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# ACTIVITY OF TRANSITION METAL SULFIDES SUPPORTED ON ALUMINA, TITANIA AND ZIRCONIA IN PARALLEL HYDRODESULFURIZATION OF 1-BENZOTHIOPHENE AND HYDROGENATION OF 1-METHYL-CYCLOHEX-1-ENE

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and hydrogenation of 1-methyl-cyclohex-1-ene (o-HYD) at 360 °C and 1.6 MPa. unconventional supports  $TiO_2$  (anatase,  $S_{BET} = 140 \text{ m}'\text{g}'$ ) and  $ZrO_2$  (baddeleyite,  $S_{BET} = 140 \text{ m}'\text{g}'$ ) Pt, and Re deposited over conventional support gamma-Al<sub>2</sub>O<sub>3</sub> ( $S_{\rm DET} = 262~{\rm m^2g^{-1}}$ ), and undoubtedly assigned by NMR spectroscopy [1]. Nevertheless, during each catalytic also ethylcyclopentene and dimethylcyclopentene isomers. All volatile isomerization 108 m<sup>2</sup>g<sup>-1</sup>) were studied in parallel hydrodesulfurization of 1-benzothiophene (HDS) Sulfided conventional transition metals Co, Ni, Mo, and W and noble metals Rh, Pd, Ir, cyclohex-lene hydrogenation (o-HYD) that proceeded together with 1-benzothiophene and methylcyclohexane, which were quantified. It was ascertained that the 1-methyl-90% the mixture of the following four compounds: 1-,3- and 4-methyl-cylohex-1-enes experiment, the model compound 1-methyl-1-cyclohex-1-ene yielded from more than chromatographic techniques including preparative gas chromatography, and were products were isolated directly from diluted reaction mixture via combination of and ZrO2 support effect (Rh/ZrO2 being an exception) and o-HYD influenced the HDS supported sulfides of Rh, Re, Pd, Pt, and Ir were highly active in HDS with low TiO2 0.4-9.3 and the o-HYD decreased the HDS activity by the factor 0.4-0.8. In contract, the and o-HYD. For these sulfides, the TiO2 and ZrO2 support effect varied by the factor [2]. Mo, W, Co, and Ni sulfided catalysts exhibited relatively low activity in both HDS found before for the HDS reaction of 1-benzothiophene without 1-methyl-cyclohex-ene the studied transition metal sulfides supported on Al2O3, TiO2 and ZrO2 which was hydrodesulfurization (HDS) did not significantly affected the HDS activity ranking of 1-Methyl-cyclohex-1-ene transformation was found to lead cyclic olefins containing activities in o-HYD. The reference bimetallic CoMo and NiMo Al<sub>2</sub>O<sub>3</sub> supported activity by the factor 0.6-1.6. Nevertheless, Rh, Re, and Pd sulfided species exhibited catalysts exhibited desirable high HDS but low o-HYD activities [3]. relatively low activities in o-HYD reaction while Ir and Pt species exhibited high

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## AI DISTRIBUTION IN TNU-9 ZEOLITE

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Zeolites are crystalline microporous aluminosilicate molecular sieves defined, ordered framework formed by corner-sharing AlO<sub>4</sub> and SiO<sub>4</sub> telru isomorphous substitution of Al atoms in the silicon framework results in the of negatively charged AlO<sub>4</sub> in the zeolite framework. This charge is balancu framework cationic species which represent active sites in a number of acid selectivity of the zeolite channel system and other zeolite properties results that zeolites represent the widest group of industrial heterogeneous catalysts,

Distribution of Al atoms in the zeolite framework determines its abillicative sites both for reactants and guest ions. This phenomenon was demons the question of FER, MOR, BEA and ZSM-5 type zeolites [1-3]. Current work zeolite [4]. Methods used to solve presented problem encompasses Vis-NIR, I NMR spectroscopy as well as introduction of probe molecules into the zeolite prepared according to carlier described methods [4]. Post synthesis muture calcined in air and converted into sodium, ammonium or cobalt balanced forms.

Analysis of prepared samples resulted in ruling out the presence of Al Al sequences reflected in the <sup>29</sup>Si MAS NMR signal. Two main Al arrangement framework thus represents: (i) single Al atoms not able to stabilize  $Co^{3t}$  | complexes and (ii) Al pairs of Al-O-(SiO)<sub>2</sub>-Al-O sequences located in one 6-ring, able to accommodate bare  $Co^{2t}$  ions. Three types of Al pairs were identification on the channel system was discussed.

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