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COMPETING REACTIONS IN HYDROTREATING COKER DISTILLATES FROM ATHABASCA BITUMEN ON UNPROMOTED AND PROMOTED CATALYSTS

R. RANGANATHAN, M. TERNAN AND B. I. PARSONS CANADIAN FOSSIL FUEL RESEARCH LABORATORY CATALYSIS RESEARCH SECTION

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COMPETING REACTIONS IN HYDROTREATING COKER DISTILLATES FROM ATHABASCA BITUMEN ON UNPROMOTED AND PROMOTED CATALYSTS

by

Ramaswami Ranganathan¹, Marten Ternan² and Basil I. Parsons³

ABSTRACT

Desulphurization, denitrogenation and hydrogenation reactions occur simultaneously in a hydrotreating process. Several unpromoted $MoO_3/alumina$ catalysts and promoted $NiO-MoO_3/alumina$ and $CoO-MoO_3/alumina$ catalysts were prepared to compare the characteristics of these competing reactions using coker kerosene distillate (193-279°C) derived from Athabasca bitumen. The denitrogenation activity was found to be higher than the desulphurization activity for all the unpromoted $MoO_3/alumina$ catalysts. Promotion of $MoO_3/alumina$ catalysts with nickel or cobalt selectively improved desulphurization so that the extents of the various reactions were in the following order:

desulphurization > denitrogenation > aromatic hydrogenation

The interactions of the denitrogenation and the desulphurization reactions at various temperatures are compared for the promoted and unpromoted catalysts. The results obtained by others for hydrotreating gas-oil and low temperature distillate from coal tar are also compared. The conversion and the interaction of the desulphurization and denitrogenation reactions were found to be dependent on the type of catalyst, the temperature of reaction and the boiling range of the feedstock.

¹Research Scientist, ²Section Head, Catalysis Research and ³Acting Chief, Energy Research Laboratories, Department of Energy, Mines & Resources, Ottawa, Canada.

DES REACTIONS CONCURRENTES DANS L'HYDROTRAITEMENT DE DISTILLATS DE COKEFACTION, PROVENANT DE BITUME DE L'A-THABASCA, SUR DES CATALYSEURS ACTIVES ET NON-ACTIVES

Ramaswami Ranganathan¹, Marten Ternan² et Basil I. Parsons³

RESUME

par

Les trois réactions, désulfuration, dénitrification et hydrogénation, ont lieu simultanément pendant un procédé d'hydrotraitement. Plusieurs catalyseurs non-activés $MoO_3/alumine$ et activés $NiO-MoO_3/alumine$ et $CoO-MoO_3/alumine$ ont été préparés afin de comparer les caractéristiques de ces réactions concurrentes utilisant un distillat de kérosène pour la cokéfaction (193-279°C) dérivé du bitume de l'Athabasca. La dénitrification a été plus active que la désulfuration pour tous les catalyseurs non-activés $MoO_3/alumine$. L'activation des catalyseurs $MoO_3/alumine$ avec du nickel ou du cobalt a amélioré la désulfuration au point où les diverses réactions eurent lieu dans l'ordre suivant:

désulfuration > dénitrification > hydrogénation aromatique A partir des catalyseurs activés et non-activés, on compare les interactions de la dénitrification et de la désulfuration à différentes températures. Les résultats obtenus avec d'autres catalyseurs lors d'hydrotraitement de gaz-huile et de distillat à basse température sont aussi comparés. On a découvert que le genre de catalyseur, la température de réaction et les limites d'ébullition de la charge d'alimentation influencent la conversion et l'interaction de la désulfuration et de la dénitrification.

¹Chercheur scientifique, ²Chef de section, Recherche sur la catalyse et ³Chef intérimaire, Laboratoires de recherche énergétique, Ministère de l'Energie, des Mines et des Ressources, Ottawa, Canada.

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INTRODUCTION

Hydrotreating crude distillates involves competing reactions such as desulphurization, denitrogenation and hydrogenation. Individual studies of these reactions have been described in the literature for several model compounds^{1, 2, 3}. In a few cases, the interaction of these competing reactions has been reported^{4, 5}. However, it is difficult to simulate all the properties of petroleum using model compounds. It was felt that a study with distillate fractions was required to determine the interaction of these competing reactions in the industrial hydrotreating process. The purpose of the present paper is to describe the interaction of the competing reactions for various catalysts and process conditions as they would be encountered in a refinery application. A coker kerosene distillate, derived from Athabasca bitumen⁶, was hydrotreated on unpromoted and promoted catalysts at various reaction temperatures.

