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Hydrotreatment of model compounds with catalysts of NiW/Al₂O₃ and NiWP/Al₂O₃ to simulate low temperature coal tar oil

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Eight characteristic model compounds were employed as hydrotreatment feedstocks to simulate low temperature coal tar oil. A series of NiW/Al₂O₃ catalysts with different Ni/W ratio and NiWP/Al₂O₃ catalysts with different phosphorus content were prepared by incipient wetness co-impregnation. Both catalysts were characterized by N₂ adsorption–desorption, X-ray diffraction, HRTEM, H₂-TPR, NH₃-TPD and FT-IR. The heteroatom removal and polyaromatic hydrogenation conversions were more than 90% over these catalysts. Ni₆W₂₀ showed the highest conversion among the NiW/Al₂O₃ catalysts due to the presence of relatively short, thick slabs when compared to the other Ni/W ratios. Phosphorus was added to adjust the acid concentration of NiW sulfides. The best hydrotreatment performance was seen with the 0.8 wt% phosphorus catalyst, which was attributed to the highest medium strength acid concentration. The results suggested the medium acid on the NiW sulfide was probably the active site for the hydrogenation of the aromatic ring in the coal tar oil hydrotreatment process. The self-prepared catalyst showed higher HDS and HDN activity than a commercial catalyst.

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1 Introduction

Coal is the main energy source in China. Poly-generation technology, combining coal pyrolysis, gasification and combustion process, can convert coal into gas, tar and steam. And then gas and tar can be converted into chemical products and fuels. Steam can be used in electricity generation, which is a promising way to deal with the problem of energy demand and environmental protection in China.¹ Coal tar oil is one of the main products of coal pyrolysis at low temperature. Therefore, coal tar oil can be transformed into clean fuel oil by hydroprocessing in this system.²-6

Coal tar oil consists of thousands of chemical compounds.⁷ Such a complex composition is hard to investigate. As such, both the understanding of coal tar oil hydrotreatment and the screening of the catalyst for this process are very difficult. It is not economical and practical to detect the transformation of every compound in coal tar oil during hydroprocessing. Fortunately, the compounds in coal tar oil can be separated into several groups with different functional groups. The hydrogenation mechanisms of the compounds in one group are almost the same. Thus, they can be represented by one characteristic compound in the group. Therefore, several characteristic

compounds can be chosen to simulate coal tar oil. Wandas et al.8 investigated the hydroprocessing of three-component model mixtures to simulate fast pyrolysis tars. They found that the interactions of naphthalene and cresols in mixtures resulted in a lower efficiency of naphthalene hydrogenation as well as cresol hydrodeoxygenation than that of the hydroconversion of the single components. Lemberton et al.9-11 reported that they had used phenanthrene, carbazole and 1naphthol as the model compounds of coal tar oil. In previous studies, three or four model compounds were generally chosen to simulate coal tar oil. However, the model compounds were not abundant enough to represent low temperature coal tar oil (LTCTO), which is composed of O-containing compounds, Ncontaining compounds, S-containing compounds, aromatics and aliphatics. Ni(Co)-Mo(W) sulfides have been extensively studied and used as hydrotreatment catalysts for years in petroleum industry. The performance of metal sulfide catalysts in coal tar oil hydroprocessing must be different from that seen in the hydroconversion of crude-derived feedstock due to much higher content of phenols, N-containing compounds and aromatics in coal tar oil. The effect of MoO₃ content in NiMo₃, Ni/W ratio, 12 zeolite 13 and fluorine 14 content in NiW catalysts on LTCTO distillate hydrotreatment has been reported. However, there is little information about these catalysts when applied in coal tar oil hydroprocessing.

In this work, eight compounds were chosen as the model mixtures to simulate LTCTO. The performance of model compounds hydrotreating over NiW/Al₂O₃ and NiWP/Al₂O₃ was

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investigated, and the effects of Ni/W ratio and P content on the physical and chemical properties of catalysts were also studied.

2 Experiment

2.1 Catalyst preparation

NiW/Al $_2$ O $_3$ and NiWP/Al $_2$ O $_3$ catalysts were prepared by incipient wetness co-impregnation on γ -Al $_2$ O $_3$. Ni(NO $_3$) $_2 \cdot 6H_2$ O and (NH $_4$) $_6$ H $_2$ W $_{12}$ O $_{40}$ were dissolved in de-ionized water. The γ -Al $_2$ O $_3$ was impregnated using an aqueous solution at room temperature for 10 h and dried at 120 °C overnight. The resulting sample was calcinated at 550 °C for 4 h. The tungsten content in NiW/Al $_2$ O $_3$ was fixed at 20 wt%, and the nickel content was 2 wt%, 4 wt%, 6 wt% and 8 wt%, with catalysts named as Ni $_2$ W $_2$ O $_3$, Ni $_4$ W $_2$ O $_3$, Ni $_6$ W $_2$ O $_3$ and Ni $_8$ W $_2$ O $_3$, respectively.

NiWP/Al₂O₃ was obtained using the same steps used to prepare NiW/Al₂O₃. Phosphoric acid was the source of phosphorus for the co-impregnation solution of Ni(NO₃)₂·6H₂O and (NH₄)₆H₂W₁₂O₄₀. The synthesized NiWP/Al₂O₃ catalysts were denoted as NiWP-x, where x represented the phosphorus content in the precursor. The nickel and tungsten content in NiWP/Al₂O₃ was fixed at 6 wt% and 20 wt%, and the phosphorus content was 0.5 wt%, 0.8 wt%, 1.2 wt% and 1.5 wt% respectively.

2.2 Catalyst characterization

With the exceptions of X-ray diffraction (XRD), FT-IR and $\rm H_2$ temperature-programmed reduction ($\rm H_2$ -TPR), all samples should be presulfided prior to catalyst characterization. The presulfidation was carried out in a fix-bed reactor at 360 °C and 3 MPa for 4 h. The presulfiding solution was 5 wt% carbon disulfide dissolved in cyclohexane. The flow rates of the $\rm H_2$ and the presulfiding solution were 100 ml min⁻¹ and 0.5 ml min⁻¹, respectively.

