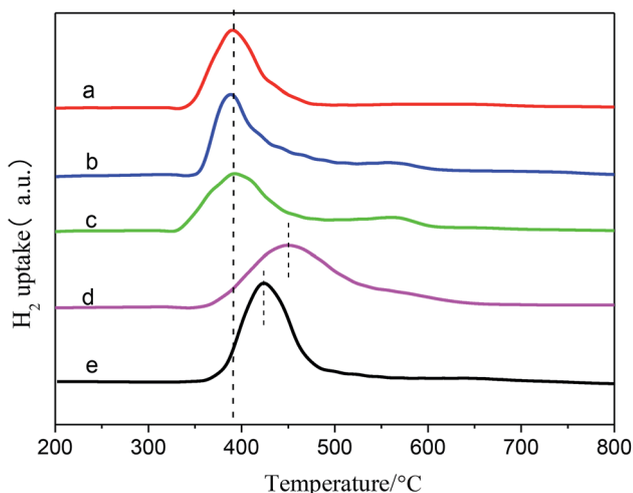


Fig. 2 H₂-TPR profiles of (a) Ni/SiO₂, (b) Ni-Ce/SiO₂, (c) Ni-Co/SiO₂, (d) Ni-Sn/SiO₂, (e) Ni-Fe/SiO₂.

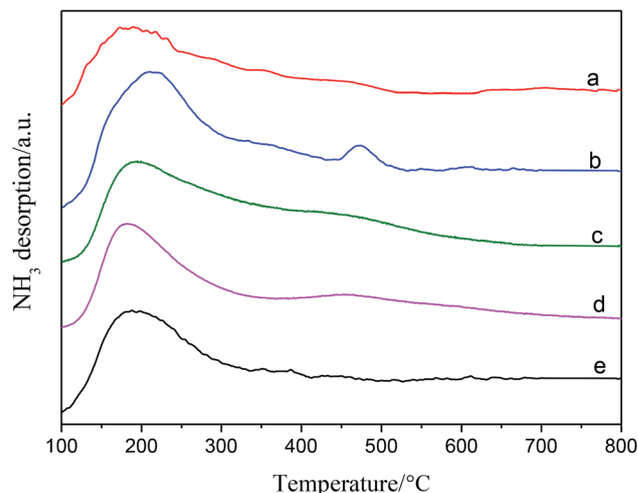


Fig. 3 NH₃-TPD profiles of (a) Ni/SiO₂, (b) Ni-Ce/SiO₂, (c) Ni-Co/SiO₂, (d) Ni-Sn/SiO₂, (e) Ni-Fe/SiO₂.



Table 2 NH₃-TPD data of the catalysts

Catalyst	Relative NH ₃ desorption amount ^a
Ni-Ce	1.52
Ni-Co	1.40
Ni-Sn	1.24
Ni-Fe	1.10

^a The NH₃ desorption amount of Ni/SiO₂ was specified as 1.00.

reduced. On the other hand, the addition of a binary metal resulted in an improvement in the dispersion of the active metal. The aggregation and mutual coverage of the active metal Ni were prevented; therefore, more Lewis acid sites could be formed.⁴⁶

Therefore, the total acid sites of the catalysts increased in the order of Ni-Ce/SiO₂ > Ni-Co/SiO₂ > Ni-Sn/SiO₂ > Ni-Fe/SiO₂ > monometallic Ni/SiO₂, which is similar to the order of metal dispersion.

3.1.5 XPS. The Ni 2p XPS lines profiles for different binary metal catalysts, shown in Fig. 4, were used to characterize the electronic properties. In the Ni 2p_{3/2} XPS profile of Ni/SiO₂, the

main peak with a binding energy (BE) value of 853.0 eV and an associated satellite peak located 6 eV higher than the main peak were attributed to metallic Ni. Additionally, Ni 2p_{1/2} at 871.0 eV is attributed to Ni⁰.^{47,48}

In comparison with monometallic Ni, shifts in BE towards lower values of about 0.1 eV to 0.5 eV were observed, from 853.0 eV (mono Ni) to 852.5 eV (Ce), 852.7 eV (Co), 852.9 eV (Sn), and 852.8 eV (Fe); this was in accordance with the electronegativity values of Ni (1.9), Ce (1.1), Co (1.8), Sn (1.7) and Fe (1.9). These shifts were mainly attributed to electron transfer from the binary metal M to nickel, which caused an increase in the electron density of the nickel atoms. In addition, the occurrence of electron transfer confirmed the interaction between Ni and the binary metal M.

In addition, the distribution of the valence states and atomic ratios on the surface are listed in Table 3. Ni²⁺ species with a small intensity peak observed at 854.8 eV (Ni/SiO₂) showed that the samples may be partially oxidized when exposed to air.⁴⁹ However, the distributions of Ni²⁺ in Ni-Ce (3.8%), Ni-Co (1.1%), Ni-Sn (0.7%) and Ni-Fe (2.1%) are lower than that observed for monometallic Ni (5.8%); this indicates that due to the interaction of Ni and the binary metal M, the phenomenon

