Trimetallic Hydrotreating Catalysts CoMoW/Al₂O₃ and NiMoW/Al₂O₃ Prepared on the Basis of Mixed Mo-W Heteropolyacid: Difference in Synergistic Effects

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Abstract—Trimetallic CoMo₃W₉/Al₂O₃ catalyst is prepared using the Keggin structure mixed heteropolyacid H₄SiMo₃W₉O₄₀ and cobalt citrate. CoMo₁₂/Al₂O₃ and CoW₁₂/Al₂O₃ catalysts based on H₄SiMo₁₂O₄₀ and H₄SiW₁₂O₄₀, respectively, are synthesized as reference samples. Sulfided catalysts are analyzed by high-resolution transmission electron microscopy and X-ray photoelectron spectroscopy. Catalytic properties are investigated in the co-hydrotreatment of dibenzothiophene (DBT) and naphthalene in a flow unit. It is shown that the catalytic activity in both DBT hydrodesulfurization and naphthalene hydrogenation (HYD) decreases in the following sequence: CoMo₁₂/Al₂O₃ > CoMo₃W₉/Al₂O₃ > CoW₁₂/Al₂O₃, and it correlates with the degree of promotion of active-phase particles by cobalt atoms. A comparison with the published data available for Ni-promoted catalysts makes it possible to reveal the general regularity for bi- and trimetallic Co(Ni)-Mo(W)S catalysts: the use of mixed Mo-W H₄SiMo₃W₉O₄₀ heteropolyacid instead of monometallic H₄SiW₁₂O₄₀ causes an increase in the degree of promotion of MoWS₂ crystallite edges for the series of catalysts is much more active than the CoMo₃W₉/Al₂O₃ counterpart. Possible reasons behind the found features are discussed.

Keywords: hydrotreating, CoMoWS, NiMoWS, dibenzothiophene, naphthalene **DOI:** 10.1134/S0965544118140104

A deep hydrotreating of hydrocarbon feedstock becomes all the more urgent in connection with the need to produce environmentally friendly fuels from heavy oil fractions and residues and because of the involvement of high-sulfur oils in refining. Therefore, the design of new catalytic materials is aimed at increasing their hydrogenation activity necessary to improve the energy efficiency of hydrocatalytic processes owing to reduction in operational costs. $Co(Ni)Mo(W)/Al_2O_3$ catalysts have been used in hydrogenation processes for more than eight decades [1-3]. The active phase in these catalysts is Co(Ni)Mo(W)S-type structures which are formed during catalyst activation (sulfiding); these are nanosized MoS₂ or WS₂ crystallites decorated by Co and/or Ni promoter atoms at the edges [4].

Catalysts based on WS_2 feature as a rule high activity in hydrogenation reactions and possess an increased stability [5]. The improvement of promoted Co(Ni)W catalyst systems is at the focus of ever growing attention [6, 7]. Previous studies showed that Co(Ni)Mo catalysts exhibit a higher catalytic activity in hydrodesulfurization (HDS) reactions compared with their Co(Ni)W analogs [3, 8-12]. In accordance with Vissenberg et al. [13], one of the reasons behind a low HDS activity of Co(Ni)W catalysts is a great difference in sulfiding conditions for supported oxide precursors, in particular, overly high sulfiding temperatures for W oxides compared with Co or Ni oxides. Therefore, during activation of Co(Ni)W catalysts, the sulfiding of Co(Ni) promoter atoms occurs initially to generate CoS_r (NiS_r) massive variable-composition sulfides inactive in catalysis and only then are WS₂ crystallites formed. Nevertheless, the authors of [13, 14] disclosed a high promoting effect of Ni related to the post-formation of NiWS sites via the redistribution of NiS_x particles, namely, the migration of a portion of promoter atoms and their localization at tungsten sulfide edges [13, 14]. Much less data are available on CoW catalysts, although attempts to gain insight

Catalyst		Content, wt %		Textural characteristics of sulfide samples			
	MoO ₃	WO ₃	CoO	surface area S_{BET} , m ² /g	pore volume $V_{\rm p}$, cm ³ /g	pore diameter, nm	
CoMo ₁₂ /Al ₂ O ₃	18.0	_	4.7	291	0.55	3.8/8.4	
CoW ₁₂ /Al ₂ O ₃	—	26.2	4.2	265	0.50	3.8/8.4	
CoMo ₃ W ₉ /Al ₂ O ₃	4.2	20.1	4.3	288	0.56	3.8/8.4	

Table 1. Composition and textural characteristics of $CoMo(W)/Al_2O_3$ catalysts

into causes of their low catalytic activity and to eliminate them to date do not cease [15-17].