 $\rm N_2$ adsorption–desorption isotherms of the catalysts were measured on a Micromeritics TriStarII3020 apparatus. The specific surface area was calculated by the Brunauer–Emmett–Teller (BET) equation. The desorption branch of the isotherm was used to calculate the average pore diameter according to the Barrett–Joyner–Halenda (BJH) method.

X-ray diffraction (XRD) of the catalysts were recorded with a X'Pert PRO diffractometer (PANalytical), using monochromatic CuK α radiation ($\lambda=1.541$ Å). The X-ray tube was set at 40 kV and 40 mA.

High resolution transmission electron microscope (HRTEM) images were obtained on a Tecnai G2 F20 S-Twin (FEI) operating at 200 kV. The samples were dissolved ultrasonically and placed on a 200-mesh copper grid coated with carbon.

 $\rm H_2$ temperature programmed reduction ($\rm H_2$ -TPR) was conducted on an AutoChem II 2920 instrument with a mixture of $\rm H_2/He$ ($\rm H_2$ 10 vol%) at a flow rate of 30 ml min $^{-1}$ (STP). A 0.1 g sample was placed into a quartz tube, and heated at a rate of 10 °C min $^{-1}$ to 900 °C.

 NH_3 temperature programmed desorption (NH $_3$ -TPD) was carried out on AutoChem II 2920 instrument as well. A NH $_3$ /He mixture (NH $_3$ 10 vol%) was absorbed on 0.1 g sample of catalyst

at 100 °C. The sample was then heated to 700 °C at a rate of 10 °C min^{-1} .

Fourier transform infrared spectroscopy (FT-IR) of absorbed pyridine was conducted on a Nicolet 6700 FT-IR Spectrometer. Self-supporting wafers with weight of 13 mg catalysts were activated at 400 °C for 4 h under 10⁻² Pa, and then cooled down to room temperature. Subsequently, pyridine vapors were admitted, and after equilibration samples were heated to 200 °C at a rate of 10 °C min⁻¹. Then the samples were outgassed for 0.5 h to evacuate the excess pyridine. Infrared spectra of catalysts were recorded with a resolution of 2.0 cm⁻¹.

2.3 Model compounds selection

Model compounds for simulating LTCTO should be carefully selected. The number of model compounds should be neither too small to represent the character of coal tar oil, nor too large to be analysed. In previous research, ^{15–18} aromatics were chosen and thoroughly studied as the model compounds to simulate the hydrogenation of high temperature coal tar oil. However, as listed in Table 1, a high content of phenol and paraffins were found in LTCTO created by coal pyrolysis, compared with the high temperature coal tar oil from coking. These differences should be considered when choosing the model compounds for representing LTCTO.

In this study, eight compounds with different functional groups were chosen as the hydrotreatment feedstock to simulate LTCTO. The component of feedstock was listed in Table 2. *N*-Cetane was chosen to represent saturates because the carbon number of alkanes in coal tar oil was 10–30. The most abundant O-containing compounds in LTCTO were phenols. *O*-Cresol was chosen for its relatively low hydrodeoxygenation activity among the three kinds of cresols. Quinoline and dibenzothiophene had the highest content of N-containing compounds and S-containing compounds in LTCTO, respectively. The reaction networks of these compounds were also characteristic in hydrodenitrogenation and hydrodesulfurization, respectively. The content of each model compound in the hydrotreatment feedstock was based on the composition analysis of LTCTO distillate by GC-MS.

2.4 Hydrotreatment unit

Catalyst activity measurements were carried out in a fixed-bed reactor with inner diameter of 12 mm. The reactor system (as shown in Fig. 1) has been described elsewhere.²¹ To establish an uniform flow distribution in the reactor and to avoid

Table 1 The difference of main compounds contents between LTCTO and high temperature coal tar oil (HTCTO)

	LTCTO ^a /wt%	HTCTO ^b /wt%
Paraffins	21	3-5
Phenols	27	2-3
Diaromatics	7	10-27
Polyaromatics	5	8-16

^a From ref. 19. ^b From ref. 20.

Table 2 The component of hydrotreatment feedstock

Compounds	Boiling point/°C	Content/ wt%	Representing compounds in coal tar oil
<i>O</i> -Xylene	144	10	Monoaromatics
O-Cresol	191	18	O-Containing compounds
1-Methylnaphthalene	240	25	Diaromatics
Biphenyl	255	10	Diaromatics
Phenanthrene	340	10	Polyaromatics
N-Cetane	287	21	Saturates
Quinoline	238	5	N-Containing compounds
Dibenzothiophene	332	1	S-Containing compounds

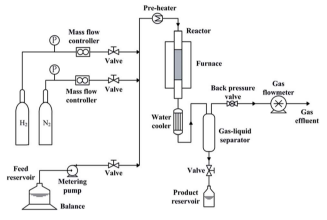


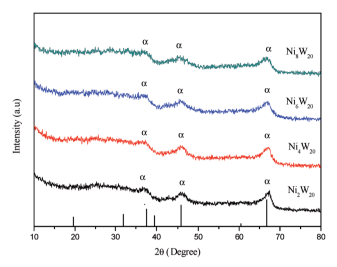
Fig. 1 Flow diagram of hydrotreatment unit.

overheating, 7.5 ml catalyst was packed into the reactor with a dilution of 17.5 ml quartz sand. Prior to activity measurement, catalysts should be in situ presulfided. The process of presulfidation was the same as that before catalyst characterization. After presulfidation, temperature and pressure were adjusted to the desired reaction values. Liquid feed was shifted to the hydrotreatment feedstock. Reaction conditions were 360 °C, 6 MPa, LHSV = 1 h^{-1} and H_2 /liquid ratio of 1000 : 1. Each run was conducted for 10 h. The first sample was collected 4 h after initiating the reaction to ensure reaction conditions were stable. The composition distribution of feedstock and products was detected by GC (Agilent 7890A) with an HP-5 capillary column (30 m \times 0.25 mm \times 0.25 μ m, Agilent). The compounds in feedstock and products were identified by GC-MS. For comparison, a commercial hydrotreating catalyst (bought from SINOPEC QILU CO. LTD) was also tested.