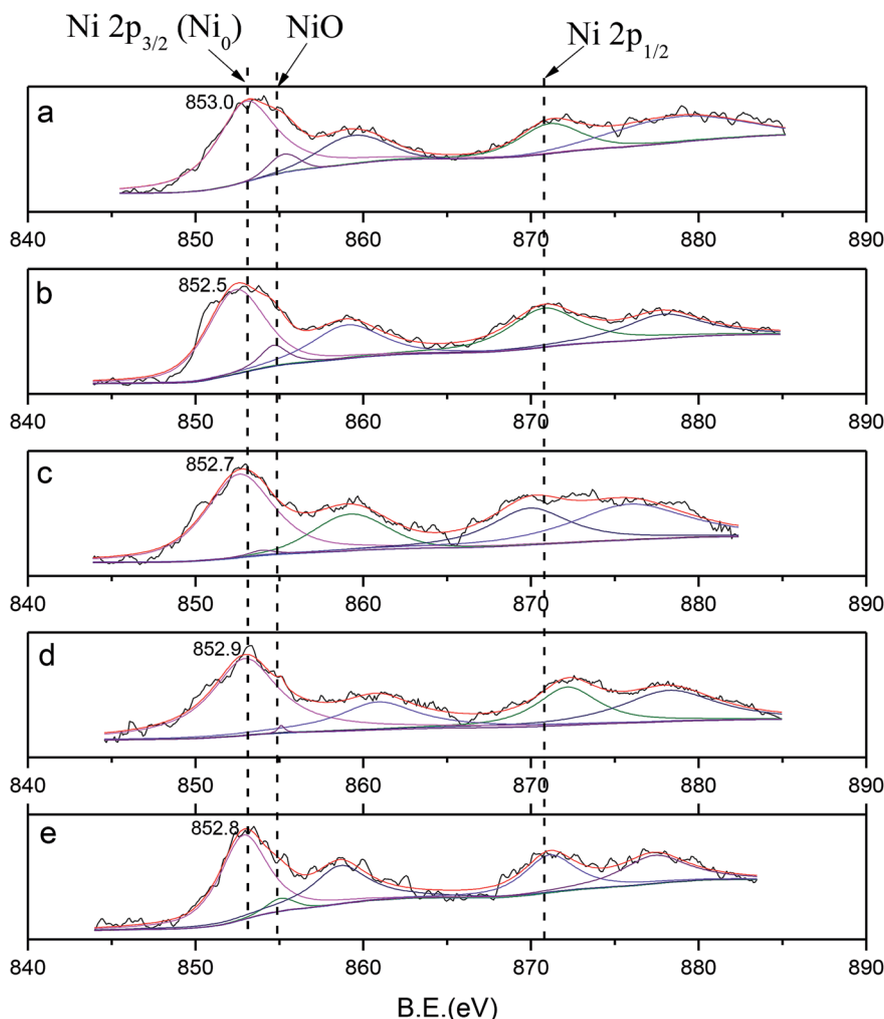


Fig. 4 XPS profiles of (a) Ni/SiO₂, (b) Ni-Ce/SiO₂, (c) Ni-Co/SiO₂, (d) Ni-Sn/SiO₂, (e) Ni-Fe/SiO₂.



Table 3 Distribution of different valence states of Ni–M and atomic ratios derived from XPS

Catalyst	Distribution of valence states (%)				Atomic ratio			
	Ni ⁰ 2p _{3/2}		Ni ⁰ 2p _{1/2}		Ni ²⁺	Ni/Si	M/Si	Ni/M
	Main	Sat.	Main	Sat.				
Ni	43.5	13.4	13.8	23.5	5.8	0.15		
Ni–Ce	37.7	18.7	23.5	16.3	3.8	0.06	0.04	1.5
Ni–Co	36.5	14.4	19.9	28.0	1.1	0.07	0.06	1.2
Ni–Sn	41.9	17.4	19.7	20.4	0.7	0.04	0.03	2
Ni–Fe	37.1	22.1	19.3	18.3	2.1	0.10	0.03	3.3

of Ni oxidation was prevented. The effect of Sn is especially obvious. In addition, compared with monometallic Ni, the Ni/Si ratios decreased after addition of M; among the metals, Sn and Ce showed the most obvious decreases. The decrease in the surface content of Ni reflects the increase in dispersion. This is in agreement with the CO uptake results and also confirms the superiority of the catalytic activity of Ni–Sn. In addition, Ni/M decreased from the theoretical 10 to 1 to 3.5, indicating that the dispersibility of Ni is better than that of M and that M atom and Ni are co-distributed on the surface.

3.2 HDO tests

3.2.1 HDO activity and selectivity of *p*-cresol on Ni–M/SiO₂.

This series of Ni–M/SiO₂ catalysts was tested in the HDO of *p*-cresol, which is considered to be a model of phenols in coal tar. The reaction conditions were as follows: the reaction temperature, pressure, LHSV, and H₂-liquid volume ratio were 370 °C, 2 MPa, 0.5 h⁻¹ and 400 : 1, respectively. The distribution of the main products is shown in Fig. 5. The conversion of *p*-cresol, the selectivity of each product and the selectivity of the target products BTX are listed in Table 4. Consistently, DDO and HYD are basically parallel processes, and *p*-cresol is hydroconverted to toluene and methylcyclohexane through these two paths, respectively.³⁴ However, based on the detection of the products, it is believed that there is a third reaction path in HDO for the

hydrogenation of *p*-cresol, namely a demethylation reaction, which ultimately results in the production of cyclohexane and benzene after the HDO of phenol. However, the xylene in the product is presumably derived from the transalkylation reaction of toluene and benzene. The main cause of the weak dealkylation and alkylation phenomena is presumed to be the Lewis acid sites provided by the oxide of the unreduced binary metal M, as characterized by NH₃-TPD and TPR.⁵⁰

The conversion over these catalysts of HDO follows the order: Ni–Ce > Ni–Sn > Ni–Co > Ni–Fe > monometallic Ni. However, the TOFs calculated and listed in Table 4 are inconsistent with the conversion.