One of the known methods to increase the activity of hydrotreating catalysts involves the incorporation of complexing agents into their composition [18–20]. Owing to the formation of stable complexes with Ni (Co), the use of chelatons makes it possible to decelerate the preliminary sulfiding of promoters or to synchronize it with the sulfiding of main metals (Mo and W). As a result, selectivity for formation Co(Ni)Mo(W)S active-phase particles grows.

Another method to enhance catalyst activity in HDS and HYD reactions includes the application of trimetallic Ni(Co)MoWS catalysts, in which a portion of tungsten atoms is replaced with molybdenum or vice versa [21-26]. The feasible synergistic effect in the co-use of Ni, Mo, and W was predicted by the density function theory (DFT) [23, 27]. As was shown in [1, 28, 29], the use of the Keggin structure mixed Mo-W heteropolycompounds as precursors of MoW/Al_2O_3 and NiMoW/Al₂O₃ hydrotreating catalysts, in which one or three W atoms of the 12 are replaced with Mo, ensures the spatial proximity of Mo and W atoms. This enables preparation of the mixed MoWS₂ phase; its composition and structure are ascertained using a scanning transmission electron microscope and a high-angle annular dark field (HAADF) detector. Given this, the degree of sulfiding tungsten grows and, as a consequence, the amount of active-phase particles increases appreciably. Moreover, the degree of promotion of active-phase edge sites by nickel on the NiMoW catalyst is $\sim 30\%$ higher than that on the NiW/Al₂O₃ catalyst [1]. Trimetallic CoMoWS hydrotreating catalysts are attracting all the more growing attention from researchers [15, 26, 30, 31], although the data on their activity are ambiguous. For example, in accordance with [17, 30], massive NiMoWS catalysts are much more active than CoMoWS analogs. However, as was reported recently by Pawelec et al. [15], owing to increased acidity (due to the partial replacement of Mo atoms with W ones), the CoMoW/Al₂O₃-TiO₂ catalyst shows a considerably higher activity in the hydrodesulfurization of DBT; this parameter is even higher than the activity of the CoMo/Al₂O₃ commercial reference sample.

The aim of this work was to explore the effect of Co promotion for the CoMoW/Al₂O₃ trimetallic catalyst. For this purpose, catalysts were synthesized using mixed $H_4SiMo_3W_9O_{40}$ heteropolyacid (HPA) and cobalt citrate as precursors of the active phase and bimetallic CoMo/Al₂O₃ and CoW/Al₂O₃ reference samples on the basis of corresponding monometallic $H_4SiMo_{12}O_{40}$ (SiMo₁₂HPA) and $H_4SiW_{12}O_{40}$ (SiW₁₂HPA) HPAs. The synthesized catalysts in the activated (sulfide) state were studied by high-resolution transmission electron microscopy (HRTEM) and X-ray photoelectron spectroscopy (XPS). Using these methods, the composition and structure of activephase nanosized particles were ascertained and the catalytic behavior of the catalysts in dibenzothiophene HDS and naphthalene HYD model reactions was explored.

EXPERIMENTAL

Precursors of the active phase of catalysts were Keggin structure heteropolyacids. Monometallic $H_4SiMo_{12}O_{40}$ and $H_4SiW_{12}O_{40}$ and mixed $H_4[SiMo_3W_9O_{40}]$ HPAs were synthesized as described in [32–34]. The composition and structure of HPAs were studied by IR and Raman spectroscopy, X-ray diffraction, and EXAFS.

CoMo/Al₂O₃, CoW/Al₂O₃, and CoMo₃W₉/Al₂O₃ catalysts with the same surface loading of metals Mo + W ~ 4 at/nm² were prepared by the single incipient wetness impregnation of the support (specific surface area, 275 m²/g; specific pore volume, 0.9 cm³/g) with the aqueous solution of corresponding HPAs and cobalt citrate. After impregnation, the samples were dried at 110°C. The content of metals in the catalysts was monitored using an EDX800HS X-ray fluorescence analyzer (Shimadzu). The compositions of the synthesized catalysts are presented in Table 1.

