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Nickel and Cobalt phosphide as effective catalysts for oxygen-removal of dibenzofuran: Role of contact time, hydrogen pressure and hydrogen/feed molar ratio

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Abstract

The catalytic activity of nickel- and cobalt phosphides, with a metallic loading of 5 wt.%, supported on silica was investigated in the hydrodeoxygenation reaction (HDO) of dibenzofuran (DBF) as model oxygenated compound at different contact times, H_2 pressures and H_2 /DBF molar ratios. The aim of the study was to understand the mechanism of the reaction and to study the impact of H_2 pressure and H_2 /DBF molar ratio on the reaction. The catalysts were characterized by N₂ adsorption–desorption isotherms at -196 °C, X-ray diffraction (XRD), X-ray photoelectron spectroscopy(XPS), CO-chemisorption, NH₃ Temperature-Programmed Desorption (NH₃-TPD), IR- Spectroscopy and H₂ Temperature-Programmed Desorption (H₂-TPD). The prepared catalysts were tested in the HDO reaction of DBF in a continuous-flow fixed-bed stainless steel catalytic reactor at pressures ranging from 1-30 bars at 275 °C. The results obtained indicate that the Ni₂P catalyst is more active than the CoP catalyst, converting more than 90% of the DBF at the highest contact time into oxygen-free products. The activity of both catalysts increases with increased contact time. At low contact times the intermediates tetrahydrodibenzofuran (THDBF) and hexahydrodibenzofuran (HHDBF) are observed as products, while an increment in the contact time led to the transformation of THDBF and HHDBF into O-free compounds, mainly bicyclohexane (BCH), indicating that the HDO of DBF follows the path: DBF \rightarrow HHDBF \rightarrow THDBF \rightarrow 2-CHP \rightarrow BCH. Further both Ni₂P and CoP catalysts are active at medium pressures with HDO degrees similar to those obtained at 30 bar. Ni₂P is less affected by changes in H₂/DBF ratio than CoP and the catalysts are more active at high H₂/DBF molar ratios.

Keywords: HDO; Ni₂P; CoP; Dibenzofuran; Silica; transition metal phosphides

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1. Introduction

The use of bioenergy as a renewable alternative to fossil fuels is nowadays attracting more and more attention. The reasons are quite obvious: our world is running out of oil reserves; the raw oil quality is decreasing;¹ and the world petroleum consumption continues to rise as a consequence of the world's high population growth, the industrialization and a big change in consumption pattern. Within this scenario, biofuels seem to be a potential energy substitute for fossil fuels since it is a renewable resource that could contribute to sustainable development and global environmental preservation².

Bio-oil is formed from fast pyrolysis of biomass. It retains most of the original oxygen in the feedstocks and the oxygen content varies in the range of 35-60 wt.% on a wet basis. First of all, the oxygenated compounds make the bio-oil polar and for this reason it will be non-miscible with non-polar petroleum fuels. The high oxygen content also gives the bio-oil other important undesirable properties. Hence, upgrading is necessary to use the pyrolysis-oil as a competitive substitute of petroleum fuels. Most important is a de-oxygenation step prior to co-processing^{3, 4}.

HDO is a promising future process for the up-grading of bio-oil. The HDO process requires the presence of hydrogen and a catalyst, and it normally operates at moderate temperatures, up to 450 °C and at a high pressure, 10-300 bar. The high pressure increases the reaction rate and decreases the formation of coke in the reactor. A good HDO catalyst should be highly active to remove oxygen in the bio-oil with a high yield and it should not produce coke⁵.

HDO is a process very similar to HDS and commercial catalysts, often used for traditional hydrotreating processes such as Co-MoS₂ and Ni-MoS₂ supported on O-Al₂O₃, have been the most frequently tested catalysts also for HDO reactions.⁶ However a problem with the above named catalysts is that co-feeding of H₂S is needed in order to regenerate the sulphide sites. Further the sulphidation agent must be separated from the products and the sulphur must be recovered from the H₂S. Another disadvantage is the corrosiveness of H₂S⁷.

Noble metal supported catalysts as for example Ru/C and Pt/C have also shown good results in HDO reactions. Noble metal catalysts in HDO are also active at low temperatures. However, the main disadvantages with noble metal based catalysts are their high cost and additionally they show a low resistance towards poisoning by the presence of sulphur⁸.

Alternative compositions that show good hydrogen transferring properties are metal carbides, metal nitrides, metal borides and transition metal phosphides⁹. Transition metal phosphides have

presented very interesting properties in hydrotreating reactions as they are bifunctional with acid and metallic catalytic functions. The metallic function is responsible for hydrogenation, while on the acidic sites, hydrogenolysis, dehydration, and isomerization reactions take place. The balance among these functions determines the observed catalytic activity. From FTIR studies of CO adsorption of the transition metal phosphide Ni₂P/SiO₂, it has been shown that there are enough electrons on the metal site to allow π -back bonding. This π -back bonding enables molecular activation as electrons fill antibonding orbitals and the hydrogenation activity is thereby enhanced. The acidity arises from acidic PO-H sites¹⁰.

In the last decade a series of studies have shown that MoP, WP, Fe₂P, CoP and Ni₂P are suitable for hydrodesulfurization and hydrodenitrogenation of fossil crude oils¹¹⁻¹⁵. More recently, these transition metal phosphides have also been applied as catalysts in HDO reactions of bio-oil ^{16, 17} as they show good hydrogen transfer properties. These kinds of catalysts are favored when supported on a less acidic support as silica, SBA-15 or MCM-41. Zhao et al.¹⁸ measured the activity for the HDO reaction of guaiacol over Ni₂P, Co₂P, Fe₂P, WP and MoP catalysts supported on SiO_2 in a fixed bed reactor with a molar ratio H_2 /guaiacol of 33. The results indicated that the type of metal has a significant effect on product distribution and a different involvement of metals during the hydrogen activation or the interaction of reactants with the catalyst surface were suggested. The activity of the different catalyst followed the order: Ni₂P >Co₂P >Fe₂P≈WP≈MoP. The activities of the transition metal catalysts were compared to the catalytic activity of the commercial catalyst CoMoS/Al₂O₃ and 5%Pd/Al₂O₃ The commercial 5% Pd/Al₂O₃ catalyst turned out to be more active than the metal phosphides at lower contact time, but the major product was catechol which is undesired. A. Iino et al.¹⁹ studied the HDO reaction of 2-metyltetrahydrofuran (2-MTHF) on a Ni₂P/SiO₂ catalyst with a continuous-flow quartz reactor. The activity and selectivity of this transition metal phosphide catalyst were compared to Pd/Al₂O₃ and Ru/C catalysts. The results showed that Ni₂P/SiO₂ had higher selectivity to *n*pentane and lower selectivity to lower hydrocarbons with carbon numbers less than 5 compared to Pd/Al₂O₃ and Ru/C. It was concluded that Ni₂P/SiO₂ catalyst has high HDO activity and low carbon bond cracking propensity.