According to the characterization analysis of XPS, the addition of binary metal M, especially Ce and Co, resulted in a decrease in the number of d-bands of Ni atom due to the electron transfer from M to Ni; this eventually led to decreases in the intrinsic reaction rates of the catalysts. However, the addition of M improved the dispersion of the active metal; therefore, the specific activity of Ni–M/SiO₂ reflected in the conversion was still higher than that of Ni/SiO₂.⁵¹

In order to evaluate the selectivity of the different reaction pathways in the HDO of *p*-cresol, $S_{\text{BTX}}/S_{\text{cycloparaffin}}$ was defined. The product distributions on Ni–Ce and Ni–Co were very different from that on the monometallic Ni catalyst. In Ni–Ce and Ni–Co, the content of toluene as the main product of the DDO path increased observably; also, the selectivity of toluene, compared to that of the monometallic Ni catalyst, increased from 63.1% to 77.8 and 71.5%, respectively. Moreover, $S_{\text{BTX}}/S_{\text{cycloparaffin}}$ increased from 2.5 to 5.2 and 4.1, respectively. Interestingly, the DDO path had a dominant advantage on Ni–Co, although the activity of the catalyst did not improve greatly. The performance of Ni–Sn differs from the above two. Although the activity of Ni–Sn was significantly improved, $S_{\text{BTX}}/S_{\text{cycloparaffin}}$ decreased sharply from 2.5 to 1.4; meanwhile, the selectivity of BTX product decreased from 66.2% (Ni) to 56.2%. This indicates that Ni–Sn promotes the aromatic hydrogenation of *p*-cresol; that is, the HYD path is enhanced, which ultimately leads to difficulties in obtaining BTX products. Unfortunately,

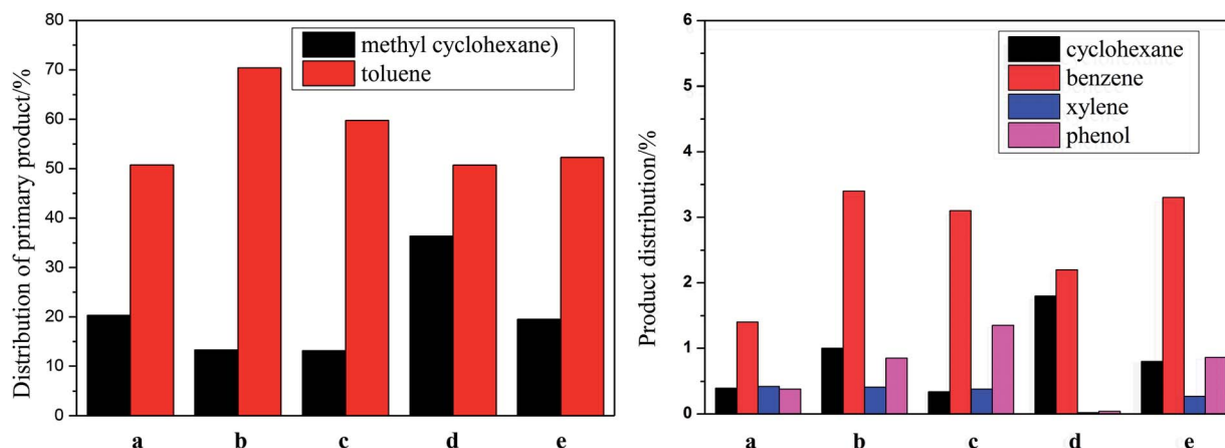


Fig. 5 Product distributions of *p*-cresol over different Ni–M/SiO₂ catalysts. (a) Ni/SiO₂; (b) Ni–Ce/SiO₂; (c) Ni–Co/SiO₂; (d) Ni–Sn/SiO₂; (e) Ni–Fe/SiO₂.



Table 4 Product selectivity comparison on different bimetallic catalysts

	Ni/SiO ₂	Ni-Ce/SiO ₂	Ni-Co/SiO ₂	Ni-Sn/SiO ₂	Ni-Fe/SiO ₂	
Conversion ^a /%	80.5	95.4	88.5	94.1	82.5	
TOF × 10 ⁻³ /S ⁻¹	8.8	6.0	6.3	7.7	7.0	
Selectivity/%						
	Cyclohexane	0.7	1.0	0.4	1.9	1.0
	Benzene	3.0	3.6	3.5	2.3	4.0
	Methyl cyclohexane	25.2	13.9	17.1	38.7	23.6
	Toluene	63.1	73.8	67.6	53.9	63.4
S _{BTX}	66.2	77.8	71.5	56.2	67.7	
S _{BTX} /S _{cycloparaffin}	2.5	5.2	4.1	1.4	2.7	

^a Each catalyst was tested in three parallel experiments with an error less than 2%.

the addition of Fe did not show good performance; the selectivity of the BTX product increased only slightly.

3.2.2 HDO process on Ni-Ce/SiO₂. Because Ni-Ce/SiO₂ showed good reactivity in the HDO of *p*-cresol and high selectivity of BTX products, it was chosen to analyze the effects of reaction temperature on conversion and product distribution. In the reaction temperature range from 330 °C to 390 °C, the amounts of products, conversion and selectivity of BTX with fresh Ni-Ce/SiO₂ are exhibited in Fig. 6.

As the temperature increases, the amount of *p*-cresol decreases from 21.9% to 3.4%; this indicates that the conversion increased from 78.1% to 96.6%. The amount of toluene obviously increased, reaching the maximum value of 74.7% at 390 °C. In contrast, the amount of methyl cyclohexane decreased gradually from 14.2% to 10.8% with increasing reaction temperature from 330 °C to 390 °C. In other words, the process of HDO of *p*-cresol, from which toluene is obtained by the cleavage of C_{aromatic}-O bonds, was enhanced with increasing reaction temperature.⁵² Meanwhile, the process of HDO of *p*-cresol to the cycloalkane through the HYD path was prevented. Similarly, the amounts of benzene and cyclohexane were similar to those of toluene and methylcyclohexane, respectively. It is possible that the high reaction temperature not only promoted the cleavage of C_{aromatic}-O bonds to improve

the selectivity of the DDO pathway, but also enhanced the dealkylation reaction. In addition, the amounts of xylene and phenol, which were lower in the product distribution, both increased initially at temperatures below 350 °C and then decreased as the reaction temperature continued to rise. In summary, the selectivity of the BTX product eventually increased from 74.8% to 81.2% as the reaction temperature increased.