EXPERIMENTAL

The catalysts used in this study were unpromoted $MoO_3/alumina$ containing 3, 6, 9 and 12 wt % of MoO_3 and promoted $MoO_3/alumina$ composed of 1.1 wt % CoO - 2.2 wt % $MoO_3/alumina$, 1.1 wt % NiO - 2.2 wt % $MoO_3/alumina$ and 3 wt % CoO - 12 wt % $MoO_3/alumina$. The catalysts were prepared by spraying aqueous solutions of metal salts on alumina powder (a mixture of 20 wt % Continental Oil Company "Catapal SB" and 80 wt % "Catapal N" alumina monohydrate) in a mix-muller⁷. The impregnated mixtures were dried in air at $110^{\circ}C$ for 3 hours and then calcined at $500^{\circ}C$ for 3 hours. The calcined powder was mixed with 2 wt % stearic acid and pressed into cylindrical pellets (L = D = 3.2 mm) in a continuous pelletizing press. The pellets were recalcined at $500-550^{\circ}C$ for 4 hours to remove the stearic acid. A commercial catalyst, 3 wt % CoO - 12 wt % $MoO_3/alumina$ (Harshaw CoMo 0603T, 3.2 mm pellets), was also used.

The feedstock used for the hydrotreating study was a coker kerosene distillate (193-279[°]C) supplied by Great Canadian Oil Sands Ltd. of Fort McMurray, Alberta. The properties of the feedstock are listed in Table 1.

The reaction measurements were carried out in a bench scale continuous flow system ⁸ with the oil and hydrogen flowing up through a fixed bed of catalyst. The reactor was 0.025 m in internal diameter, 0.305 m long, and was made of 316 stainless steel. The reactor was filled sequentially from the bottom with 42 ml of berl saddles, 100 ml of catalyst pellets and 13 ml of berl saddles. The axial temperature profile was measured by a movable thermocouple in a thermocouple well located centrally in the reactor. The gas and liquid reaction products were separated in one of the two down-stream vessels. When steady-state conditions had prevailed for 1 hour, the product flow was routed to the second vessel where the liquid product was collected for subsequent analysis.

For all the experiments the reaction pressure, liquid space velocity and hydrogen flow rate were kept constant at $1.39 \times 10^7 \text{N/m}^2$ (2000 psig), $2h\overline{r}^1$ and $890.5 \text{ m}^3\text{H}_2/\text{m}^3$ oil (5000 scf H₂/bbl oil) respectively. The reaction temperatures were varied in the following sequence: 400, 420, 360, 320 and 400°C . The second run at 400°C was undertaken to check the series and confirm the stability of the catalyst.

The product samples were analysed for sulphur, nitrogen, aromatics, olefins and saturates. The conversions of higher boiling fractions to lower boiling fractions were determined by atmospheric distillation and the sulphur concentration by X-ray fluorescence⁹. The sulphur apparatus was calibrated using a series of oil samples analysed by the bomb sulphur technique⁷. The nitrogen content was measured using a hydrogenation-microcoulometric apparatus developed and manufactured by the Dohrmann Division of Envirotech Corp., Mountain View, California¹⁰. The Dohrmann procedure was developed for oils containing 10-100 ppm. The instrument was calibrated using pyridine, carbazole and acetanilide. The samples having higher nitrogen were diluted to reduce the nitrogen concentration to the working range of the instrument. The aromatics, olefins and saturates were separated using the ASTM standard FIA method¹¹. The conversion of fractions boiling above 220°C to fractions boiling below 220°C was calculated using the distillation data in the following equation:

(1)

$$X_{BP} = (100-F) - (100-P)$$

100 - F

where

P = wt % of product boiling below $220^{\circ}C$ F = wt % of feed boiling below $220^{\circ}C$

RESULTS AND DISCUSSION

Catalyst Pretreatment:

Preliminary studies showed that the catalyst pretreatment conditions are important in obtaining stable activity¹². The oxides in the catalysts are slowly converted to sulphides by the hydrogen sulphide produced from the sulphur compounds in the feed. Large variations in catalyst activity have been related to differences in the extent of catalyst sulphiding. It was found that a pretreatment with the coker kerosene feedstock at reaction conditions of $1.39 \times 10^7 \text{N/m}^2$ (2000 psig), 673°K , $2h^{-1}$ and $890.5 \text{ m}^3\text{H}_2/\text{m}^3$ oil (5000 cu ft/bbl) for 2 hours produced an adequate extent of catalyst stabilization.