3 Results and discussion

Catalyst characterization

3.1.1 NiW catalysts. Fig. 2 shows the XRD patterns of NiW/ Al_2O_3 catalysts. Peaks representing Al_2O_3 support ($2\theta = 37^\circ, 46^\circ,$ and 67°) can be observed. The bars at the bottom of the figure



XRD patterns of NiW catalysts.

show the characteristic peaks of γ-Al₂O₃ (pdf#29-0063). No nickel oxide or tungsten oxide peaks were detected, indicating that metal had a good dispersion on the support and the particle diameter was small.

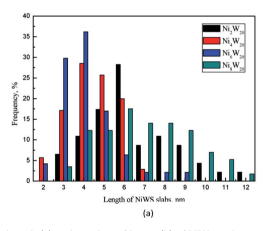
The transmission electron microscopy images of four NiW catalysts are shown in Fig. 3. Black lines representing the layered NiWS slabs could be observed clearly. The length and number of layers of slabs were determined statistically through more than 10 images and calculated from measurement of more than 50 slabs. The distribution of the slab length and the number of slab layers are shown in Fig. 4. As shown in Fig. 4(a), the slab length with the highest frequency on Ni₂W₂₀ and Ni₈W₂₀ was 6 nm, while the corresponding value on Ni₄W₂₀ and Ni₆W₂₀ was 4 nm. More than 90% of the slabs on Ni₄W₂₀ and Ni₆W₂₀ had a length of 3-6 nm. However, the distributions of slab length on Ni₂W₂₀ and Ni₈W₂₀ were more even. Some slabs on Ni₂W₂₀ and Ni₈W₂₀ showed a length of more than 10 nm. From Fig. 4(b), the frequency of two layers was the highest on Ni_2W_{20} and Ni_8W_{20} . More slabs were observed with three layers rather than two layers on Ni₄W₂₀. The frequencies of slabs with two layers and three layers were almost the same on Ni_6W_{20} .

The average length and number of layers of NiWS slabs are given in Table 3. As shown in TEM images and calculation results, Ni₂W₂₀ and Ni₈W₂₀ exhibited longer and thinner NiWS slabs. Ni₄W₂₀ and Ni₆W₂₀ had the highest average number of layers and the lowest average length of NiWS slabs, respectively, among the four NiW catalysts.

Fig. 5 shows the H₂-TPR curves of NiW catalyst precursors. Two peaks and a peak shoulder could be seen in each curve. The low temperature peak at 400-600 °C was regarded as the reduction of NiO, while the high temperature peak at 700-800 °C was considered as the reduction of WO₃.²² The low temperature reduction peak shifted to the low temperature zone as the Ni/W ratio increased from 0.1 to 0.3. NiO reduction could occur at a temperature as low as 300 °C when the Ni/W ratio was 0.3 and 0.4. However, the peak at high temperature shifted to higher temperature when Ni/W ratio increased. The results indicated that as Ni/W increased, NiO reduction was easier to achieve Paper

(a) 10 mm (b) (c) 10 mm (d)

Fig. 3 TEM images of NiW catalysts. (a) Ni_2W_{20} , (b) Ni_4W_{20} , (c) Ni_6W_{20} , (d) Ni_8W_{20}



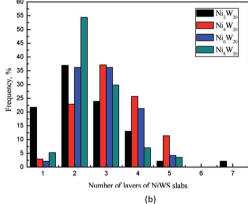


Fig. 4 The slabs length (a) and number of layers (b) of NiW catalysts.

while WO₃ reduction was not. More nickel in the catalyst caused the separation of the high temperature reduction peak and the peak shoulder, which might represent the reduction of Ni-W-O

Table 3 The average length and number of layers of slabs on NiW catalysts

	Ni_2W_{20}	Ni_4W_{20}	$\mathrm{Ni_6W_{20}}$	Ni ₈ W ₂₀
Average length/nm	6.39	4.46	4.17	6.96
Average number of layers	2.46	3.20	2.89	2.49

phase. The shifting of the peak shoulder to a lower temperature was attributed to a high amount of nickel, weakening the strength of the interaction between Ni and W.

The physical properties of NiW catalysts were listed in Table 4. As shown, Ni_6W_{20} had the highest BET surface area and pore volume. The average pore diameter decreased with the increase of Ni/W ratio.

3.1.2 NiWP catalysts. Fig. 6 shows the XRD patterns of Ni_6W_{20} and NiWP catalysts. Similar to the trend of NiW catalysts, Ni, W or P oxide peaks were not detected from the patterns, indicating that the addition of phosphorus did not influence the phases on NiW catalysts.

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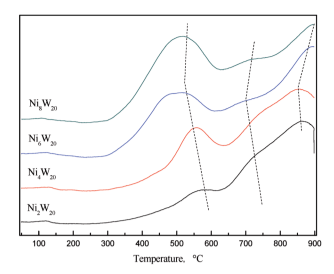


Fig. 5 H₂-TPR curves of NiW catalyst precursors.

Table 4 The physical properties of NiW catalysts

	BET surface area (m² g ⁻¹)	Pore volume (cm³ g ⁻¹)	Average pore diameter (nm)
Ni ₂ W ₂₀	112.1	0.193	5.18
Ni_4W_{20}	103.6	0.191	5.16
Ni_6W_{20}	133.4	0.212	4.47
Ni_8W_{20}	129.8	0.182	4.16

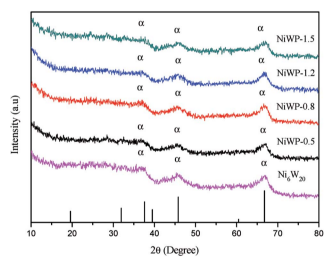


Fig. 6 XRD patterns of Ni₆W₂₀ and NiWP catalysts.