Fig. 7 shows the amounts of *p*-cresol and HDO products as well as the conversion and selectivity of BTX, with LHSV values of 0.25 h⁻¹ to 1.0 h⁻¹ at 370 °C.

As LHSV increased, the content of *p*-cresol increased from 2.7% to 16.9%, indicating that the conversion decreased from 97.3% to 83.1%; this is mainly because a higher LHSV results in shorter contact times of the reactants with the catalyst sites.

The amounts of cyclohexane and methylcyclohexane decreased respectively from 2.7% and 21.2% to 0.6% and 9.9% with increasing LHSV; this also indicates that the decrease in residence time decreased the depth of hydrogenation, resulting in the suppression of aromatic saturation. The amounts of benzene and toluene, as the main BTX products, showed changes of an initial increase followed by a decrease with increasing LHSV, reaching maximum values of 3.4% and 70.4% at 0.5 h⁻¹, respectively. It can be speculated that the decreased

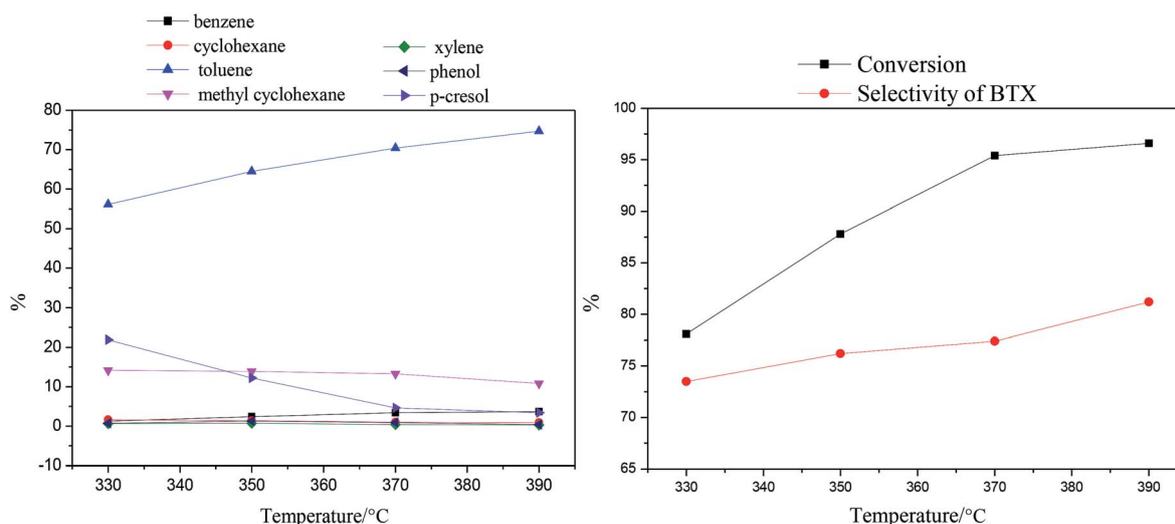


Fig. 6 Effects of reaction temperature on the products and *p*-cresol content via HDO (LHSV = 0.5 h⁻¹).



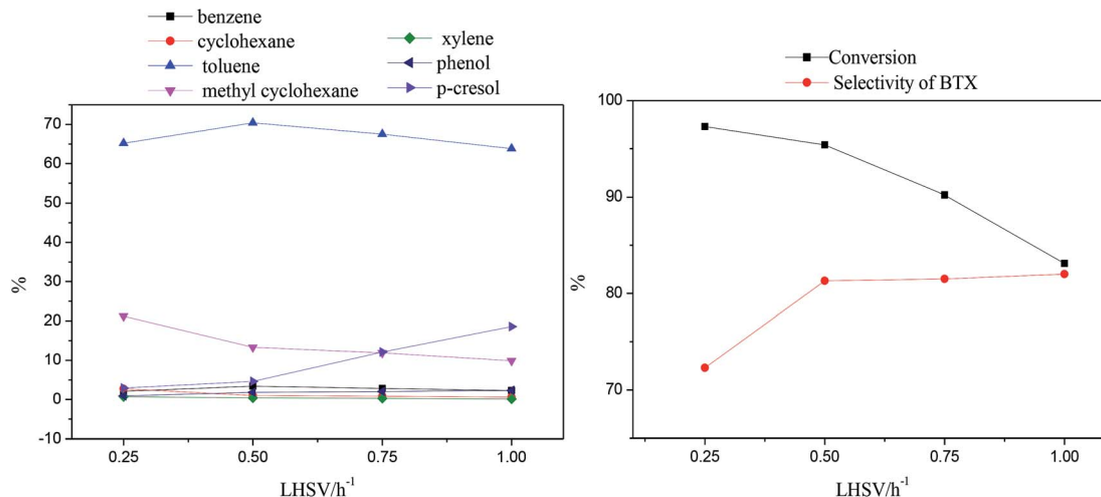


Fig. 7 Effects of LHSV on the products and *p*-cresol content via HDO.