Kinetics:

Mass transfer effects in catalysts are known to be important, particularly when hydrotreating high boiling petroleum fractions¹³. However, for the system used in this work there were strong indications that mass transfer was not controlling. The fact that the different catalysts had markedly different activities suggests that the effect of external mass transfer was negligible. Other experiments undertaken in this laboratory with a higher boiling gas-oil over extruded catalysts, having different dimensions (0.317 mm and 0.159 mm) but the same chemical composition, also produced the same activity¹⁴. This indicated that pore diffusional resistance was not controlling. Additional evidence for minimal diffusional effects is obtained when one compares the desulphurization and denitrogenation reaction data of two catalysts having the same metals content (Table 2). For 1.1 wt % NiO - 2.2 wt % MoO2/alumina, the denitrogenation conversion improved relative to desulphurization conversion with increasing temperature. For 3 wt % Mo0,/alumina, the desulphurization improved relative to denitrogenation conversion with increasing temperature. Such changes indicate that the differences in desulphurization and denitrogenation are not due to pore diffusion effects and can be attributed to kinetics.

The kinetics of denitrogenation, desulphurization and hydrogenation were studied on 3 wt % CoO - 12 wt % MoO₃/alumina catalyst. Denitrogenation followed first order kinetics (Figure 1 A) as reported by others¹⁵. The desulphurization followed second order kinetics (Figure 1 B). Similar desulphurization kinetics have been observed by others for high sulphur crudes ¹⁶. However, for low sulphur crudes, first order kinetics have also been reported ¹³. Beuther et al ¹⁶ and Schuit et al ¹⁷ suggest that desulphurization is first order for individual sulphur compounds and appears to be second order in a feedstock containing different types of sulphur compounds.

In the literature, the hydrogenation of aromatics is reported to follow reversible first order kinetics¹⁸. However, in the work discussed here the data did not fit zero order, first order irreversible, first order reversible or second order irreversible kinetics. This is probably due to the inhibition of the hydrogenation process by sulphur compounds. Voorhoeve and Stuiver¹⁹ reported that the hydrogenation of cyclohexane deviated from first order kinetics when carbon disulphide was present. They suggest that the hydrogenation sites are blocked by the preferential adsorption of CS₂. In the present case it is entirely possible that the sulphur compounds in the feedstock complicated the hydrogenation kinetics in this way.

Effects of Metal Oxide Concentration:

Several catalysts of varying Mo_3 concentration were evaluated. Figure 2 A shows that the % conversion of sulphur increases with Mo_3 concentration and reaches a plateau after 9 wt % Mo_3 . The denitrogenation results (Figure 2 B) show similar trends. However, at 320° C the rate of increase of conversion with metal oxide concentration was considerably greater for denitrogenation. At all temperatures studied the denitrogenation conversion is higher than desulphurization for the whole series of unpromoted $Mo_3/alumina$ catalysts. Qader et al²⁰ also reported that the denitrogenation conversion was higher than desulphurization conversion for low temperature distillate from coal tar (200 - 325° C, 0.83 wt % S and 0.40 wt % N) on a WS₂ catalyst at pressures above 1000 psig. However, Williams et al²¹ found that desulphurization conversions were greater than denitrogenation conversions for a heavy gas-oil (345 - 525° C, 3.59 wt % S and 0.38 wt % N) on unpromoted $Mo0_3/alumina$ catalysts at 2000 psig. The combined results indicate that the nitrogen in the heavier fractions is much more difficult to remove.

Hydrogenation of aromatics in coker kerosene distillate showed a trend different from denitrogenation and desulphurization (Figure 2 C). A 3 wt % MoO₃/alumina catalyst showed significantly higher hydrogenation conversion than the pure alumina support. However, further increases in MoO₃

concentration only slightly improved the hydrogenation of aromatics.