The physical properties of NiWP catalysts with different phosphorus contents are shown in Table 5. With the increase of phosphorus content in catalyst, the BET surface area of NiWP catalysts increased and reached a maximum of 138.2 m 2 g $^{-1}$ at NiWP-0.8, then decreased significantly to 100.0 m 2 g $^{-1}$ when phosphorus content was 1.5 wt%. Pore volume showed the same trend as that of BET surface area. It could be deduced from the trend of BET surface area and pore volume that pores were blocked when phosphorus content was more than 1 wt%.

Table 5 The physical properties of Ni₆W₂₀ and NiWP catalysts

	BET surface area (m² g ⁻¹)	Pore volume (cm ³ g ⁻¹)	Average pore diameter (nm)
Ni ₆ W ₂₀	133.4	0.212	4.47
NiWP-0.5	120.0	0.186	4.54
NiWP-0.8	138.2	0.206	4.25
NiWP-1.2	122.6	0.188	4.48
NiWP-1.5	100.0	0.154	4.47

Fig. 7 shows the H_2 -TPR curves of Ni_6W_{20} and NiWP catalyst precursors. From the Fig. 7, the curves of NiWP-0.5, NiWP-0.8 and NiWP-1.2 were almost the same. However, the peaks at low temperature and high temperature both shifted to a higher temperature with further addition of phosphorus. A 50 °C shifting of the low temperature peak was observed between the curves of NiWP-1.5 and the other three curves. The variation showed that the reduction of metal oxides would be difficult when the phosphorus content on the catalyst was 1.5 wt% or more. This may have been due to a stable phase combining with phosphorus and metal, which resulted in the formation of a Al_2O_3 support.

Fig. 8 shows the $\mathrm{NH_3}\text{-}\mathrm{TPD}$ curves of $\mathrm{Ni_6W_{20}}$ and NiWP catalysts. Three or four peaks were observed in the curves. Compared with $\mathrm{Ni_6W_{20}}$, the peak at low temperatures (<200 °C) on NiWP catalysts was almost the same. However, a large difference was seen on the peaks at higher temperatures (>300 °C) after the addition of phosphorus. The peak at 500 °C on the $\mathrm{Ni_6W_{20}}$ curve shifted to a lower temperature by about 100 °C. Desorption peaks became broader and some peak shoulders appeared when phosphorus was added in the catalyst. The results revealed that the addition of phosphorus weakened the strength of medium acid sites and new acid sites were generated.

The acid sites concentration of Ni₆W₂₀ and NiWP catalysts obtained from the Gaussian deconvolution are reported in

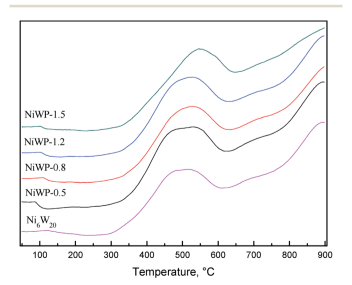


Fig. 7 H₂-TPR curves of Ni₆W₂₀ and NiWP catalyst precursors.

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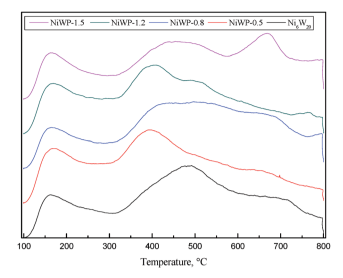


Fig. 8 NH₃-TPD curves of Ni₆W₂₀ and NiWP catalysts.

Table 6 Acid sites concentration distribution on Ni₆W₂₀ and NiWP (unit: mmol q^{-1})

	Weak acid (<300 °C)	Medium acid (300–550 °C)	Strong acid (>550 °C)	Total
Ni ₆ W ₂₀	0.110 (26) ^a	0.220 (52)	0.094 (22)	0.423
NiWP-0.5	0.157 (37)	0.201 (48)	0.064 (15)	0.422
NiWP-0.8	0.085 (18)	0.239 (50)	0.151 (32)	0.475
NiWP-1.2	0.117 (29)	0.163 (41)	0.118 (30)	0.397
NiWP-1.5	0.096 (23)	0.195 (47)	0.124 (30)	0.415

^a The value in the parentheses is the percentage of various strength of

Table 6. According to the maximum temperature, the acid sites were classified into the weak (<300 °C), medium (300-550 °C) and strong (>550 °C). As can be seen in Table 6, the medium acid was dominant on all catalysts. After adding 0.5 wt% phosphorus, the weak acid concentration on the catalyst increased, accompanied with a decrease of medium and strong acid concentrations. As phosphorus content further increased, the concentration of medium and strong acids first increased, and then decreased. The observed medium acid concentration trend was: NiWP-0.8 > Ni_6W_{20} > NiWP-0.5 > NiWP-1.5 > NiWP-1.2. The change of acid site distribution of Ni₆W₂₀ and NiWP catalysts indicated that an increase of phosphorus content on the catalysts resulted in the increase of the weak acid concentration, followed by an increase of the medium and strong acid concentrations. However, the improvement on acid concentration by adding phosphorus was only slight when the phosphorus content was higher than 1 wt%.