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When it comes to the reaction parameters, Mortensen et al.²⁰ describe in their review of catalytic upgrading of bio-oil that a high pressure is beneficial in the HDO process. A high pressure increases the solubility of hydrogen in the oil and consequently there will be higher availability of hydrogen in the proximity to the catalyst. This increases the reaction rate and counteracts carbon deposition on the surface of the catalyst. However, a great amount of hydrogen interferes with

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the adsorption of reactant on the surface¹⁹. In the work of A. Iino et al.¹⁹ the results showed that the conversion decreases as H₂ partial pressure increases. These experiments were run under H₂ partial pressures in the range 75-418 kPa. In a recent study of T.A. Le et al.²¹ the HDO process of 2-furyl methyl ketone (FMK) catalyzed by CoP/ γ -Al₂O₃, the complete HDO of FMK into methyl cyclopentane and methane was achieved at 400 °C and at atmospheric pressure. The authors urge that positive catalytic results can be achieved even at atmospheric pressure (low pressure).

In this work, the focus is on the activity of nickel phosphide catalyst (Ni₂P) and cobalt phosphide catalyst (CoP), both supported on silica and tested in the HDO of dibenzofuran (DBF). In a previous work the HDO of DBF on Ni₂P with different metallic loadings and P/Ni molar ratios was studied. In the present paper an evaluation of the impact of the experimental parameters on the catalytic response of these catalysts is carried out (contact time, H₂ pressure and H₂/DBF molar ratio) to better understand the reaction mechanism of the two phosphide-type catalysts.

2. Experimental

2.1. Materials

The catalytic support used in this study was commercial fumed silica from Sigma-Aldrich. The reagents used for the preparation of the precursor solution were nickel(II) hydroxide (Ni(OH)₂), cobalt(II) hydroxide (Co(OH)₂, 95%) and phosphorous acid (H₂PO₃H, 99%), all supplied by Sigma Aldrich. In the catalytic reactivity study the chemicals used were dibenzofuran(C₁₂H₈O, 98%) and decahydronaphtalene, cis+trans (C₁₀H₁₈, 98%), both from Alfa Aesar. The gases used in the experimentation, H₂, N₂, CO, NH₃ had a purity of 99.9999% and were supplied by Air Liquide.

2.2. Preparation of catalysts

Nickel phosphide and cobalt phosphide catalysts containing 5 wt.% of nickel and cobalt, respectively, supported on fumed commercial silica ($S_{BET} = 218 \text{ m}^2 \text{ g}^{-1}$ and $Vp = 0.52 \text{ cm}^3 \text{ g}^{-1}$) were prepared. Precursor solutions of nickel(II) dihydrogenphosphite (Ni(HPO_3H)_2) and cobalt(II) dihydrogenphosphite (Co(HPO_3H)_2) were prepared as described elsewhere^{14, 15}. The precursor solutions were incorporated into the silica support by the incipient wetness impregnation method to prepare catalysts containing 5 wt.% of metal. To convert the precursor phosphite into phosphide the impregnated solids were dried at 60°C for 24 hours and treated in hydrogen by heating at a linear temperature ramp (3 °C min⁻¹) in flowing H₂ (100 ml min⁻¹), from rt. to 620 °C for nickel phosphide catalysts and from rt. to 700 °C for cobalt phosphide catalysts. The obtained catalysts will hereafter be named as Ni₂P and CoP.

2.3. Characterization of catalysts

The textural properties (S_{BET}, V_p, d_p) of the fresh catalysts were obtained from the N₂ adsorptiondesorption isotherms at -196 °C measured with a Micromeritics ASAP 2020 apparatus. Prior to the measurements the samples were outgassed at 200 °C and 0.01 kPa overnight. Surface areas were determined by using the Brunauer-Emmet-Teller equation and a nitrogen molecule cross section of 16.2 Å² (3). The pore size distribution was calculated by applying the Barrer-Joyner-Halenda method (BJH) to the desorption branch. The total pore volume was calculated from the adsorption isotherm at P/P₀ = 0.95.

X-ray diffraction patterns (XRD) of the samples were obtained with aX 'Pert PRO MPD Philips diffractometer (PANanalytical), equipped with a Ge (111) primary monochromator using CuK α 1 (1.5406 Å) radiation. The X-ray tube was set at 45 kV and 40 mA and the measurements were carried out from 10° to 70° (2theta) for 12 minutes.

CO chemisorption analyses were performed under static volumetric conditions in a Micromeritics ASAP 2020 apparatus. Samples were reduced ex situ and transferred into an inert atmosphere. Prior to measurement samples were re-reduced in situ in H₂ at 300 °C and evacuated at 25 °C for 10 h. The chemisorption isotherm was obtained by measuring the amount of CO adsorbed between 10 and 600 mmHg at 35°C. After completing the initial analysis, the reversibly adsorbed gas was evacuated and the analysis repeated to determine only the chemisorbed amounts.

X-ray photoelectron spectra were collected using a Physical Electronics PHI 5700 spectrometer with non-monochromatic Al K_{α} radiation (300 W, 15 kV, and 1486.6 eV) with a multi-channel detector. Spectra of pelletized samples were recorded in the constant pass energy mode at 29.35 eV, using a 720 μ m diameter analysis area. Charge referencing was measured against adventitious carbon (C 1s at 284.8 eV). A PHIACCESS ESCA-V6.0 F software package was used for acquisition and data analysis. A Shirley-type background was subtracted from the signals. Recorded spectra were always fitted using Gaussian-Lorentzian curves in order to determine the binding energy of the different element core levels more accurately. Reduced catalysts were stored in sealed vials with an inert solvent. The sample were prepared in a dry box under a N₂ flow and analyzed without previous treatment.