The conversion of fractions boiling above 220° C to fractions boiling below 220° C (equation 1) was measured for all catalysts studied. At 400° C, the per cent conversion increased with increasing metal oxide concentration up to 9 wt % $Mo0_3$ and decreased above 9 wt % (Figure 2 D). At 320° C, the maximum was at 6 wt % $Mo0_3$. The findings of Seshadri et al²² may provide a partial explanation for the maximum in conversion. They found that, for $Mo0_3$ /alumina catalysts, the catalytic activity in vapour-phase aldol condensation of n-butyraldehyde and the esr signals (indicating Mo^{5+} concn.) showed maxima at 9 wt % in $Mo0_3$, suggesting that Mo^{5+} centres may be responsible for molecular weight reduction. Desulphurization, denitrogenation and hydrogenation conversions did not show these maxima. It would appear that quite different sites are involved for molecular weight reduction than for the refining reactions.

Effects of Promoters:

At low reaction temperatures the addition of promoters to $MoO_3/$ alumina catalysts affected the desulphurization, denitrogenation and hydrogenation processes differently. Comparing catalysts containing 2.2 wt % $MoO_3/$ alumina and 1.1 wt % CoO - 2.2 wt % $MoO_3/$ alumina, the conversions at $320^{\circ}C$ for desulphurization, denitrogenation and aromatic hydrogenation increased from 20 to 76.7%, 30 to 48% and 31 to 43% respectively (Figures 1 A, 1 B and 1 C and Table 2). A laboratory catalyst containing 12 wt % $MoO_3/$ alumina was compared with a commercial catalyst having 3 wt % CoO - 12 wt % $MoO_3/$ alumina. In this case the conversions improved from 38.8 to 93.5%, 57.2 to 79.3% and 40.5 to 49.2% respectively. The promotion of $MoO_3/$ alumina with nickel showed similar increases in desulphurization, denitrogenation and hydrogenation (Table 2). At $320^{\circ}C$ the increase in conversions due to promoter addition varied in the following order:

desulphurization > denitrogenation > aromatic hydrogenation The results show that the addition of cobalt or nickel to molybdenum/alumina significantly increases the selectivity for the desulphurization reaction in particular. The promoter would appear to change the chemisorption characteristics of sulphur compounds in such a way as to selectively improve desulphurization.

Similar results were obtained by Williams et al²¹ showing that the addition of a promoter selectively increased the desulphurization conversion. They found that, at 380° C for 3 wt % CoO - 12 wt % MoO₃ catalysts, the desulphurization and denitrogenation conversions were 78% and 24% respectively compared with 35% and 16% for 12 wt % MoO₃ catalysts. The important point to note is that the different types of compounds present in the low boiling coker kerosene and the heavy gas-oil feedstocks did not affect the promotion of desulphurization by cobalt or nickel.

Comparison of Denitrogenation and Desulphurization at Different Temperatures:

The rate constants calculated using the first order equations for denitrogenation and the second order equations for desulphurization are shown in Arrhenius plots (Figures 3 & 4). The results obtained for both promoted and unpromoted catalysts are given. It is seen from Figure 3 that the activation energies (proportional to the slopes of the lines) for denitrogenation decrease with increasing concentrations of unpromoted MoO3 on alumina. On the other hand, for desulphurization (Figure 4) the activation energies increase with increasing MoO $_3$ concentration. Cremer²³ suggests that one of the reasons for variations in activation energies and frequency factors is the difference in the strength of adsorption. In the present work, the variations in activation energies also suggest that the differences in denitrogenation and desulphurization conversions on MoO3/alumina are caused by differences in strengths of chemisorption of sulphur and nitrogen compounds on the catalyst. When Ni or Co promoter was added the activation energies for both denitrogenation and desulphurization decreased, but there was a large increase in the frequency factor for the desulphurization reaction compared with the denitrogenation reaction.

Satterfield et al⁴ suggest that the interaction between desulphurization and denitrogenation is temperature-dependent. They studied the desulphurization of model compounds such as thiophene and the denitrogenation of pyridine on commercial CoMo, NiMo and NiW sulphided catalysts at temperatures up to 425° C and pressures up to $1.1 \times 10^{3} \text{ kN/m}^{2}$ (11 atm). At low temperatures, thiophene inhibited denitrogenation and at high temperatures the sulphur compounds enhanced the denitrogenation. The results obtained with distillate fractions and the <u>promoted catalysts</u> both in the present work and by Williams et al²¹ support the observations of Satterfield et al⁴. With the coker distillate over Co and Ni promoted catalysts, the desulphurization was much

higher than denitrogenation at 320[°]C, but the denitrogenation conversion improved relative to desulphurization with increasing temperature (Table 2). Also, Williams et al found that for similar promoted MoO₃/alumina catalysts (Table 3), the differences between the conversions for denitrogenation and desulphurization decreased at higher reaction temperatures.