Fig. 9 shows the FT-IR spectrogram of pyridine adsorbed on Ni₆W₂₀ and NiWP catalysts outgassed at 200 °C. According to literature, 23,24 the band at 1540 cm⁻¹ was due to pyridine absorbed on Brønsted acid sites, and the band at 1450 cm⁻¹ could be assigned to coordinately bound pyridine interacting with Lewis acid sites. As for the band at 1490 cm⁻¹, it was

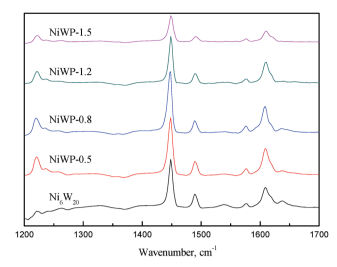


Fig. 9 FT-IR spectrogram of pyridine adsorbed on Ni₆W₂₀ and NiWP catalysts outgassed at 200 °C.

associated with an interaction of pyridine with both Brønsted and Lewis acid sites.23 The results indicated that both the Ni₆W₂₀ and NiWP catalysts showed Brønsted and Lewis acid sites while the height of peaks at corresponding bands were quite different. For Ni₆W₂₀, the peak at 1540 cm⁻¹ could be observed which represented the Brønsted acid sites. After the addition of phosphorus the peak was less distinct and it could hardly be observed when the phosphorus loading was 1.5 wt%. As for the band at 1450 cm⁻¹, the height of the peak increased first and then decreased as the loading of phosphorus and the same trend was observed at 1490 cm⁻¹.

The relative acidic amount of Brønsted and Lewis acid sites were determined by calculating the areas of pyridine adsorption bands at 1540 and 1450 cm⁻¹, and were summarized in Table 7. We could see from the table that the Brønsted acid amount decreased obviously after the introduction of phosphorus. This might be due to the loss of tetrahedral framework aluminum.25 The Lewis acid amount increased first and then decreased, during which the highest was reached on NiWP-0.5. As for total acid amount, the highest was obtained on NiWP-0.8 because of its relatively higher Brønsted acid amount compared with other NiWP catalysts. This was in accordance with the results in Table 6. From the ratio of Brønsted acid to Lewis acid we could also find that the acid type was changed significantly after adding phosphorus. Therefore, the addition of phosphorus not only changed the total acid sites concentration, but also modified the types of acid sites on the surface of catalysts by decreasing Brønsted acid sites and generating new Lewis acid sites.

3.2 NiW catalysts activity

3.2.1 The effect of Ni/W ratio on heteroatom removal. In LTCTO, heteroatom compounds include O-containing, Ncontaining, and S-containing compounds. In the model mixture, these are represented by o-cresol, quinolone and dibenzothiophene (DBT), respectively. No O-containing or Scontaining intermediates were detected in the product.

Table 7 Acid properties of Ni₆W₂₀ and NiWP determined by FT-IR of absorbed pyridine (unit: µmol/g)

	Brønsted acid amount	Lewis acid amount	Total acid amount	B/L
Ni_6W_{20}	60.03	293.88	353.91	0.204
NiWP-0.5	28.94	423.22	452.16	0.068
NiWP-0.8	42.88	412.04	454.92	0.104
NiWP-1.2	11.27	236.45	247.72	0.048
NiWP-1.5	4.87	178.03	182.90	0.027

Therefore, hydrodeoxygenation (HDO) and hydrodesulfurization (HDS) conversion were equal to the value of the conversion of o-cresol and DBT, respectively. However, tetrahydroquinoline, one of the intermediates in hydrodenitrogenation (HDN), was found in product. Consequently, HDN conversion was calculated as the ratio of the molar concentration of non-N-containing compounds in the HDN network and the molar concentration of quinoline in feedstock.

The HDO, HDS, and HDN conversion as a function of Ni/W ratio is shown in Fig. 10. From Fig. 10, HDO, HDS and HDN conversion on all catalysts approached or exceeded 90%, which shows the high hydrotreatment activity of NiW catalysts. HDO conversion approached 100% when the Ni/W ratio was higher than 0.1. HDO conversion was higher than HDS and HDN conversion, which mostly reside in the 90-95% range. With an increasing Ni/W ratio, HDO conversion first increased and then decreased slightly. HDN conversion decreased to less than 90% at a Ni/W ratio of 0.2, and then went up to about 93%. In contrast, HDS conversion decreased linearly when the Ni/W ratio increased, though the variation was less than 2%.

Complex reaction networks with several reaction pathways were reported on cresol HDO,26 DBT HDS,27 and quinoline HDN.28 The hydrogenation (HYD) pathway, which is hydrogenation followed by deoxygenation, was the main pathway for ocresol HDO. However, the direct desulfurization (DDS) pathway dominated DBT HDS. In quinoline HDN, the hydrogenation of aromatic rings was followed by C-N bond breaking. The decrease of DBT HDS conversion with the increase of Ni/W ratio showed that direct desulfurization activity was not enhanced by the addition of Ni in this range. The HDO conversion increased notably as the Ni/W ratio increased from 0.1 to 0.2, indicating that the hydrogenation activity of the aromatic ring was improved. The decrease of HDN conversion in the same Ni/W range might be due to the low denitrogenation activity, since it was more difficult for C-N bond to break than the C-O bond.

3.2.2 The effect of Ni/W ratio on aromatics hydrogenation. Four aromatic compounds were investigated in the hydrotreatment of model mixtures. The conversions of these compounds over the change of Ni/W ratio are exhibited in Fig. 11. The conversion of biphenyl was calculated using the assumption that the selectivity of biphenyl in DBT HDS was 90%,29 because biphenyl was the product of DBT HDS as well. As shown in Fig. 11, o-xylene representing monoaromatics showed the lowest conversion behaviour of 10-15%. The conversion of biphenyl was less than 50%, while the conversions of 1-methylnaphthalene and phenanthrene were above 90%. Nearly constant conversion of o-xylene in the range of all Ni/W ratios indicated that the catalyst surface structure and activity change by Ni addition made little impact on the hydrogenation of monoaromatics. However, the conversion of diaromatic and triaromatic compounds increased with the increase of Ni/W ratio, reaching a maximum at the Ni/W ratio of 0.2 or 0.3, and then decreased with further increase of Ni/W ratio. The variation of biphenyl and 1-methylnaphthalene conversions was larger than that of phenanthrene. This is because the hydrogenation of phenanthrene into dihydrophenanthrene was easier to achieve under this reaction condition, resulting in a small effect from the catalysts.