The textural characteristics of the catalyst samples were investigated by low-temperature nitrogen adsorption on a Quantachrome Autosorb-1 adsorption porosimeter. The specific surface area was calculated according to the Brunauer–Emmett–Teller (BET) model for $P/P_0 = 0.05-0.3$. The total pore volume and pore size distribution were calculated from the desorption curve in terms of the Barrett–Joyner– Halenda model.

The XPS studies of sulfide catalysts were conducted on a Kratos Axis Ultra DLD spectrometer using Al K_{α} radiation (hv = 1486.6 eV). The binding energy (E_b) scale was preliminarily calibrated against position of the peaks of core levels Au4 $f_{7/2}$ (84.0 eV) and $Cu2p_{3/2}$ (932.67 eV). The samples were applied on a double-sided nonconducting adhesive tape. The charging effect arising during the photoemission of electrons was minimized through irradiation of the sample surface by low-energy electrons of a special source (flood gun). Calibration was made against the C1s line (284.8 eV) of carbon occurring on the surface of the catalyst. The energy step was 1 eV for the survey spectrum and 0.1 eV for individual lines C1s, A12p, S2p, Mo3d, and W4f. The decomposition of the Mo3d and W4f lines was conducted in accordance with parameters described in [35-37]. For all sulfide catalysts, the relative concentrations of $W^{6+}(Mo^{6+})$ and Co²⁺ particles in the oxide environment, oxysulfides WS_xO_v (MoS_xO_v), and sulfides WS₂ (MoS₂), CoW(Mo)S, Co_9S_8 were determined. For example, the relative concentration of WS₂ particles was calculated through the following equation:

$$[WS_2](\%) = \frac{A_{WS_2}}{A_{WS_2} + A_{WS_2O_y} + A_{W^{6+}}} \times 100, \qquad (1)$$

where A_X is the area under the peak of particles x.

The amount of cobalt in the Co(Mo)WS phase was determined via the following equation:

$$c_{\rm Co(Mo)WS} = [\rm CoMo(W)S]c_{\rm Co}, \qquad (2)$$

where $C_{\rm Co}$ is the total content of Co measured using XPS, wt %.

The degree of promotion of active-phase crystallites was calculated by the following equation:

$$\left(\frac{\text{Co}}{\text{Mo} + W}\right)_{\text{slab}} = \frac{c_{\text{Co(Mo)WS}}}{c_{\text{W(Mo)Sp}}},$$
(3)

where C_X is the absolute concentration of Co(Mo + W) in Co(Mo)WS particles, at %.

The degree of promotion of the edges of activephase crystallites was determined in accordance with [37] through the following equation:

$$\left(\frac{\text{Co}}{\text{Mo} + \text{W}}\right)_{\text{edge}} = \frac{(\text{Co}/(\text{Mo} + \text{W}))_{\text{slab}}}{D},$$
 (4)

where *D* is the dispersity of the active phase calculated using the HRTEM data.

The catalysts were analyzed by HRTEM on a Tecnai G2 20 instrument equipped with a LaB6 cathode at an accelerating voltage of 200 kV. The average length of particles (\overline{L}) and the number of NiMo(W)S layers per stack (\overline{N}) were determined taking into account on the order of 400-600 particles occurring in 10-15 different areas of the catalyst surface. The dispersity (*D*) of active-phase particles was calculated in terms of the Kasztelan hexagonal model [38] using the following equation:

$$D = \frac{M_{\rm e} + M_{\rm c}}{M_{\rm T}} = \frac{\sum_{i=1..t} 6n_i - 6}{\sum_{i=1..t} 3n_i^2 - 3n_i + 1},$$
 (5)

where M_e is the number of Mo(W) atoms on the edges of middle crystallite CoMo(W)S, M_c is the number of Mo(W) atoms on the edges of middle crystallite CoMo(W)S, M_T is the total number of Mo(W) atoms in the middle particle of the active phase, *n* is the number of molybdenum atoms along one side of the MoS₂ crystallite defined by its length, and *t* is the total number of layers in the crystallite calculated from the HRTEM data.