With the <u>unpromoted catalysts</u>, on the other hand, the hydrotreating data shows a reverse trend. The results for the unpromoted catalysts (Figures 1 A, 1 B and Table 2) with the coker distillate and of Williams et al on gas-oil (Table 3) show that with increasing reaction temperature, the desulphurization reaction conversion improves relative to denitrogenation conversion. A comparison of relative conversions for desulphurization and denitrogenation at different temperatures definitely indicates that the interaction of these reactions depends on whether the catalyst is promoted or not.

CONCLUSIONS

Comparison of molecular weight reduction and the refining reactions on unpromoted MoO₃ catalysts indicates that quite different active sites are involved for molecular weight reduction than for the refining reactions. Variations in trends for apparent activation energies of desulphurization and denitrogenation on unpromoted catalysts indicate differences in chemisorption characteristics of nitrogen and sulphur compounds.

With the promoted catalysts, the kinetics observed with the distillate fractions generally agrees with the results of the pure compound studies. The trend of the data was the same regardless of the boiling range of the distillate. However, in the case of unpromoted catalysts, the boiling range of the feed-stock has a considerable influence on the reaction rates. Comparison of the high pressure data for coker kerosene reported here, the high pressure data of Williams et al²¹ and the low pressure data reported by Satterfield et al⁴ indicates that with increasing reaction temperature, the presence of sulphur compounds enhances denitrogenation relative to desulphurization on promoted catalysts but not on unpromoted catalysts.

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TABLE 1

Properties		Coker Kerosene Distillate
Boiling range,	°c	193 - 279
Specific Gravity 60/60	° _F	0.871
Sulphur,	wt %	2.32
Nitrogen,	ppm	430
Pour Point,	° _F	Below -60
Cloud Point,	° _F	Below -60
Flash Point,	°F	116
Vanadium,	ppm	0.40
Nickel,	ppm	0.36
Iron,	ppm	0.50
Ramsbottom Carbon		
Residue (10% bottoms),	wt %	0.29
	.	50
Aromatics + olefins,	vol %	58
Saturates,	vol %	42

PROPERTIES OF THE FEEDSTOCK

TABLE 2

DESULPHURIZATION, DENITROGENATION AND HYDROGENATION OF COKER KEROSENE ON PROMOTED AND UNPROMOTED CATALYSTS

Catalysts			Unpromoted											
Concn metal oxides	s 1.1 wt % CoO-2.2 wt % MoO ₃ /alumina 1.1 wt % NiO-2.2 wt % MoO ₃ /alumina 3									3 wt % Mo0 ₃ /alumina				
Reaction temperature, ^O C	320	360	400	420	320	360	400	420	320	360	400	420		
% desulphurization	76.7	92.7	96.6	96.6	71.1	94.0	96.1	96.1	26.7	56.5	87.1	92.2		
% denitrogenation	48.1	82.3	95.3	97.7	50.9	86.1	96.3	98.6	34.4	68.6	91.2	94.0		
% hydrogenation	43.2	54.1	56.6	56.8	42.9	52.2	55.8	57.5	34.9	43.4	55.3	56.3		