There was a relationship between the aromatic hydrogenation activity and NiWS slab features on NiW catalysts. Ni₄W₂₀ and Ni₆W₂₀ had smaller average slab lengths and a larger average number of layers, showing better aromatics hydrogenation performance than Ni₂W₂₀ and Ni₈W₂₀. The thickness

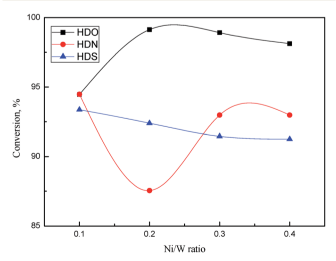


Fig. 10 HDO, HDS, and HDN conversions as a function of Ni/W ratio. Reaction conditions: 360 °C, 6 MPa, LHSV = $1 h^{-1}$, H₂/oil ratio 1000 : 1.

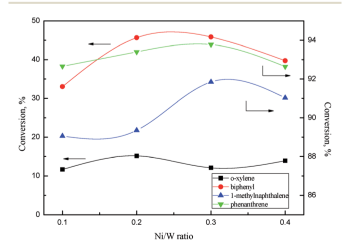


Fig. 11 Aromatics conversions as a function of Ni/W ratio. Reaction conditions: 360 °C, 6 MPa, LHSV = 1 h⁻¹, H₂/oil ratio 1000 : 1.

and length of slabs strongly affected the hydrogenation activity of the NiW catalyst. Yao $et~al.^{30}$ reported that more diaromatics were hydrogenated into monoaromatics over NiMo sulfide with larger number of slab layers. Wang $et~al.^{31}$ and Vradman $et~al.^{32,33}$ believed that multilayered structure facilitated the π adsorption of aromatic ring on slabs, resulting in the high hydrogenation activity.

3.2.3 Product composition. After hydrotreatment, the model mixtures simulating LTCTO were transformed from dark brown into transparent. Table 8 presents the hydrotreatment product composition over Ni₆W₂₀ catalyst. As shown in Table 8, the product was mainly composed of cycloalkane, partially saturated diaromatics and triaromatics, and cetane. Methylcyclohexane, which was the product of o-cresol HDO, had the highest content in monocyclic compounds. The content of methyltetralin was much higher than methyldecalin, the other product of methylnaphthalene, indicating that complete saturation of diaromatics could not be accomplished. Though exhibiting the highest conversion, phenanthrene was hydrogenated into many product compounds. Octahydrophenanthrene had the highest content among phenanthrene hydrogenating products, while tetrahydrophenanthrene had the lowest. This result revealed that compounds with one unsaturated aromatic ring were the most stable hydrogenating compounds under this reaction condition.

3.3 NiWP catalysts activity

By investigating a series of catalysts with different Ni/W ratio, we thought the catalyst with Ni/W ratio of 0.3 had best hydrotreating activity considering the heteroatom removal and aromatics hydrogenation. Thus we fixed the nickel and tungsten content in NiWP/Al $_2$ O $_3$ at 6 wt% and 20 wt% and further investigated the influence of phosphorus content on hydrotreatment activity.

3.3.1 The effect of phosphorus content on heteroatom removal. The HDO, HDS, and HDN conversions as a function of the phosphorus content of NiWP catalysts are displayed in Fig. 12. The HDO, HDS, and HDN conversions over NiWP-0.5 and NiWP-0.8 were similar with those of Ni_6W_{20} , in which no

phosphorus was added. As the phosphorus content on NiWP catalysts further increased, HDO conversion went down dramatically, while HDN conversion went up slightly. However, HDS conversion increased to a maximum with NiWP-1.2 and decreased quickly with NiWP-1.5. Atanasova *et al.*³⁴ also reported the wave-shape change of thiophene HDS conversion with the increase of phosphorus content added in NiW/Al₂O₃.

Across the range of phosphorus content on catalysts, HDO, HDS, and HDN conversion showed different trends. Additionally, the corresponding phosphorus contents with the highest conversion of HDO, HDS, and HDN were not the same. This was probably due to the difference of HDO, HDS, and HDN reaction networks, which caused many active sites on catalysts to be responsible for heteroatom removal, rather than one or two sites. The addition of phosphorus might increase a certain kind of active site, thus increasing the activity of the corresponding reaction. However the reaction networks of HDO, HDS and HDN were combination of many kinds of reactions. The change of

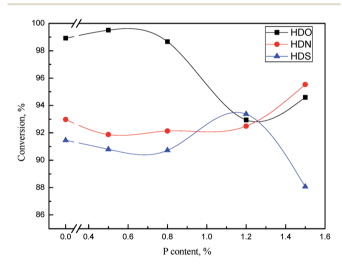


Fig. 12 HDO, HDS, and HDN conversions as a function of phosphorus content. Reaction conditions: 360 °C, 6 MPa, LHSV = 1 h^{-1}, H_2/oil ratio 1000 : 1.

Table 8 Hydrotreating product composition over Ni₆W₂₀^a

GC retention time/min	Compounds	Content/wt%	GC retention time/min	Compounds	Content/wt%
-	•			•	
3.332	Methylcyclohexane	12.48	18.462	Tetrahydroquinoline	0.40
3.604	Toluene	2.01	18.898	1-Methylnaphthalene	2.55
3.867	Ethylcyclohexane	0.31	19.245	Phenylcyclohexane	4.6
4.194	Ethylbenzene	0.26	21.635	Biphenyl	6.99
5.206	<i>O</i> -Xylene	8.6	25.184	Tetradecahydrophenanthrene	2.39
5.813	Propylcyclohexane	3.4	29.418	Butyltetralin	0.24
6.211	Propylbenzene	0.39	31.548	Cetane	22.52
6.392	Butylcyclohexane	0.29	34.972	Octahydrophenanthrene	3.77
6.812	Butylbenzene	0.29	35.393	DBT	0.12
8.665	<i>O</i> -Cresol	0.19	35.707	Tetrahydrophenanthrene	1.59
11.444	Methyldecalin	3.34	36.123	Phenanthrene	0.76
18.372	Methyltetralin	19.5			

^a Reaction conditions: 360 °C, 6 MPa, LHSV = 1 h⁻¹, H₂/oil ratio 1000 : 1.

catalytic activity caused by the phosphorus content exhibited three different trends during the removal of three kinds of heteroatoms. When the phosphorus content was higher than 1 wt%, the HDN and HDS activity was relatively high while the HDO activity was low. The content of phenols in LTCTO were much higher than that of N-containing compounds and S-containing compounds, thus the phosphorus content should be lower than 1 wt% considering the total removal rate of heteroatoms.