Temperature-programmed desorption (TPD) experiments were carried out to determine the acidity of the samples by using ammonia (NH₃-TPD) and the hydrogen activation capability was determined by using Hydrogen (H₂-TPD). For NH₃-TPD experiments, 80 mg of catalyst precur-

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sor was reduced at atmospheric pressure using the same procedure described in the section 2.2. After flushing with helium and adsorption of ammonia at 100 °C, NH₃-TPD was performed using a helium flow and raising the temperature from 100 to 800 °C at a heating rate of 10 °C min⁻¹. Prior to raising the temperature, physisorbed ammonia was removed flushing with helium. For the H₂-TPD measurements 300 mg of catalyst precursor were reduced using the same procedure as described in section 2.2. The catalyst was cooled under flowing helium from the reduction temperature down to 100 °C and then exposed to H₂ gas with a flow of 60 mL min⁻¹ at 100 °C for 30 minutes, followed by flushing in argon for another 30 minutes in order to remove physisorbed hydrogen, and then heated in argon to 800 °C. The evolved gas was analyzed by online gas chromatography (Shimadzu GC-14A) equipped with a TCD.

Fourier transform infrared (FTIR) spectra of the samples after reduction and outgassing at 400°C as well as FTIR spectra of adsorbed CO were recorded with a Nicolet Nexus instrument, using a conventional cell connected to a gas-handling system. All spectra were obtained after subtraction of the spectrum corresponding to the degassed sample. Prior to adsorption, pressed disks of powder samples with a diameter of 1.5 cm of sample powders (ca.20 mg) were thermally treated within the cell at 400 °C Under H₂ (67 kPa) for 1 h. CO adsorption (0.13 kPa) experiments were performed at -140 °C and spectra were recorded in the range -140 to 10 °C while degassing.

2.4. Catalytic Activity

Dibenzofuran (DBF) was chosen as a model compound for the HDO reaction of bio-oil. Furan derivate compounds are common template molecules used to evaluate the catalyst performance in HDO reactions including dibenzofuran which is a very low reactive compound. The reactions were carried out in a high-pressure fixed-bed continuous-flow stainless steel catalytic reactor (9.1 mm in diameter and 230 mm in length). The concentration of the organic feed was in the range of 1-4 wt.% and decalin was used as solvent. The catalysts were reduced ex-situ and saved in a close vial containing cyclohexane to avoid oxidation. For the activity tests, an amount of 0.25 g of catalyst were used (particle size 0.85-1.00 mm) diluted with quartz sand to a total volume of 3 cm³. Prior to the catalytic tests, the catalyst was left to dry overnight in a hydrogen flow of 100 mL min⁻¹ with a temperature ramp starting at 40 °C and finishing at 400°C with a heating rate of 3 °C min⁻¹. After drying and before catalytic testing, the catalysts were left in a nitrogen flow of 20 mL min⁻¹ at 40 °C to avoid oxidation. Catalytic activities were measured at 275 °C under a 0.1-3.0 MPa pressure, with a flow rate of H₂ of 100 mL min⁻¹ and with hourly space velocities (WHSV) in the range of $0.4 - 2.6 \text{ h}^{-1}$. The evolution of the reaction was monitored by collecting liquid samples every 60 minutes for 6 hours. A volume of 0.33 ml of product was col-

lected and diluted up to 1 ml volume with pure decalin and 20 µl of nonane were added as an internal standard. These samples were kept in sealed vials and subsequently analyzed by gas chromatography (Shimadzu GC-14B, equipped with a flame ionization detector and a capillary column, TBR-14, coupled to an automatic Shimadzu AOC-20i injector). In order to identify intermediates samples were also analyzed by CG-MS (MODEL) with a TBR-14 capillarity column.

The catalytic selectivity was calculated considering all the reaction products observed, biphenyl (BP), cyclohexylbenzene (CHB), tetrahydrodibenzofuran (THDBF), hexahydrodibenzofuran (HHDBF), 2-cyclohexylphenol (2-CHP), bicyclohexyl-2-en-3-ol (BCH-2-en-3-ol) and cyclopen-tylmethylcyclohexane (C-PE-ME-CH). Carbon balances are in the range of 1.00 +/- 0.05

The HDO conversion was calculated by the following equation;

$$HDO_{Conv.}(\%) = \frac{C_{DBF.Ini} - C_{DBF.Fin} - C_{Int}}{C_{DBF.Ini}} \times 100$$
(1)

where $C_{DBF.Ini}$ is the DBF concentration in the feed, $C_{DBF.Fin}$ is the DBF concentration in the HDO liquid product and C_{IntHyd} is the concentration of O-containing intermediates (THDBF, HHDBF, 2-CHP, BCH-3-en-2-ol).

The turnover frequency was calculated using equation (2):

$$TOF = -\frac{F/W}{M} \times \ln(1 - X)$$
(2)

where F is the molar rate of reactant (mol min⁻¹), W is the catalyst weight (g), X is the conversion and M is the mole of sites loaded (mol g⁻¹) calculated from CO-chemisorption. This formula was used because the conversions were close to 100% so were far from differential conditions, and an integral analysis was needed. This equation, in which $-\ln(1-X)$ substitutes for X assumes a pseudo first-order reaction which may be justified by the large excess of hydrogen.

In order to determine if internal diffusion is limiting the reaction the Weisz-Prater criterion can be used. It uses measured values of the rate of reaction to determine if the reaction suffers from problem of internal diffusion. Internal mass transfer effect can be neglected when the value of CWP is less than 1^{16} . The detail of the calculations made is found as suplementary information (S1). All the values of C_{WP} obtained under the experimental conditions studied were in the order of 10^{-2} - 10^{-3} . This indicates that internal mass transfer effects can be neglected at the conditions employed for this reactivity study.