The catalytic properties of the synthesized samples were examined in dibenzothiophene (0.86 wt %)hydrodesulfurization and naphthalene (3 wt %) hydrogenation reactions during the co-hydrotreating of the model mixture in toluene in the flow unit equipped with a microreactor. A steel reactor was charged with the catalyst (0.25-0.5 mm fraction, 0.2 g) diluted with carborundum at a ratio of 1 : 4. Tests were run under the following conditions: temperature, 280°C; pressure, 3 MPa; a feed space velocity, 40 h⁻¹; and H₂/feedstock = 500 nL/L. Before testing, the catalysts were activated in situ by holding with a mixture of dimethyl sulfide (2 wt % sulfur) in decane at 240°C for 10 h and at 340°C for 8 h. The products were identified by gas-liquid chromatography on a Kristall-5000 chromatograph. The steady-state activity of the test samples was observed after continuous testing for 7–10 h.

The rate constants for DBT hydrodesulfurization and naphthalene hydrogenation were determined under the assumption that these are first-order reactions and were calculated by the equations

$$k_{\text{HDS}} = -\frac{F_{\text{DBT}}}{m} \ln(1 - x_{\text{DBT}}) \text{ and}$$

$$k_{\text{HYD}} = -\frac{F_{\text{Naft}}}{m} \ln(1 - x_{\text{Naft}}),$$
(6)

where k_{HDS} and k_{HYD} are the rate constants (mol/(g h)) of DBT hydrodesulfurization and naphthalene hydrogenation, respectively; x_{DBT} and x_{Naph} are the conversions (%) of DBT and naphthalene; F_{DBT} and F_{Naph} are the molar consumptions (mol/h) of reagents; and *m* is the catalyst weight (g).

Taking into consideration that the DBT hydrodesulfurization occurs via two routes, catalyst selectivity was estimated using Eq. (7) as the ratio between the total concentration of the products of reaction following the dibenzothiophene hydrogenation route to form

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Fig. 1. TEM images of sulfided catalysts: (a) $CoMo_{12}/Al_2O_3$, (b) CoW_{12}/Al_2O_3 , and (c) $CoMo_3W_9/Al_2O_3$.

tetrahydrodibenzothiophene (THDBT), bicyclohexyl (BCH), and cyclohexylbenzene (CHB) and the concentration of diphenyl (DP) formed via the direct hydrodesulfurization (DS) route:

$$S_{\rm HYD/DS} = \frac{C_{\rm THDBT} + C_{\rm BCH} + C_{\rm CHB}}{C_{\rm DP}},$$
 (7)

where C_x are the concentrations of corresponding reaction products (mol %).

RESULTS AND DISCUSSION

For the test catalysts in the sulfide form, the specific surface area changed insignificantly within $265-291 \text{ m}^2/\text{g}$ and the pore volume was $\sim 0.50-0.55 \text{ cm}^3/\text{g}$. The bimodal distribution of pores with an effective diameter of 3.8 and 8.4 nm is related to coke particles formed during the sulfiding of samples containing citric acid and the effective pore size of the initial support (Table 1).

The TEM images of the sulfided catalysts are shown in Fig. 1. Black threadlike bands correspond to the layers of $Mo(W)S_2$ crystallites with a characteristic interplanar distance of about 0.65 nm [39].

The length of active-phase particles decreases in the sequence $CoW_{12}/Al_2O_3 > CoMo_{12}/Al_2O_3 > CoMo_3W_9/Al_2O_3$ (Table 2). The dispersity of particles varies within 0.31–0.42; the CoW_{12}/Al_2O_3 sample exhibits the minimum dispersity among the promoted catalysts.

The relative contents of Mo, W, and Co particles on the surface of sulfided CoMo(W)/Al₂O₃ catalysts and the degrees of promotion of Mo(W)S₂ crystallites according to the XPS data are summarized in Table 3. All of the catalysts possess comparable sulfiding depths. However, the content of Co in the Co(Mo)WS phase for the CoMo₁₂/Al₂O₃ sample was higher (0.92 wt %) compared with CoMo₃W₉/Al₂O₃ (0.51 wt %) and $\text{CoW}_{12}/\text{Al}_2\text{O}_3$ (0.26 wt %) catalysts. Probably, these results may be explained by the fact that for tungsten oxide the temperature of sulfuding is much higher than that for molybdenum and cobalt oxides [40]. As a result, for the CoW_{12}/Al_2O_3 sample, the proportion of MoS_2 is higher than that of WS_2 and the content of individual Co_9S_8 is maximal (79 rel. %). For the mixed catalyst, the selectivity for formation of the Co(Mo)WS active phase and the degree of promo-