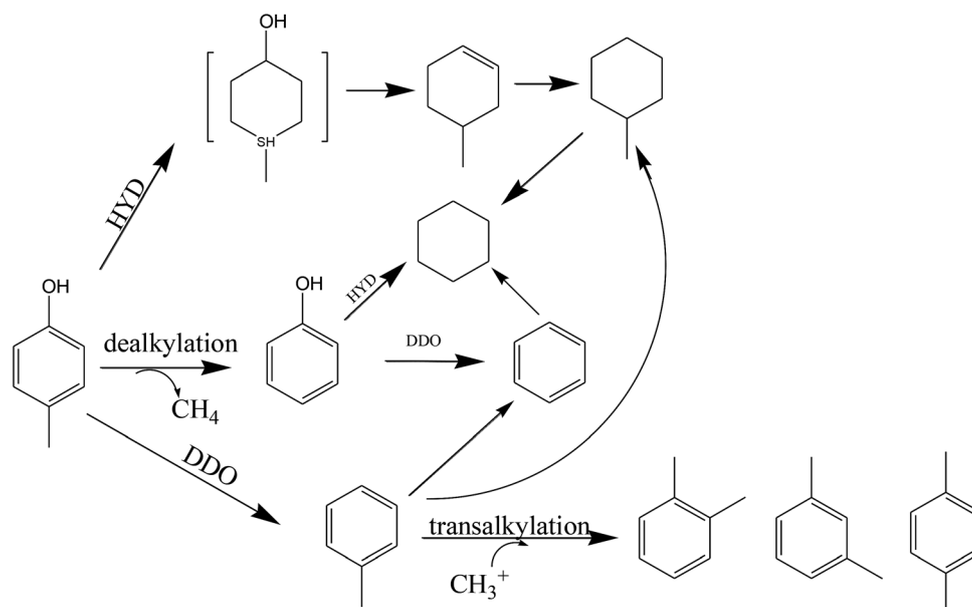
amounts of BTX products caused by the high LHSV corresponds to low conversion; however, at 0.3 h^{-1} , a low LHSV led to a large amount of aromatic ring consumption through full hydrogenation, which resulted in the conversion of aromatic hydrocarbons to naphthenic products.⁵³

Moreover, the amount of xylene slightly decreased but the content of phenol obviously increased from 0.9% to 2.3% with increasing LHSV. It is possible that the longer residence time corresponding to a low LHSV was beneficial to the transalkylation reaction but also facilitated the consumption of phenol by further hydrodeoxygenation.

Previous studies have been conducted on the path of methylphenol hydrodeoxygenation.^{11,33,54} Odeunmi⁵⁵ obtained similar reaction paths for the hydrodeoxygenation of cresol with Co-Mo/ γ -Al₂O₃ catalyst. They concluded that cresol first

produced an intermediate and then continued to produce toluene (k_1), methylcyclohexane (k_2) and phenol (k_4), respectively, and that there was interconversion between toluene and methylcyclohexane (k_3), showing $(k_1, k_3) \gg k_2, k_4$ on fresh catalyst. Kirby⁵⁶ also speculated that the reaction pathway of dibutylmethyl phenol hydrodeoxygenation on a novel organometallic catalyst precursor included dealkylation, direct deoxygenation, hydrodeoxygenation, and alkyl isomerization. By analyzing the product distributions at different reaction temperatures and LHSVs, a reaction conversion pathway of *p*-cresol on Ni-Ce/SiO₂ is proposed in Scheme 1.

It can be seen from the HDO test that the addition of a small amount of different binary metal M results in distinct reaction path selectivity. Supported Ni, a group VIII metal, especially when used for HDO on phenolic compounds, can adsorb the



Scheme 1 Reaction path of *p*-cresol on Ni-M/SiO₂ catalysts.



aromatic ring at the bridge site and repel C–O bond adsorption simultaneously, resulting in relatively strong hydrogenation reactivity and saturation ability for aromatic rings or C=C bonds.

After the addition of the binary metal M, the electron density of the nickel atoms changes significantly, which may cause a change in the adsorption performance of the catalyst surface for the groups of *p*-cresol and activate cleavage of the C–O bonds; that is, the priority of the hydrogenation reactions changes between the aromatic ring and C–O bond.⁵⁷ The increased electron density of the nickel atoms, which is related to the transfer of electrons from the binary metal M to Ni, tends to attack the C–O bonds and repel the electron-rich aromatic rings. This results in a preference for the hydrogenation process to involve direct deoxygenation after cleavage of the C–O bond rather than saturation of the aromatic ring.^{58,59} Moreover, changes in the acidity of the catalyst also affect the selectivity of HDO products on phenols. The low oxidation states of Co and Ce (observed from XRD and TPD) added to Ni/SiO₂ at lower reduction temperatures leads to the formation of more Lewis acid sites, resulting in optimization of hydrogenation performance.²⁹ In addition, the structural effects imparted by Ce and Co may cause changes in selectivity. Yang *et al.*⁵⁷ observed that for monometallic Ni catalyst, the nickel atom arrangement is continuous, so that the aromatic ring of phenol can be adsorbed in a plane form but the C–O bond is rejected; therefore, the cycloalkane is more favorable as the saturated product. However, after the binary metal entered the nickel alloy, the above continuity was destroyed, resulting in surface roughening. This created a rough atomic plane to prevent the adsorption of the benzene ring; therefore, aromatic ring hydrogenation was less likely to occur, and the adsorption probability of the C–O bond greatly increased. Other factors may exist in the effects of Fe on the increased selectivity of the DDO path. We can speculate that the combination of Fe, an oxophilic metal, with nickel not only enhanced the attraction of the catalyst to C_{aromatic}–O but also caused rejection of the π -electron cloud of the aromatic ring.⁴⁴ This resulted in an increasing tendency of C_{aromatic}–O bond cleavage and a weakened tendency of aromatic ring hydrogenation during the hydrogenation process.^{28,60} In contrast, Ni–Sn exhibited saturation of the phenolic ring. Due to the strong interactions between Sn and Ni, under the low load of Sn, an alloy state such as Ni₃Sn is formed, which was detected by XRD. Some previous studies showed that the formation of Ni₃Sn affects the hydrogenation priority of different groups. Rodiansono⁶¹ *et al.* found that the presence of Ni₃Sn alloy in Ni–Sn alloy catalysts increased the hydrogenation selectivity of C=O over that of C=C groups in unsaturated compounds. Therefore, we can speculate that Ni₃Sn was more favorable for promoting the hydrogenation of aromatic rings than the cleavage of C–O bonds in the process of hydrodeoxygenation of oxygen-containing phenolic compounds. Because the tin atoms are likely to be located in the vicinity of the steps, the influence on the continuity of the atomic surface is weak; thus, the surface readily adsorbs and is enriched by a large amount of hydrogen, and the phenol ring is thus more inclined to be saturated.^{50,57,61}