3. Results and discussion

3.1. Catalytic study

The HDO activity and selectivity of the reduced catalysts were investigated by using DBF as a model molecule present in bio-oil. Several experimental parameters were evaluated in order to better understand the reaction mechanism of the HDO reaction of DBF and to find the optimal experimental conditions to transform DBF into O-free products. Several types of products can be formed during HDO reaction, presented in Scheme 1, depending on the route followed. There is a hydrogenation pathway 1, in which oxygen removal is preceded by hydrogenation of benzene rings; a direct deoxygenation pathway 3, in which oxygen removal occurs by direct cleavage of bonds linking oxygen to aromatic carbons; and an intermediate pathway 2, in which oxygen removal is preceded by hydrogenation of one benzene ring only.

In the present work a further study to investigate the reaction mechanism of HDO of DBF on Ni_2P and CoP catalysts is performed by changing several operational parameters; contact time, reaction pressure and H_2 /DBF ratio feed.

3.1.1 Contact time analysis

Contact-time experiments were carried out to improve the understanding of the reaction network and therefore the hydrogen gas volume flow and the feed (DBF in decalin) volume flow were adjusted. The hydrogen pressure, H₂/DBF molar ratio and the reaction temperature maintained constant. The catalytic results in terms of total conversion and HDO conversion are presented in Figure 1 in a contact time range of 1.8-12 s. It is observed that both total (dotted lines) and HDO conversion (solid lines) rise with increased contact time. Total conversion is always higher than HDO conversion, since the former only considers DBF transformation; instead, the later considers DBF transformation into O-free compounds. It is clearly seen that the longer the catalyst is in contact with the reactants the more of them can react and transform into oxygen free products. The Ni₂P catalyst turns out to be more active than CoP, whereas only at low contact times CoP is slightly more active than Ni₂P. Ni₂P catalyst presents the highest HDO conversion values at a contact time of 6 s, where the conversion values are reaching 98%. Instead, on CoP catalyst the highest value (80%) is attained at a contact time of 12 s. At low contact times the catalysts exhibit a low catalytic activity (less than 15% conversion). With regard of to the lower conversion value observed at very high contact time for Ni₂P catalyst, in a recent work of A. Lino et al¹⁹ about HDO of 2-MTHF on Ni₂P, these authors have observed a linear relationship in the plot of

conversion versus contact time. These author have pointed that in this situation each reactant molecule is reacting independently, meaning that each reactant molecule sees essentially a fresh surface. On Ni_2P , the linearity is maintained until a contact time of 6 s where the conversion is close to 100%. By increasing the contact time the conversion lowers perhaps to surface coverage with other organic species. If there were coverage, the rate of adsorption would be retarded by those species and therefore decreasing the conversion.

The reaction network is studied by observing the trends of product selectivity at different contact times. Figure 2 shows the product selectivity as a function of contact time for Ni₂P catalyst and CoP catalyst, respectively. The products are classified into three groups; saturated oxygen-free cyclic compounds, unsaturated oxygen-containing cyclic compounds and furans.

At low contact times the intermediates tetrahydrodibenzofuran (THDBF) and hexahydrodibenzofuran (HHDBF) are observed as products along with 2-cyclohexylphenol (2CHP) and bicyclohexyl-2-en-3-ol (BCH-2-en-3-ol). It should be noted that on Ni₂P, the selectivity to 2-CHP did not change obviously, and the selectivity to BCH-2-en-3-ol firstly increased and then decreased, indicating that BCH-2-en-3-ol is an intermediate for Ni₂P catalyst. This is not the case for CoP one: as the contact time increased, the selectivities to 2-CHP and BCH-2-en-3-ol did not change monotonously.

By increasing the contact time the transformation of O-containing intermediates into O-free compounds occurs. Bicyclohexane (BCH) is the main O-free compound product formed in all reactions.

Based on the observation of formed intermediates and products and reviews of earlier studies 9,22 , the reaction mechanism of HDO on CoP and Ni₂P catalysts follows the hydrogenation pathway 1. In the first step one of the benzene rings in DBF is hydrogenated and the furan ring opens up:

DBF→THDBF→HHDBF→2-CHP

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In the next step the second benzene ring is hydrogenated:

2-CHP→BCH-3-en-2-ol→BCHol

Then the oxygen removal takes place by dehydration and cycloalkane formation, which is the last step when both aromatic rings are completely saturated.

BCHol→BCH + C-PE-CH + C-PE-ME-CH

From these results it can be concluded that the catalysts have both metallic and acidic properties. They show good hydrogenation properties which can be explained by the high metallic disper-

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sion and they also present acidic properties which shows in their hydroisomerization ability leading to C-PE-CH and C-PE-ME-CH. Moreover a contact time of 6 s seems to be adequate to assess the transformation of DBF into O-free products by following the route 1 presented in Scheme 1.

3.1.2 H₂ Pressure study

Another parameter evaluated was the reaction H₂ pressure, an important parameter in hydrogenation reactions. For this aim the reaction was carried out at three different H_2 pressures, 1, 15 and 30 bar, while the other parameters were kept constant; temperature 275 °C, H_2/DBF molar ratio 69 and the contact time 6 s. Figure 3 shows the total conversion and the HDO conversion as a function of the H₂ pressure. The plots of Figure 3 indicate that the activity is very low when the reaction is performed at atmospheric pressure (the conversion is below 5% with both catalysts). Nonetheless, Ni₂P is more active than CoP at medium and high pressures. Moreover, both Ni₂P and CoP are active at medium pressures with HDO degrees similar to those attained at 30 bar in the case of CoP. These results indicate that the reaction can be carried out at moderate pressures and that high pressures are not required to perform O-removal as in S- N- removal, where high pressures are always required. A higher H₂ pressure is beneficial also in hydrodeoxygenation as by increasing the pressure the thermodynamics would shift further towards complete conversion. However, too much H₂ interferes with the adsorption of reactants on the surface¹⁹. This could explain why the conversion is slightly higher at the 15 bar pressure compared to the 30 bar pressure for CoP catalyst. Wang et al²³ observed an improvement in DBF conversion via the hydrogenation pathway by increasing the pressure from 0.5 to 5.0 MPa on Pt/SBA-15 catalysts. In our case the excellent activity of metallic phosphides, even being highly active at medium pressures, should be highlighted in this type of reaction.