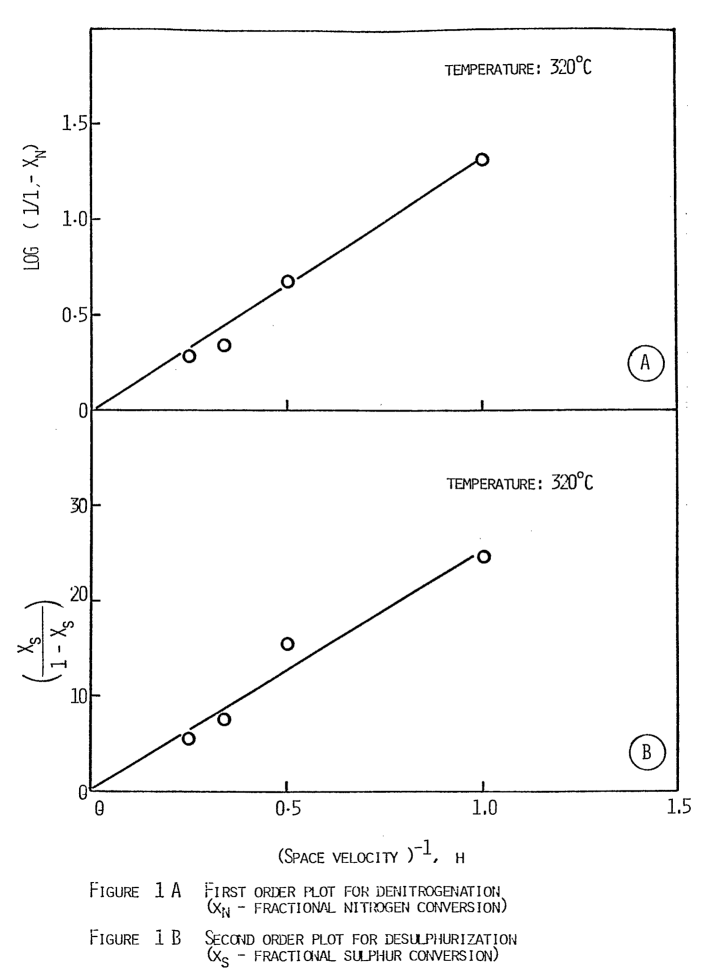
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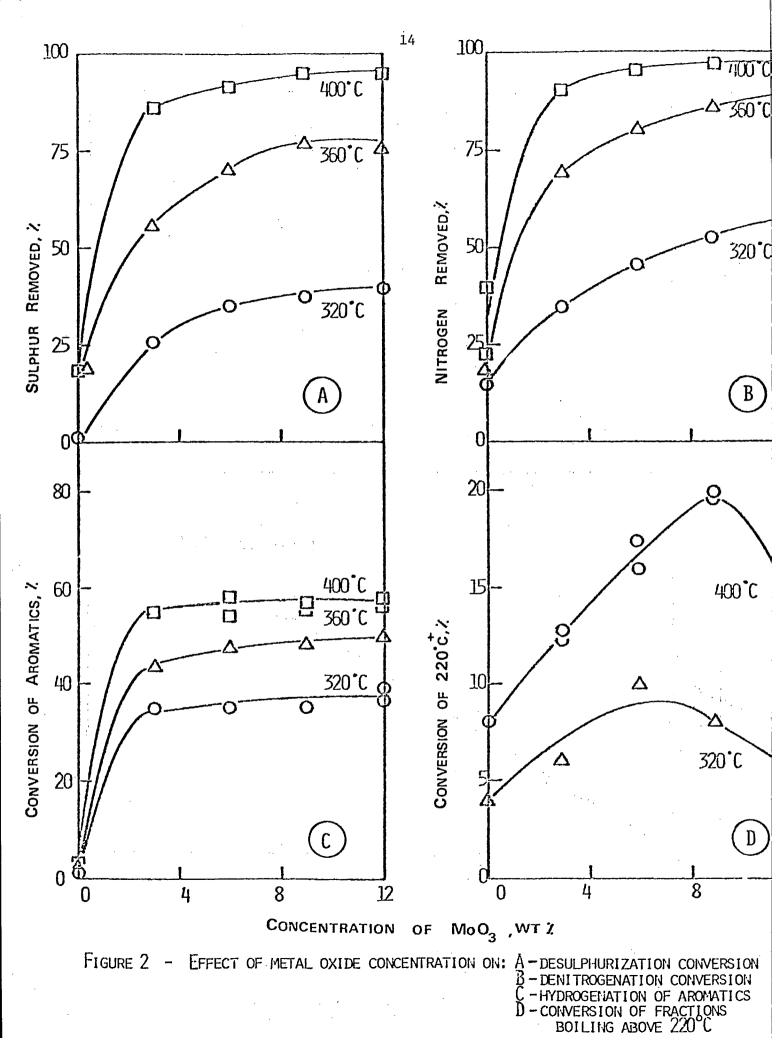
TABLE 3