4 Conclusions

A series of bimetallic catalysts, Ni–M/SiO₂, were prepared and evaluated in a reduced state by HDO with *p*-cresol as a model. In the hydrodeoxygenation process, deoxygenation and trans-alkylation mainly occur, yielding saturated products such as methylcyclohexane and cyclohexane as well as unsaturated products such as toluene and benzene along with small amounts of phenol and xylene. The catalyst activity was as follows: Ni–Ce > Ni–Sn > Ni–Co > Ni–Fe > monometallic Ni. After the addition of the binary metal M, in comparison with the monometallic nickel, the dispersibility of the metal was observed to improve; meanwhile, the electron density also changed, which may be a factor contributing to the improvement of the activity.

Different binary metals exhibit great differences in the distribution of reaction products; Ce, Co, and Fe effectively increased the BTX selectivity and $S_{\text{BTX}}/S_{\text{cycloparaffin}}$, while Sn had the opposite effect. The reason for the difference in selectivity may be the joint effect of Lewis acid sites, the electron-rich phenomenon of nickel and reconstruction of the surface atomic arrangements. These changes may have a congenerous effect which not only strengthens the adsorption and cleavage of the polar C–O bond but also weakens the adsorption and saturation of the aromatic ring.

Conflicts of interest

There are no conflicts to declare.

References

- 1 T. Kan, X. Sun, H. Wang, *et al.*, Production of Gasoline and Diesel from Coal Tar via Its Catalytic Hydrogenation in Serial Fixed Beds, *Energy Fuels*, 2012, **26**, 3604–3611.
- 2 H. Clough, Gasoline Production from Coal Tar Oils, *Ind. Eng. Chem.*, 1957, **49**(4), 673–678.
- 3 D. Li, Z. Li, W. Li, *et al.*, Hydrotreating of low temperature coal tar to produce clean liquid fuels, *J. Anal. Appl. Pyrolysis*, 2013, **100**, 245–252.
- 4 S. Caplan, J. Ross, G. M. Sevag, *et al.*, Examination of the phenols of a low-temperature coal tar, *Ind. Eng. Chem., Anal. Ed.*, 2002, **6**(1), 379–380.
- 5 C. Karr, P. M. Brown, P. A. Estep, *et al.*, Identification and Determination of Low-Boiling Phenols in Low Temperature Coal Tar, *Anal. Chem.*, 1958, **30**(8), 1413–1415.
- 6 R. Wang, M. Sun, Q. Liu, *et al.*, Extraction and GC/MS analysis of phenolic compounds in low temperature coal tar from Northern Shaanxi, *J. China Coal Soc.*, 2011, **36**(4), 664–669.
- 7 M. Niu, X. Sun, R. Gao, *et al.*, Effect of Dephenolization on Low-Temperature Coal Tar Hydrogenation To Produce Fuel Oil, *Energy Fuels*, 2016, **30**(12), 10215–10221.
- 8 L. Wang, D. Li, F. Han, *et al.*, Experimental optimization and reactor simulation of coal-derived naphtha reforming over Pt–Re/ γ -Al₂O₃ using design of experiment and response