Considering the O-free product yield (Figure 4) at the different pressures, it is possible to observe that there is no major changes in product yield distribution by increasing the pressure from 15 to 30 bar in the case of the CoP sample, however, the Ni₂P catalyst yielded greater amounts of BCH at high pressures. It is worth noting that the maximum yield of a certain intermediate was <4% in all experiments. At pressures of 15 and 30 bar the yield of the final product, BCH, is much higher for Ni₂P. These observations are consistent with the mechanism and the conclusion is that CoP catalyst is less active than Ni₂P.

3.1.3 H₂/DBF molar ratio study

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With the aim to study how the reaction is influenced by different H_2/DBF molar ratios, the feed was prepared with three different compositions (H_2/DBF molar ratios of 34, 69 and 139). Figure 5 shows the total conversion and the HDO conversion as a function of H_2/DBF molar ratio.

The first thing observed is that the Ni₂P catalyst is much more active than CoP, and also that the Ni₂P sample is less affected by H₂/DBF molar ratios than the CoP catalyst. In fact, at low H₂/DBF molar ratios the activity of the Ni₂P sample is close to that observed at high molar ratios. On the contrary the CoP sample shows a sharp increase in activity by increasing the H₂/DBF molar ratio. The highest conversion of all the experiments, when keeping the contact time at 6 s, is achieved at the highest H₂/DBF molar ratio for both catalysts.

The yields of the different reaction products for this set of experiments are depicted in Figure 6. It is observed that the Ni₂P sample shows high yields to BCH (>80%) regardless the H₂/DBF molar ratio employed, which demonstrates its high efficiency to convert DBF into O-free compounds under poorer H₂ conditions. The CoP samples only attain high yields of BCH at high H₂/DBF molar ratios.

In fact, when the HDO conversion evolution with time on stream is observed (Figure 7), at low H_2/DBF molar ratio (Figure 7A), cobalt catalyst suffers from a fast deactivation after the first hour of reaction. Its initial HDO conversion is close to 80%, meanwhile after 6 h on stream the HDO conversion attained is down at 35%. However HDO conversion for the Ni₂P sample under the same conditions slightly decreases and reaches the steady state after 6 hours with a conversion of 89%.Contrary and as expected, at $H_2/DBF=139$ (Figure 7.B) both catalysts are rather stable, which confirms previous statement; higher H_2 contents are required to improve the activity of CoP.

3.1.4. Intrinsic activity of the prepared sample-TOF number

In order to evaluate the intrinsic activity of Ni_2P and CoP catalysts at the different conditions employed, TOF numbers were calculated according to Eq. 2 and is defined as the moles of reactant converted per mol of active sites per unit of time. This value gives information about the activity per active site. Table1 compiles the calculated TOF numbers for contact time, pressure effect and H_2/DBF effect experiments, respectively.

From this table, it is observed how the TOF number is higher for the Ni_2P catalyst except at low contact time contact time. Moreover, the TOF number increases for the Ni_2P catalyst when the contact time is increased up to 6 s, for longer contact times the TOF number decreases. However,

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no great changes in TOF number are observed for the CoP sample, what could indicate that the activity per active site is hardly affected by changes in contact time; it only increases at the highest contact time studied. Contrary the Ni₂P sample is much more sensitive to these changes with the TOF number increasing with about 400% from 1.8 to 6.0 s, in accordance to the sharp increase in the conversion. From H₂ pressure effect experiments it is observed that at low H₂ pressure, the activity per active site is very low for both samples. Nevertheless at 15 and 30 bar, the Ni₂P sites are much more active than the CoP ones. Finally, H₂/DBF experiments also showed that the Ni₂P sites had the highest activity, presenting much higher TOF values than the CoP samples. Wang et al.²³ tested Pt, Pd and Ru catalysts in HDO of DBF and the most active catalyst presented a TOF value, at 30% HDO conversion, lower than the TOF value presented by CoP catalysts at the same conversion value (15 bar), 0.42 s^{-1} vs. 2.55 s^{-1} , respectively.

It should be highlighted that the TOF value reached under H₂-poor conditions (4 wt.%DBF) is the highest 23.7 s⁻¹ and much higher than those found in the literature for noble-metal based catalysts²²⁻²⁴.

3.2. Catalyst characterization

The textural properties of the prepared catalysts are included in Table 2. As showed in this table, BET surface area of the SiO₂ support was 218 m² g⁻¹ and that of the fresh Ni₂P and CoP catalysts in pellets form were 109 and 126 m² g⁻¹, respectively. The decrease in specific surface area indicates that pores are partly blocked due to the presence of phosphides particles on the support which partially impede the access of gas molecules^{25, 26}. The textural properties of Ni₂P and CoP prepared catalysts were similar.

The active phases formed after reduction were firstly evaluated from XRD experiments. Ni₂P and CoP catalysts do not show any well-defined diffraction lines (diffractograms not shown here). This is due to the low metallic loading of the catalysts (5 wt.%) and because of the high dispersion of the active phase as observed in earlier studies⁹. The diffractogram of Ni₂P shows some weak diffraction signals centred at 20 (°) = 40.7, 44.6 and 55.0 and assigned to the Ni₂P phase (PDF 01-089-4864). Meanwhile, CoP catalyst shows a very weak diffraction line located at 20 (°) = 48.3 and assigned to the CoP phase (JCPDS No. 29-0497).

CO chemisorption capacity of the reduced catalysts provides with information about the number of surface metal atoms and gives an estimation of the number of active sites. Table 2 compiles the CO uptake at room temperature for cobalt phosphide and nickel phosphide catalysts. Both samples chemisorb similar amounts of CO, i.e., the number of active sites is quite similar in both catalysts.