3.3.2 The effect of phosphorus content on aromatics hydrogenation. Fig. 13 shows the conversions of four aromatic compounds over NiWP-*x* as a function of phosphorus content on catalysts. From Fig. 13, all aromatics conversion first increased, and then decreased with the increase of phosphorus content, showing a large peak. NiWP-0.8 showed the best performance among the NiWP and Ni₆W₂₀ catalysts. A 5% improvement on *o*-xylene and biphenyl conversion was observed for NiWP-0.8. The conversions of diaromatics and triaromatics decreased with increased addition of phosphorus, and were even lower than the catalyst without phosphorus.

As revealed in Table 6, the acid concentrations on catalysts were impacted by the addition of phosphorus. Fig. 14 shows the relation of the aromatics conversion over NiWP and Ni₆W₂₀ catalysts and the medium acid concentration on catalysts. It can be clearly seen from Fig. 14 that the conversions of monoaromatics, diaromatics, and triaromatics were all linear with the medium acid concentration on catalysts. NiWP-0.8 had the highest medium acid concentration (see Table 6), showing the best performance on aromatic hydrogenation. The results indicated that the medium acid site on the NiW catalysts was probably the active site for the hydrogenation of aromatic ring. The conversion of aromatics and other compounds, whose reaction network included a rate-controlling step of aromatic ring hydrogenation, would be improved with the increase of medium acid concentration. Therefore, increased medium acid concentration was valuable to increasing the activity of hydrotreated LTCTO, since aromatic compounds were dominant in the coal tar oil. With regard to NiW sulfide catalysts for LTCTO

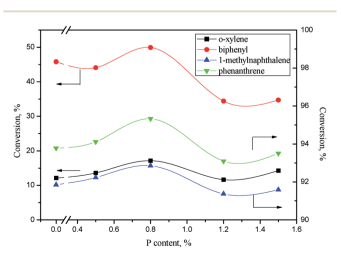


Fig. 13 Aromatics conversions as a function of phosphorus content. Reaction conditions: $360\,^{\circ}\text{C}$, $6\,\text{MPa}$, LHSV = $1\,\text{h}^{-1}$, H₂/oil ratio 1000:1.

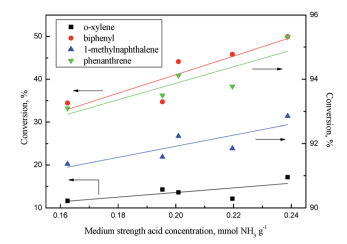


Fig. 14 Aromatics conversions as a function of medium strength acid concentration on catalysts. Reaction conditions: 360 °C, 6 MPa, LHSV = $1 \, h^{-1}$, H_2/oil ratio 1000 : 1.

hydrotreatment, performance could be improved by addition of phosphorus with an optimal content of 0.8 wt%.

3.3.3 Activity comparison between NiWP-0.8 commercial catalyst. Fig. 15 shows the comparison of compounds conversion between NiWP-0.8 and commercial catalyst. From Fig. 15, it can be seen that the conversion of oxylene and biphenyl on commercial catalyst was superior to that of NiWP-0.8, indicating the improved performance of commercial catalyst on the hydrogenation of light aromatics. However, the self-prepared NiWP-0.8 showed a much higher HDS and HDN activity than commercial one. During the hydroprocessing of LTCTO, high HDS and HDN activity was required to produce clean fuel oil, because the S and N content in the feedstock was very high. Furthermore, deactivation would be observed on the catalyst in subsequent reactors due to the Scontaining and N-containing compounds which were not

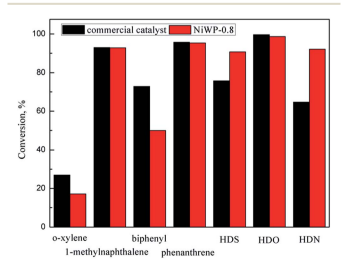


Fig. 15 Activity comparison between NiWP-0.8 and commercial catalyst. Reaction conditions: 360 °C, 6 MPa, LHSV = 1 h^{-1} , H₂/oil ratio 1000 : 1.

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removed in the hydrotreatment process.35 Therefore, NiWP-0.8, with its high HDS and HDN activity, was a promising catalyst for the hydrotreatment of LTCTO.