- surface methodology, *React. Kinet., Mech. Catal.*, 2018, **125**(1), 245–269.
- 9 X. Wei Xu and E. Chen Jiang, “BTX” from guaiacol HDO under atmospheric pressure: the effect of support and the carbon deposition, *Energy Fuels*, 2017, **31**(3), 2855–2864.
- 10 X. Xu, E. Jiang, Y. Du, *et al.*, BTX from the gas-phase hydrodeoxygenation and transmethylation of guaiacol at room pressure, *Renewable Energy*, 2016, **96**, 458–468.
- 11 Y. Romero, F. Richard and S. Brunet, Hydrodeoxygenation of 2-ethylphenol as a model compound of bio-crude over sulfided Mo-based catalysts: promoting effect and reaction mechanism, *Appl. Catal., B*, 2010, **98**(3–4), 213–223.
- 12 S. B. Gevert, M. Eriksson, P. Eriksson, *et al.*, Direct hydrodeoxygenation and hydrogenation of 2,6- and 3,5-dimethylphenol over sulphided CoMo catalyst, *Appl. Catal., A*, 1994, **117**(2), 151–162.
- 13 W. Tang, M. Fang, H. Wang, *et al.*, Mild hydrotreatment of low temperature coal tar distillate: product composition, *Chem. Eng. J.*, 2014, **236**, 529–537.
- 14 T. Kan, H. Wang, H. He, *et al.*, Experimental study on two-stage catalytic hydroprocessing of middle-temperature coal tar to clean liquid fuels, *Fuel*, 2011, **90**(11), 3404–3409.
- 15 W. Cui, W. Li, R. Gao, *et al.*, Hydroprocessing of Low-Temperature Coal Tar for the Production of Clean Fuel over Fluorinated NiW/Al₂O₃-SiO₂ Catalyst, *Energy Fuels*, 2017, **31**(4), 3768–3783.
- 16 E. Laurent, A. Centeno, and B. Delmon, Coke Formation during the Hydrotreating of Biomass Pyrolysis Oils: Influence of Guaiacol Type Compounds, in *Studies in Surface Science and Catalysis*, ed. B. Delmon and G. F. Froment, Elsevier, 1994, pp. 573–578.
- 17 Y. Yoshimura, T. Sato, H. Shimada, *et al.*, Influences of oxygen-containing substances on deactivation of sulfided molybdate catalysts, *Appl. Catal.*, 1991, **73**(1), 55–63.
- 18 O. İ. Şenol, E. M. Ryymin, T. R. Viljava, *et al.*, Effect of hydrogen sulphide on the hydrodeoxygenation of aromatic and aliphatic oxygenates on sulphided catalysts, *J. Mol. Catal. A: Chem.*, 2007, **277**(1), 107–112.
- 19 T. R. Viljava, R. S. Komulainen and A. O. I. Krause, Effect of H₂S on the stability of CoMo/Al₂O₃ catalysts during hydrodeoxygenation, *Catal. Today*, 2000, **60**(1), 83–92.
- 20 P. M. de Souza, L. Nie, L. E. P. Borges, *et al.*, Role of Oxophilic Supports in the Selective Hydrodeoxygenation of m-Cresol on Pd Catalysts, *Catal. Lett.*, 2014, **144**(12), 2005–2011.
- 21 C. A. Fisk, T. Morgan, Y. Ji, *et al.*, Bio-oil upgrading over platinum catalysts using *in situ* generated hydrogen, *Appl. Catal., A*, 2009, **358**(2), 150–156.
- 22 A. Gutierrez, R. K. Kaila, M. L. Honkela, *et al.*, Hydrodeoxygenation of guaiacol on noble metal catalysts, *Catal. Today*, 2009, **147**(3–4), 239–246.
- 23 K. Li, R. Wang and J. Chen, Hydrodeoxygenation of Anisole over Silica-Supported Ni₂P, MoP, and NiMoP Catalysts, *Energy Fuels*, 2011, **25**(3), 854–863.
- 24 M. D. Romero, J. A. Calles and A. Rodríguez, Influence of the Preparation Method and Metal Precursor Compound on the Bifunctional Ni/HZSM-5 Catalysts, *Ind. Eng. Chem. Res.*, 1997, **36**(9), 3533–3540.
- 25 J.-S. Moon, E.-G. Kim and Y.-K. Lee, Active sites of Ni₂P/SiO₂ catalyst for hydrodeoxygenation of guaiacol: a joint XAFS and DFT study, *J. Catal.*, 2014, **311**, 144–152.
- 26 Y. Yang, A. Gsilbert and C. Xu, Hydrodeoxygenation of bio-crude in supercritical hexane with sulfided CoMo and CoMoP catalysts supported on MgO: a model compound study using phenol, *Appl. Catal., A*, 2009, **360**(2), 242–249.
- 27 H. Y. Zhao, D. Li, P. Bui, *et al.*, Hydrodeoxygenation of guaiacol as model compound for pyrolysis oil on transition metal phosphide hydroprocessing catalysts, *Appl. Catal., A*, 2011, **391**(1), 305–310.
- 28 S. Leng, X. Wang, X. He, *et al.*, NiFe/γ-Al₂O₃: a universal catalyst for the hydrodeoxygenation of bio-oil and its model compounds, *Catal. Commun.*, 2013, **41**, 34–37.
- 29 P. M. Mortensen, J.-D. Grunwaldt, P. A. Jensen, *et al.*, Screening of Catalysts for Hydrodeoxygenation of Phenol as a Model Compound for Bio-oil, *ACS Catal.*, 2013, **3**(8), 1774–1785.
- 30 G. Bai, L. Niu, Z. Zhao, *et al.*, Ni-La-B amorphous alloys supported on SiO₂ and γ-Al₂O₃ for selective hydrogenation of benzophenone, *J. Mol. Catal. A: Chem.*, 2012, **363–364**, 411–416.
- 31 A. Y. Bunch, X. Wang and U. S. Ozkan, Adsorption characteristics of reduced Mo and Ni-Mo catalysts in the hydrodeoxygenation of benzofuran, *Appl. Catal., A*, 2008, **346**(1), 96–103.
- 32 S. A. Khromova, A. A. Smirnov, O. A. Bulavchenko, *et al.*, Anisole hydrodeoxygenation over Ni-Cu bimetallic catalysts: the effect of Ni/Cu ratio on selectivity, *Appl. Catal., A*, 2014, **470**, 261–270.
- 33 L. Nie, P. M. de Souza, F. B. Noronha, *et al.*, Selective conversion of m-cresol to toluene over bimetallic Ni-Fe catalysts, *J. Mol. Catal. A: Chem.*, 2014, **388–389**, 47–55.
- 34 L. Li, W. Yi, T.-W. Liu, *et al.*, Hydrogenation of 3-Hydroxypropanal to 1,3-Propanediol over Cu-V/Ni/SiO₂ Catalyst, *New J. Chem.*, 2017, **41**, 8965–8976.
- 35 Z. Wang, Q. Liu, J. Yu, *et al.*, Surface structure and catalytic behavior of silica-supported copper catalysts prepared by impregnation and sol-gel methods, *Appl. Catal., A*, 2003, **239**(1), 87–94.
- 36 F. Yang, D. Liu, Y. Zhao, *et al.*, Size Dependence of Vapor Phase Hydrodeoxygenation of m-Cresol on Ni/SiO₂ Catalysts, *ACS Catal.*, 2018, **8**(3), 1672–1682.
- 37 M. G. Prakash, R. Mahalakshmy, K. R. Krishnamurthy, *et al.*, Studies on Ni-M (M = Cu, Ag, Au) bimetallic catalysts for selective hydrogenation of cinnamaldehyde, *Catal. Today*, 2016, **263**, 105–111.
- 38 M. V. Bykova, D. Y. Ermakov, V. V. Kaichev, *et al.*, Ni-based sol-gel catalysts as promising systems for crude bio-oil upgrading: guaiacol hydrodeoxygenation study, *Appl. Catal., B*, 2012, **113–114**, 296–307.
- 39 O. Clause, L. Bonneviot and M. Che, Effect of the preparation method on the thermal stability of silica-supported nickel oxide as studied by EXAFS and TPR techniques, *J. Catal.*, 1992, **138**(1), 195–205.