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A more detailed information about the phases formed on the catalyst surface was obtained by XPS. The corresponding binding energies values are summarized in Table 3. Both, Ni 2p core level spectrum and Co 2p core level spectrum are decomposed into three contributions, i.e., three doublets containing the $2p_{3/2}$ and $2p_{1/2}$ components. The first contribution at lower binding energy is due to Ni₂P (853.1 eV) and CoP phase (777.3 eV); the second contribution centered at 856.6 eV for Ni₂P catalyst and at 780.7 eV for CoP samples is due to Me²⁺ species; meanwhile, the third contribution observed is due to the shake-up satellite assigned in the literature to divalent species. The core level spectra of P 2p were also analyzed (See Fig S1, Supplementary Information) and decomposed. Both catalysts show two doublets (P $2p_{3/2}$ -solid lines and P $2p_{1/2}$ dash lines) were observed indicating that, at least, two phosphorus species are present on the catalyst surface. The band close to 134.0 eV is due to phosphate species and the other band at 129.3 eV for Ni₂P sample and at 129.0 eV for CoP one is due to the phosphide phase. No unreduced HPO₃H⁻ species are observed, which could be a result of the high reduction temperature employed³¹. A much more detailed discussion of XPS spectra for these phases can be found in previous papers^{27, 28}.

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Table 3 includes the percentage of phosphorous as PO_4^{3-} and as phosphide (Ni₂P or CoP). It is clearly seen the much greater amount of phosphate species on Ni₂P rather than on CoP, indicating a surface enrichment of P in the former. In line with this, the corresponding surface atomic ratios indicate that the Ni/P molar atomic ratio is much lower than the expected values due to the formation of the Ni₂P phase. On the contrary, for CoP sample, the Co/P ratio is higher than the expected stoichiometric value due to the presence of CoP which indicates that there is a greater concentration of metallic atoms on the surface.

These results are explained considering that the same amount of P-precursor has been added in both catalysts. Nonetheless Ni₂P is a metallic rich phosphide and CoP is a stoichiometric phosphide, therefore more P is included into the CoP structure. Moreover the reduction temperature to form CoP is higher than that for Ni₂P, 700 °C vs 620 °C. Therefore it is expected thatat greater quantities of P are released during the reduction process of CoP. As a result, the remaining extra-P is expected to be higher in the Ni₂P catalyst.

The acidic properties of the catalysts were measured by temperature-programmed desorption of ammonia (NH₃-TPD) between 100 and 800 °C. Figure 8 shows the corresponding profiles while

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Table 4 compiles the amount of NH₃ desorbed (μ mol of NH₃ g⁻¹) for the reduced catalysts. By comparing both profiles and the data compiled in Table 3 one sees that the amount of NH₃ released is much higher on Ni₂P than on CoP and Ni₂P sample releases NH₃ molecules at temperatures lower than 400 °C (weak acidity) while CoP sample releases NH₃ molecules at very high

tures lower than 400 °C (weak acidity) while CoP sample releases NH₃ molecules at very high temperatures (550-700 °C), indicating the presence of strong acid sites on this CoP sample. Previously the presence of both Lewis and Brönsted acid sites has been reported for the Ni₂P catalyst²⁹. Lewis acidity was ascribed to unreduced Ni²⁺ and Brönsted acidity due to PO-H species. No much Co²⁺ or Ni²⁺ species are expected to be present here, due to both the high reduction temperature applied. Therefore, the acidity would be expected to come from PO-H groups. XPS data show that the Ni₂P sample presents a surface enrichment of P species which could account for its higher acidity.

FT-IR spectra of the samples after reduction and outgassing at 400°C will also give information about the presence of surface hydroxyl groups. The corresponding spectra are represented in Figure 9. These clearly show a main band at 3742 cm⁻¹ due to stretching of isolated silanol species together with a weaker band centered at 3670 cm⁻¹ due to PO-H stretching modes, thus confirming that the acidity detected over these samples arises mainly from the Phosphorouscontaining species. The frequency of this band is completely consistent with the frequency values reported for phosphoric acid supported on several oxides such as Al₂O₃, SiO₂, ZrO₂, forming hydrogenphosphate species^{30, 31}. A shoulder at 3700 cm⁻¹ also seems to be present, which could be due to another kind of OH groups showing intermediate acidic properties, possibly some silanols affected by nearby phosphate species. Moreover, the intensity ratio I₃₆₇₀/I₃₇₄₂ for the two silanol and P-OH bands is higher in the Ni₂P sample spectrum, as a further evidence that this catalyst shows an higher surface concentration of P-OH species, as it was previously suggested by XPS data, and also justifying the greater acidity presented by this catalyst. In line with this, IR analysis of adsorbed CO at liquid nitrogen temperature over Ni₂P sample corroborates the presence of P-OH groups on this catalyst. The corresponding subtraction spectra are reported in Figure 10, in the CO stretching region the main bands of the spectrum are centered at 2187, 2158 and 2130 cm⁻¹, shoulder. The main band at 2158 cm⁻¹ is assigned to CO weakly interacting with surface OH groups³² through H- bonds and this band is readily disappearing at decreasing CO coverage. Its assignation is also confirmed by the negative band appearing in the OH stretching region (inset) due to the consumption of isolated OH groups after interaction with CO. From these data it seems that silanols are involved first (highest frequency band), but also P-OH are interacting with CO which behaves as weak base (negative component at 2168 cm⁻¹). Decreasing

CO coverage leads to a partial restoring of silanols, but on the other hand the negative band due to P-OH group is still clearly detectable, thus the hydrogen bonding of CO with the surface hydroxy groups of hydrogenphosphate is significantly stronger than that with silanol, in agreement with P-OH higher acidic strength.

On the other hand the band detected at 2187 cm⁻¹ is due to carbonyls coordinated over Ni-ions, acting as Lewis acidic sites at the catalyst surface^{29, 32}. Upon outgassing this band decreases in intensity and shifts towards higher frequencies, as expected. The weaker component at 2130 cm⁻¹ can be assigned to carbonyls coordinated over partially reduced Ni ions (possibly Ni⁺), whereas only a very weak and broad absorption from 2050 to 2000 cm⁻¹ could be assigned to carbonyls coordinated over a very dispersed nickel metal phase³³. The presence of some metallic nickel particles cannot be discarded, possibly arising from Ni ions strongly interacting with the support and not involved with P-species during reduction to form the phosphide phase.