Catalyst	Average length \overline{L} , nm	Average number of Mo(W)S ₂ layers in crystallite \overline{N}	Dispersity of Mo(W)S ₂ particles D^1
CoMo ₁₂ /Al ₂ O ₃	3.0	1.6	0.38
CoW ₁₂ /Al ₂ O ₃	3.8	2.0	0.31
CoMo ₃ W ₉ /Al ₂ O ₃	2.7	1.5	0.42

Table 2. Morphology of active-phase particles of CoMo(W)/Al₂O₃ catalysts

¹ The dispersity of $Mo(W)S_2$ particles was calculated from the HTEM data using Eq. (5).

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Table 3.	Content of molybdenum,	tungsten, and nicke	l particles on the s	surface of sulfide ($CoMo(W)/Al_2O_3 c$	atalysts and
degree o	f promotion of crystallites	according to the XPS	S data			

Catalysts	Mo content, rel. %			W content, rel. %		Co content, rel. %						
	MoS_2	MoS_xO_y	Mo^{6+}	^z SM	WS _x O _y	-9M	Co(Mo)WS	Co_9S_8	Co^{2+}	C _{Co(Mo)WS} , wt % ¹	$\left(\frac{Co}{Mo+W}\right)_{slab}^2$	$\left(\frac{\text{Co}}{\text{Mo} + \text{W}}\right)_{\text{edge}}^{3}$
CoMo ₁₂ /Al ₂ O ₃	80	11	9	_	_	_	51	29	20	0.92	0.33	0.88
CoW ₁₂ /Al ₂ O ₃	—	_	_	68	8	24	8	79	13	0.26	0.12	0.40
CoMo ₃ W ₉ /Al ₂ O ₃	86	8	24	70	8	22	12	76	12	0.51	0.20	0.47

 1 Co content in the Co(Mo)WS phase was calculated from the XPS data using Eq. (2).

 2 The degree of promotion of Co/(Mo+W) particles was calculated from the XPS data through Eq. (3).

³ The degree of promotion of Co/(Mo+W) particle edges was calculated from XPS and TEM data through Eq. 4.

Table 4. Catalytic behavior of $CoMo(W)/Al_2O_3$ catalysts in DBT hydrodesulfurization and naphthalene dehydrogenation

Catalyst	Conversion	n, %	Rate constants, ×10	Selectivity Superco	
	dibenzothiophene	naphthalene	$k_{ m HDS}$	$k_{ m HYD}$	Servering SHYD/DS
CoMo ₁₂ /Al ₂ O ₃	46.8	6.2	32.2	2.3	0.10
CoW ₁₂ /Al ₂ O ₃	6.6	1.1	3.2	0.4	0.01
CoMo ₃ W ₉ /Al ₂ O ₃	22.0	3.3	11.2 (10.5)*	1.2 (0.9)*	0.07

* The values of rate constants calculated by the additivity method are shown in parentheses.

tion of active-phase particles grow compared with CoW_{12}/Al_2O_3 .

The data on the catalytic activity of the synthesized catalysts are presented in Table 4.

The conversion of reagents was varied from 6.6 to 46.8%. The lowest activity was exhibited by the CoW₁₂/Al₂O₃ catalyst in both DBT hydrodesulfurization and naphthalene hydrogenation. Among the tested catalysts the highest activity was observed for $CoMo_{12}/Al_2O_3$. The selectivity of the dibenzothiophene preliminary hydrogenation route for the tested catalysts varied within 0.01-0.10 and was lower compared with monometallic Mo_{12}/Al_2O_3 and W_{12}/Al_2O_3 catalysts (0.7 and 0.9, respectively [28]). This fact provides evidence for the generation of new promoted sites. In addition, the rates constants for HDS and HYD over the $CoMo_3W_9/Al_2O_3$ catalyst were calculated by the additivity method. The experimental value for DBT hydrodesulfurization was slightly above the calculated one (by 6%); for naphthalene hydrogenation, by 25%. The highest rates constants for DBT hydrodesulfurization and naphthalene hydrogenation were attained in the case of the $CoMo_{12}/Al_2O_3$ catalyst.