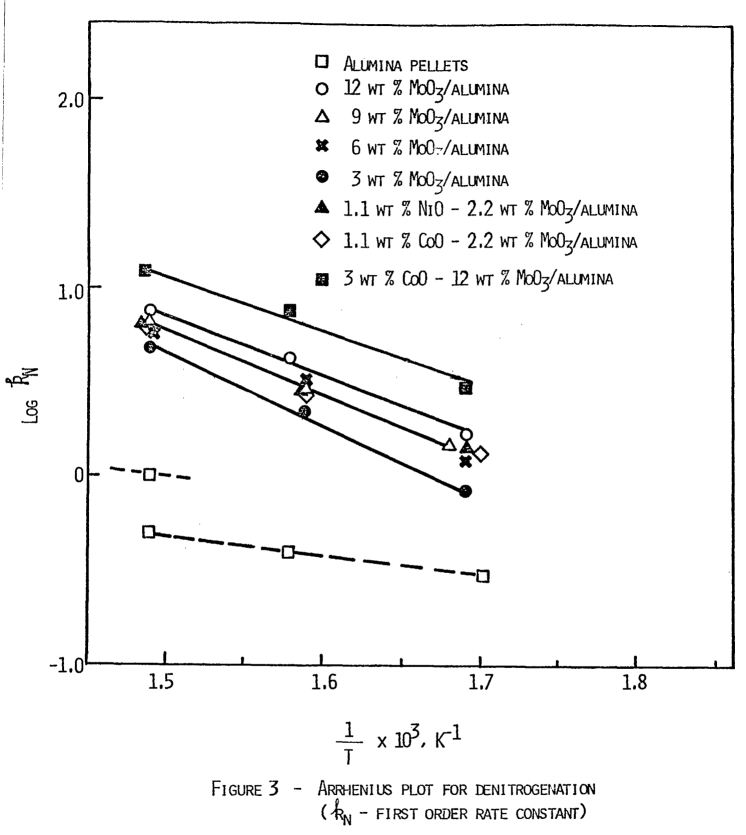
GAS OIL HYDROTREATING ON PROMOTED AND UNPROMOTED CATALYSTS

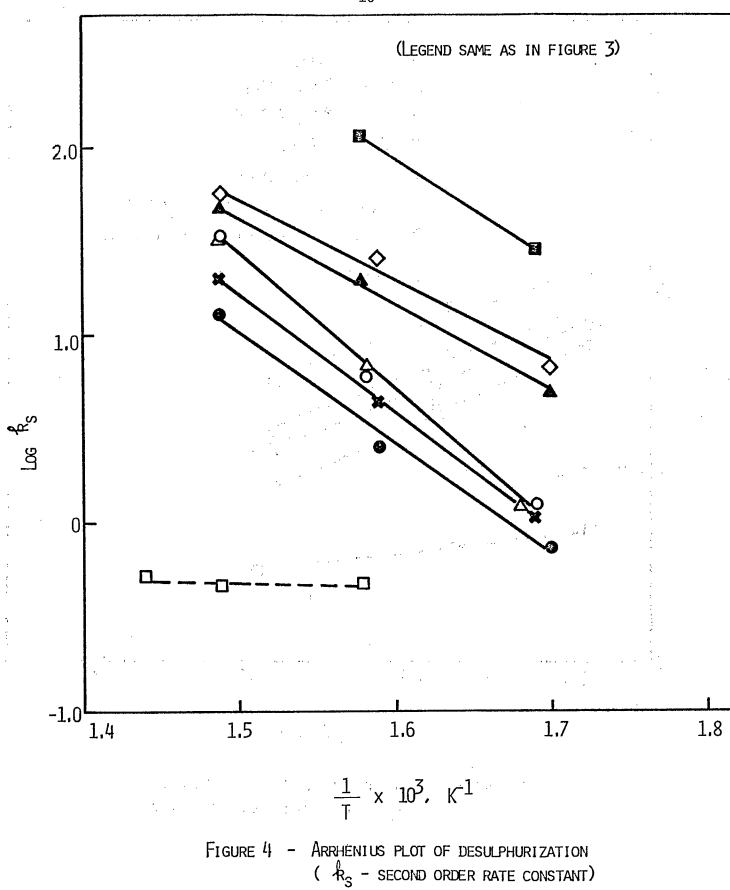
Williams et al (21)

Catalyst	Promoted				Unpro	omoted					
Con c n metal oxides	3 wt % Co0 -	12 wt % MoO ₃	3 wt	^{% Mo0} 3	6 wt 2	^{% MoO} 3	9 wt	% Mo0 ₃	12 wt % MoO ₃		
Reaction temperature, ^O C	380	450	380	450	380	450	380	450	380	450	
% desulphurization	78	93	26	74	29	81	36	87	35	85	
% denitrogenation	24	73	0	24	0	45	8	61	16	63	
					,						









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