As for the CoP sample, the strongest acidic sites detected by NH₃-TPD could be assigned to the Co ions unreduced still exposed at the catalyst surface. In order to ascertain this hypothesis we performed CO adsorption and indeed the IR spectrum recorded at room temperature, i.e. when the strong band due to H-bound CO is not detectable, shows a weak carbonyl band at 2200 cm⁻¹ which can be assigned to CO coordinated over Co ions (insert in Figure 10).

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Finally, when testing a catalyst for hydrotreating reactions, it is important to consider its capability to activate hydrogen. To this end, H₂-TPD experiments on Ni₂P and CoP reduced samples were performed and their corresponding profiles are shown in Figure 11. In general, the temperature at which H-species are released helps to understand the nature of these species. The H₂desorption below 250 °C is ascribed to the adsorbed H₂ on metal Ni or Co sites, while the H₂ desorption between 250°C and 450°C is ascribed to the H₂ from the metal-support interface. Finally, desorption at higher temperatures is usually ascribed to H₂ spill over species. At first glance, when comparing the H₂-TPD profiles of the two different catalysts it is clear that the Ni₂P catalyst desorbs greater amounts of hydrogen species, indicating a greater capacity to activate H₂ compared to CoP. Moreover, the Ni₂P catalyst mainly desorbs hydrogen at low temperature, that is, coming from the metallic surface. Besides, greater amounts of H-spillover are also present on this catalyst. Literature data indicate that H₂-desorption at intermediate temperatures on Ni₂P catalysts could be related to P-OH groups at the metal support interface³⁴ and catalysts with greater amount of P-OH groups desorbs a greater amount of H₂³⁵. In fact, the Ni₂P catalyst desorbs greater amounts of H₂ from both metal surface and metal-support interface, revealing the

greater capability of Ni₂P to activate hydrogen and also the greater amount of P-OH present, contributing to H₂-desorption at intermediate temperatures.

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3.3. Discussion

Catalytic results have shown the greater activity of the Ni₂P catalyst with regard to the CoP one. The most important results are that both catalysts show important conversion levels at medium pressure (15 bar) and that Ni₂P catalyst is hardly affected by changes in the H_2/DBF ratio, meanwhile the CoP one requires greater amounts of hydrogen to be active.

The main cause for catalyst deactivation in HDO reactions is considered to be carbon deposition on the surface²⁰. Brønsted acid sites are responsible for coke formation as they act as proton donors that form carbocations, therefore coking increases with increasing acidity of the catalyst. However the HDO reaction does require acidity and a balance should be obtained to have acid sites capable to activate oxygen containing molecules but whose strength will not be too strong to give rise to coke formation³⁶. As stated before, the Ni₂P sample, presents a higher amount of acid sites of weak nature, probably related to its higher amount of P-OH groups on the surface. The CoP sample presents an important concentration of very strong acid sites that could be responsible for coke formation³⁶, this could explain the low activity at low H₂/DBF molar ratio where the concentration of C-species is expected to be higher. By increasing the H₂ content in the feed, hydrogen acts as a cleaner hydrogenating C-species on the surface and the coke formation is minimized³⁷.

Another reason for catalyst deactivation is the formation of water during reaction. The water formed could interact with oxidized metal phosphide particles. It has been observed in a previous study⁹ that Ni₂P catalysts with an initial P/Ni atomic ratio of 2 or higher are more stable than those with the P/Ni atomic ratio of 1. The oxidation resistance of Ni₂P phase under water molecules has been explained in the literature³⁸ by the dissociation of H₂O on Ni₂P surface (0 0 1) which generates O atoms that interact with both P and Ni atoms, but preferably with P. This leads to the formation of oxy-phosphide species with electron transfer from P to O. Hence P shows a ligand effect and the P species may inhibit the interaction of Ni with O, improving the oxidation resistance of Ni₂P. In a previous work, we observed this fact on Ni₂P catalysts, that is, catalysts with greater P content were more resistant to deactivation due to H₂O interacting presumably with P-species⁹. By looking at XPS and IR data, it is clearly seen how the Ni₂P sample

has a higher P-concentration on the surface compared to CoP mainly in the form of P-OH. This could be another reason why Ni₂P shows the highest activity of the two catalysts.

The better results obtained with Ni₂P vs CoP is not only related to the higher intrinsic activity of Ni₂P but also extra P-species that play an important role in these systems. Characterization results have evidenced the presence of much more P-on the surface of Ni₂P, mainly in the form of P-OH groups and as a consequence this catalyst possesses better acidity and H₂-desorption capacity, mainly at low temperature. P-species provide P-OH groups that can act as acid site and as a source of hydrogen species.

The role of P was also evaluated in CoP catalysts presenting different P/Co molar ratios and tested in the catalytic hydrodechlorination of chlorobenzene³⁷. The most active catalysts were those containing P/Co>2 and showing better acidity values and greater amount of H-species from H₂-TPD measurements. In line with this, in a recent manuscript of Chen et al³⁹, it has been evaluated the promoting effect of water and oxygen pretreatment in the hydrodechlorination activity of Ni₂P/SiO₂. H₂O or O₂ pretreatment increased the amount of P-OH groups on the surface and improved the catalytic response of these systems. So the activity of Ni₂P was explained in terms of synergism between Ni atoms and P-OH groups.

4. Conclusions

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The catalytic activity of Nickel and Cobalt phosphides on silica was tested in hydrodeoxygenation of DBF as a model oxygenate compound under different experimental conditions. The results obtained pointed out that the Ni₂P catalyst is more active than the CoP one. Moreover it is less affected by hydrogen pressure changes and presenting very high conversion values even at low H₂/DBF molar ratios. Both catalysts perform the HDO reaction through the hydrogenation route: DBF \rightarrow THDBF \rightarrow HHDBF \rightarrow 2-CHP \rightarrow BCH-3-en-2-ol \rightarrow BCHol \rightarrow BCH + C-PE-CH + C-PE-ME-CH. Characterization results have indicated that the better activity of the Ni₂P phase is related to its greater intrinsic activity, and the greater presence of extra-P. Extra P-OH species can prevent metal phase oxidation; act as weak acid sites that can interact with O-species; but they can also act as H-donors to perform reaction, meanwhile spillover species could act as surface cleaners avoiding deactivation by coke formation.