Conclusions

The hydrotreatment performance of eight model compounds that simulate LTCTO using NiW/Al2O3 and NiWP/Al2O3 catalysts was examined. HDO, HDS, and HDN conversion showed different trends when Ni/W ratio changed since main pathways in their reaction networks were not the same. Higher Ni/W ratios facilitated the reduction of metal oxides on the catalysts. Aromatic hydrogenation conversions on Ni₄W₂₀ and Ni₆W₂₀ were higher than that on Ni₂W₂₀ and Ni₈W₂₀ because shorter and thicker multilayer slabs were formed on Ni₄W₂₀ and Ni₆W₂₀ samples. The addition of phosphorus changed the acid concentrations and acid types on the catalysts. NiWP-0.8 exhibited the best hydrotreatment performance among all catalysts, with a 5% improvement of monoaromatics and diaromatics conversions than the catalysts without phosphorus. Further addition of phosphorus would inhibit the reduction of catalyst precursor and decrease the acid concentration and BET surface area of the catalysts. The linear relation between aromatics conversion and the medium acid concentration on catalysts suggested that medium acid was probably the active site for the hydrogenation of aromatic rings. NiWP-0.8 showed higher HDS and HDN conversion than a commercial catalyst, indicating that NiWP-0.8 was a promising catalyst for LTCTO hydrotreatment.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 Z. Guo, Q. Wang, M. Fang, Z. Luo and K. Cen, Appl. Energy, 2014, 113, 1301-1314.
- 2 T. Kan, H. Wang, H. He, C. Li and S. Zhang, Fuel, 2011, 90, 3404-3409.
- 3 K. Tao, X. Sun, H. Wang, C. Li and U. Muhammad, Energy Fuels, 2012, 26, 3604-3611.
- 4 J. Kusy, L. Andel, M. Safarova, J. Vales and K. Ciahotny, Fuel, 2012, 101, 38-44.
- 5 D. Li, Z. Li, W. Li, Q. Liu, Z. Feng and Z. Fan, J. Anal. Appl. Pyrolysis, 2013, 100, 245-252.
- 6 H. Wang, Y. Cao, D. Li, U. Muhammad, C. Li, Z. Li and S. Zhang, J. Renewable Sustainable Energy, 2013, 5, 13101.
- 7 P. Na, D. Cui, R. Li, S. Quan, K. H. Chung, H. Long, Y. Li, Y. Zhang, S. Zhao and C. Xu, *Energy Fuels*, 2012, 26, 5719–5728.

- 8 R. Wandas, J. Surygala and E. Śliwka, Fuel, 1996, 75, 687-694.
- 9 J. L. Lemberton, M. Touzeyidio and M. Guisnet, Appl. Catal., 1989, 54, 91-100.
- 10 J. L. Lemberton, M. Touzevidio and M. Guisnet, Appl. Catal., 1989, 54, 101-109.
- 11 J. L. Lemberton, M. Touzeyidio and M. Guisnet, Appl. Catal., A, 1991, **79**, 115–126.
- 12 Y. G. Wang, H. Y. Zhang, P. Z. Zhang, D. P. Xu, K. Zhao and F. J. Wang, J. Fuel Chem. Technol., 2012, 40, 12014.
- 13 H. Y. Zhang, Y. G. Wang, P. Z. Zhang, X. C. Lin and Y. F. Zhu, J. Fuel Chem. Technol., 2013, 41, 1085-1091.
- 14 W. Cui, W. Li, R. Gao, H. Ma, D. Li, M. Niu and X. Lei, Energy Fuels, 2017, 31, 3768-3783.
- 15 W. H. Wiser, S. Singh, S. A. Qader and G. R. Hill, Prepr. Pap. -Am. Chem. Soc., Div. Fuel Chem., 1969, 13(4), 350-357.
- 16 R. Rosal, F. V. Diez and H. Sastre, Ind. Eng. Chem. Res., 1992, 31, 1007-1012.
- 17 M. J. Girgis and B. C. Gates, *Ind. Eng. Chem. Res.*, 1994, 33(5), 1098-1106.
- 18 M. J. Girgis and B. C. Gates, Ind. Eng. Chem. Res., 1994, 33(10), 1098-1106.
- 19 M. Sun, X. X. Ma, Q. X. Yao, R. C. Wang, Y. X. Ma, G. Feng, J. X. Shang, L. Xu and Y. H. Yang, Energy Fuels, 2015, 25, 1140-1145.
- 20 C. Li and K. Suzuki, Resour., Conserv. Recycl., 2010, 54, 905-915.
- 21 W. Tang, M. Fang, H. Wang, P. Yu, Q. Wang and Z. Luo, Chem. Eng. J., 2014, 236, 529-537.
- 22 A. Duan, Z. Gao, Q. Huo, C. Wang, D. Zhang, M. Jin, G. Jiang, Z. Zhao, H. Pan and K. Chung, Energy Fuels, 2010, 24, 796-803.
- 23 X. Zhu, S. Liu, Y. Song and L. Xu, Appl. Catal., A, 2005, 288, 134-142.
- 24 S. Qu, G. Liu, F. Meng, L. Wang and X. Zhang, Energy Fuels, 2011, 25, 2808-2814.
- 25 M. Göhlich, W. Reschetilowski and S. Paasch, Microporous Mesoporous Mater., 2011, 142, 178-183.
- 26 J. Horáček, G. Šťávová, V. Kelbichová and D. Kubička, Catal. Today, 2013, 204, 38-45.
- 27 D. Valencia and T. Klimova, Catal. Commun., 2012, 21, 77-81.
- 28 O. Y. Gutiérrez, A. Hrabar, J. Hein, Y. Yu, J. Han and J. A. Lercher, J. Catal., 2012, 295, 155-168.
- 29 Y. Saih and K. Segawa, Appl. Catal., A, 2009, 353, 258-265.
- 30 S. Yao, Y. Zheng, S. Ng, L. Ding and H. Yang, Appl. Catal., A, 2012, 435-436, 61-67.
- 31 L. Wang, B. Shen, F. Fang, F. Wang, R. Tian, Z. Zhang and L. Cui, Catal. Today, 2010, 158, 343-347.
- 32 L. Vradman, M. V. Landau and M. Herskowitz, Fuel, 2003, 82,
- 33 L. Vradman, M. V. Landau, M. Herskowitz, V. Ezersky, M. Talianker, S. Nikitenko, Y. Koltypin and A. Gedanken, J. Catal., 2003, 213, 163-175.
- 34 P. Atanasova, T. Tabakova, C. Vladov, T. Halachev and A. L. Agudo, Appl. Catal., A, 1997, 161, 105–119.
- 35 P. M. Mortensen, J. D. Grunwaldt, P. A. Jensen, K. G. Knudsen and A. D. Jensen, Appl. Catal., A, 2011, 407, 1-19.