- 40 S. Zhang, S. Muratsugu, N. Ishiguro, *et al.*, Ceria-Doped Ni/SBA-16 Catalysts for Dry Reforming of Methane, *ACS Catal.*, 2013, **3**(8), 1855–1864.
- 41 S. Hwang, J. Lee, U. G. Hong, *et al.*, Hydrogenation of carbon monoxide to methane over mesoporous nickel-M-alumina (M=Fe, Ni, Co, Ce, and La) xerogel catalysts, *J. Ind. Eng. Chem.*, 2012, **18**(1), 243–248.
- 42 T. Ishihara, K. Eguchi and H. Arai, Hydrogenation of carbon monoxide over SiO₂-supported Fe-Co, Co-Ni and Ni-Fe bimetallic catalysts, *Appl. Catal.*, 1987, **30**(2), 225–238.
- 43 A. Onda, T. Komatsu and T. Yashima, Characterizations and catalytic properties of fine particles of Ni-Sn intermetallic compounds supported on SiO₂, *J. Catal.*, 2004, **221**(2), 378–385.
- 44 S. Sitthisa, W. An and D. E. Resasco, Selective conversion of furfural to methylfuran over silica-supported NiFe bimetallic catalysts, *J. Catal.*, 2011, **284**(1), 90–101.
- 45 C. Minchev, S. A. Zubkov, V. Valtchev, *et al.*, Nature of the active sites and catalytic activity of SAPO-5 synthesized in the presence of nickel cations, *Appl. Catal.*, A, 1994, **119**(2), 195–204.
- 46 K. Fang, J. Ren and Y. Sun, Effect of nickel precursors on the performance of Ni/AlMCM-41 catalysts for n-dodecane hydroconversion, *J. Mol. Catal. A: Chem.*, 2005, **229**(1), 51–58.
- 47 S. G. Zhang, K. Yorimitsu, S. Nohara, *et al.*, Surface analysis of an amorphous MgNi alloy prepared by mechanical alloying for use in nickel-metal hydride batteries, *J. Alloys Compd.*, 1998, **270**(1), 123–126.
- 48 H. W. Nesbitt, D. Legrand and G. M. Bancroft, Interpretation of Ni 2p XPS spectra of Ni conductors and Ni insulators, *Phys. Chem. Miner.*, 2000, **27**(5), 357–366.
- 49 S. J. Sawhill, D. C. Phillips and M. E. Bussell, Thiophene hydrodesulfurization over supported nickel phosphide catalysts, *J. Catal.*, 2003, **215**(2), 208–219.
- 50 A. Borgna, B. Anderson, A. Saib, *et al.*, Pt-Co/SiO₂ Bimetallic Planar Model Catalysts for Selective Hydrogenation of Crotonaldehyde, *J. Phys. Chem. B*, 2004, **108**(46), 17905–17914.
- 51 M. H. Peyrovi, T. Rostamikia and N. Parsafard, Competitive Hydrogenation of Benzene in Reformate Gasoline over Ni Supported on SiO₂, SiO₂-Al₂O₃, and Al₂O₃ Catalysts: Influence of Support Nature, *Energy Fuels*, 2018, **32**(11), 11432–11439.
- 52 Y. Li, J. Fu and B. Chen, Highly selective hydrodeoxygenation of anisole, phenol and guaiacol to benzene over nickel phosphide, *RSC Adv.*, 2017, **7**(25), 15272–15277.
- 53 X. Xu, Z. Li, Y. Sun, *et al.*, High quality fuel from the upgrading of heavy bio-oil by the combination ultrasonic with mutual solvent, *Energy Fuels*, 2018, **32**, 3477–3487.
- 54 E. Furimsky, Catalytic hydrodeoxygenation, *Appl. Catal.*, A, 2000, **199**(2), 147–190.
- 55 E. O. Odeunmi and D. F. Ollis, Catalytic hydrodeoxygenation: I Conversions of o-, p-, and m-cresols, *J. Catal.*, 1983, **80**(1), 56–64.
- 56 S. R. Kirby, C. Song and H. H. Schobert, Hydrodeoxygenation of O-containing polycyclic model compounds using a novel organometallic catalyst precursor, *Catal. Today*, 1996, **31**(1), 121–135.
- 57 F. Yang, D. Liu, H. Wang, *et al.*, Geometric and electronic effects of bimetallic Ni-Re catalysts for selective deoxygenation of m-cresol to toluene, *J. Catal.*, 2017, **349**, 84–97.
- 58 X. Wang and J. Chen, Effects of indium on Ni/SiO₂ catalytic performance in hydrodeoxygenation of anisole as model bio-oil compound: suppression of benzene ring hydrogenation and C-C bond hydrogenolysis, *Chin. J. Catal.*, 2017, **38**(11), 1818–1830.
- 59 C. Li, Y. Chen, S. Zhang, *et al.*, Ni-In Intermetallic Nanocrystals as Efficient Catalysts toward Unsaturated Aldehydes Hydrogenation, *Chem. Mater.*, 2013, **25**(19), 3888–3896.
- 60 A. N. Mansour, D. E. Sayers, J. W. Cook, *et al.*, X-ray absorption studies of some platinum oxides, *J. Phys. Chem.*, 1984, **88**(9), 1778–1781.
- 61 Rodiansono, S. Khairi, T. Hara, *et al.*, Highly efficient and selective hydrogenation of unsaturated carbonyl compounds using Ni-Sn alloy catalysts, *Catal. Sci. Technol.*, 2012, **2**(10), 2139.