The presented results seem to indicate that extra P-OH groups on Ni_2P could be those governing the excellent results presented by the Ni_2P phase. Ni_2P sample should be considered a good candidate to be employed in hydrotreating units to perform HDO reactions.

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DBF: Dibenzofuran, THDBF: 2,3,4,9 Tetrahydrodibenzofuran, HHDBF: 2,3,4,4a,9,9a Hexahydrodibenzofuran, 2-CHP: 2-cyclohexylphenol, BCH-3-en-2-ol isomers:
1-1'bi(cyclohexan)-3-en-2-ol, BCHol: By(cyclohexan)-2ol, BCH: Bicyclohexane, C-PE-CH: Cyclopentylcyclohexane, C-PE-ME-CH: (Cyclopentylmethyl)cyclohexane,
2-PP: 2-phenylphenol, 2P-1CHol: 2-Phenyl-1-cyclohexanol, CHB: Cyclohexylphenzene, CH: Cyclohexane, B: Benzene, BP: Biphenyl

Scheme 1. Mechanism of HDO of DBF

Mechanism of HDO of DBF

Table 1. TOF number for Ni_2P/SiO_2 and CoP/SiO_2 catalysts at T = 275 °C, T.o.S.=6 h under dif-

	Ni ₂ P/SiO ₂	CoP/SiO ₂		
Contact time (s)	TOF $x \ 10^2 \ (s^{-1})^a$			
1.8	1.69	3.33		
2.4	2.67	2.27		
3.6	6.68	1.40		
6.0	8.42	2.18		
12.0	4.30	3.15		
H ₂ Pressure	TOE ~ 10	$2_{(a}-1)b$		
(Bar)	$10F \times 10^{-1}(s^{-1})^{-1}$			
1	0.33	0.14		
15	7.95	2.55		
30	8.42	2.18		
H ₂ /DBF	TOF <i>x</i> 10 ²	$(s^{-1})^{c}$		
34.3	23.7	3.01		
69.2	8.42	2.18		
139.0	6.09	2.93		
3 D 20 D 111/D	DE (0			

ferent experimental conditions

^a P=30 Bar and $H_2/DBF=69$

^b Contact time= 6 s and H₂/DBF=69

^c P= 30 Bar and Contact time= 6 s

	$\frac{S_{BET}}{(m^2 g^{-1})}$	$\frac{V_p^{b}}{(cm^3 g^{-1})}$	CO Chemisorption ^c (µmol g ⁻¹)
Support SiO ₂	218	0.52	
Ni ₂ P Pellets	109	0.57	32.9
CoP Pellets	126	0.67	36.7

Table 2. Textural Properties^b and CO Chemisorption^c properties of the prepared samples

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Sample]	Atomic ratio				
	Ni 2 <i>p</i> _{3/2} /	/Co 2 <i>p</i> _{3/2}	P 2 <i>p</i> _{3/2}		Me/P	Me/Si
	Co ²⁺	CoP	PO4 ³⁻	CoP		
CoP	780 7	777 3	134.1	129.0	1.65	0.04
	/80./	111.5	(48.6)*	(51.4)*		
	Ni ²⁺	Ni ₂ P	PO_4^{3-}	Ni ₂ P		
Ni ₂ P	856.6	856.6 853.1	134.4	129.3	0.41	0.01
	050.0		(25.5)*	(74.5)*		

Table 3. Binding energy (eV) of core electrons and surface atomic ratios of fresh CoP and Ni_2P catalysts

*In parenthesis the percentage of each specie

Γable 4. Quantity of NH ₃	released (µmol g ⁻¹)) from NH ₃ -TPD	experiments
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	μmol NH ₃ g ⁻¹					
Catalyst			Tem	perature (°C)		
Cataryst	Total	100-200	200-300	300-400	400-500	500-800
Ni ₂ P	481	155	202	98	20	6
CoP	211	57	71	35	8	41

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Figure 1. Total conversion and HDO conversion of DBF on Ni_2P/SiO_2 and CoP/SiO_2 as a function of contact time. Experimental conditions: T = 275 °C; P = 30 Bar; H₂/DBF ratio 69; T.o.S= 6 h



Figure 2. Product selectivity on Ni₂P/SiO₂ and CoP/SiO₂ as a function of contact time. Experimental conditions: T = 275 °C; P = 30 Bar; H₂/DBF molar ratio 69; T.o.S= 6 h



Figure 3. Total conversion and HDO conversion as a function of reaction pressure for Ni_2P/SiO_2 and CoP/SiO_2 catalyst. Experimental conditions: T = 275 °C; contact time 6 s; H₂/DBF molar ratio 69; T.o.S= 6 h

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Figure 4. Yield distribution of the products as a function of H_2 pressure for Ni₂P/SiO₂ and CoP/SiO₂ catalyst. Experimental conditions: T = 275 °C; contact time 6 s; H₂/DBF molar ratio 69; T.o.S= 6 h



Figure 5. Total conversion and HDO conversion as a function of H_2/DBF molar ratio for Ni_2P/SiO_2 and CoP/SiO_2 catalysts. Experimental conditions: T = 275 °C; P = 30 bar; contact time 6 s; T.o.S.= 6 h

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Figure 6. Yield distribution of the products as a function of H_2/DBF molar ratio for Ni_2P/SiO_2 and CoP/SiO₂ catalysts. Experimental conditions: T = 275 °C; P = 30 Bar; contact time 6 s; T.o.S.=6 h



Figure 7. Evolution of conversion as a function of time on stream for Ni_2P/SiO_2 and CoP/SiO_2 catalysts tested at two different H₂/DBF molar ratios, A) 34 and B) 139. Experimental conditions: T = 275 °C; P = 30 bar; contact time 6 s; T.o.S. = 6 h



Figure 8. NH₃-TPD profiles of CoP and Ni₂P catalysts



Figure 9. FT IR spectra of Ni₂P and CoP samples after activation at 450°C. OH stretching region

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Figure 10. FT IR subtraction spectra of surface species arising from CO adsorption and outgassing at liquid nitrogen temperature on Ni_2P catalyst and CO adsorption on CoP at room temperature. The activated surface has been subtracted. Inset: OH stretching region



Figure 11. H₂-TPD profiles of CoP and Ni₂P catalysts