Figure 2 shows the dependence of the rate constant of dibenzothiophene hydrodesulfurization on the con-

tent of Co(Ni) in particles of the Co(Ni)(Mo)WS active phase for catalysts promoted by Co and Ni. At a comparable content of the promoter in active-phase particles on NiMo₃W₉/Al₂O₃ and CoMo₃W₉/Al₂O₃ samples, the catalyst promoted by nickel showed a higher activity. Moreover, for (Ni)CoMo₁₂/Al₂O₃ catalysts, activity analogous to that of Mo₃W₉/Al₂O₃ may be achieved only at a higher promoter content in the Co(Ni)(Mo)WS phase (Fig. 3). This synergism observed when Ni was used as a promoter for the mixed NiMoWS active phase is probably associated with a more optimum value of metal-sulfur binding energy (ΔE_{MS}) than that in the bimetallic samples and CoMoWS [23].

Analysis of the dependence of the rate constant of DBT hydrodesulfurization on the degree of promotion of edges of the (Mo)WS₂ phase (Fig. 3) makes it possible to conclude that the replacement of SiW₁₂ HPA with MoW-containing SiMo₃W₉ HPA leads to a rise in the degree of decoration of MoWS₂ crystallite edges for the series of catalysts promoted by both Co and Ni. In addition, the use of Ni entails a higher degree of promotion for edges of active-phase particles compared with Co. For the NiMo₃W₉/Al₂O₃ catalyst, the degree of promotion is commensurable with that of NiMo₁₂/Al₂O₃ and CoMo₁₂Al₂O₃ samples. These



Fig. 2. Rate constant of DBT hydrodesulfurization vs. the content of Co(Ni) in particles of the Co(Ni)(Mo)WS active phase in the catalysts promoted by Co and Ni. For the series of Ni-promoted catalysts, calculated from [1]; for W_{12}/Al_2O_3 , calculated from [28].

results may be attributed to the fact that the atoms of promoters (Co or Ni) are localized on different edges of tungsten sulfide: Ni on the metal edge (Me edge) or on both edges (Me and S edges), while Co predominantly inserts via the S edge [41–43]. A high degree of



Fig. 3. Rate constant of DBT hydrodesulfurization vs. the degree of promotion of edges of Co(Ni)(Mo)WS activephase particles in the catalysts promoted by Co and Ni. For the series of Ni-promoted catalysts, calculated from [1]; for W_{12}/Al_2O_3 , calculated from [28].

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Ni promotion may also be associated with the abovementioned mechanism of post-formation of NiWS sites via the redistribution of NiS_x particles. It is evident that, in the case of nickel, the probability of relocalization on both edges will be higher than that for cobalt on the S edge. Moreover, a higher stability of the nickel complex with citric acid compared with its cobalt counterpart [44] is favorable for the synchronous formation of the promoted NiWS active phase.

Thus, the effect of Co promotion in the $CoMo_3W_9/Al_2O_3$ mixed trimetallic catalyst has been compared with $CoMo_{12}/Al_2O_3$ and CoW_{12}/Al_2O_3 catalysts, and the influence of the nature of the promoter on the physicochemical and catalytic properties of biand trimetallic sulfide catalysts has been explored.

It has been shown that the catalytic activity in both DBT dihydrodesulfurization and naphthalene hydrogenation decreases in the following sequence: $CoMo_{12}/Al_2O_3 > CoMo_3W_9/Al_2O_3 > CoW_{12}/Al_2O_3$, and it correlates with the degree of promotion of active-phase particles by cobalt atoms.

The general regularity which makes itself evident in that the use of mixed $H_4SiMo_3W_9O_{40}$ HPA instead of monometallic $H_4SiW_{12}O_{40}$ HPA causes an increase in the degree of decoration of edges of MoWS₂ crystallites for the series catalysts promoted by both Co and Ni has been established. It is shown that the use of Ni as a promoter facilitates a higher degree of promotion of edges of active-phase particles compared with Co apparently owing to differences in the stability of citrate complexes and, hence, in the rates of their sulfiding and different tendencies toward localization on the edges of $Mo(W)S_2$ crystallites.

The activity of bimetallic (Ni)CoMo₁₂/Al₂O₃ catalysts comparable with that of trimetallic NiMo₃W₉/Al₂O₃ samples may be achieved only at a higher content of the promoter in particles of the active phase. This synergistic effect is probably provided by the metal-sulfur binding energy (ΔE_{MS}) in NiMoWS which is more optimal for catalysis than that in bimetallic CoMoS (NiMoS) counterparts.

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