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Derbyshire, F.J.; de Beer, V.H.J.; Abotsi, G.M.K.; Scaroni, A.W.; Solar, J.M.; Skrovanek, D.J.

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THE INFLUENCE OF SURFACE FUNCTIONALITY ON THE ACTIVITY OF CARBON-SUPPORTED CATALYSTS

F.J. DERBYSHIRE, V.H.J. de BEER, G.M.K. ABOTSI, A.W. SCARONI, J.M. SOLAR and D.J. SKROVANEK

Department of Materials Science and Engineering, Fuel Science Program, The Pennsylvania State University, University Park, PA 16802, USA.

aLaboratory for Inorganic Chemistry and Catalysis, Eindhoven University of Technology, P.O. Box 513, 5600 MB Eindhoven, The Netherlands.

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ABSTRACT

The aim of this research is to investigate how the presence of surface functional groups can influence the activity of carbon-supported MoS2 catalysts for coal asphaltene hydrogenation. Porous carbons were subjected to various chemical treatments in order to introduce oxygen and nitrogen surface functionalities prior to impregnation with ammonium tetrathiomolybdate. Supports and catalysts were examined by FTIR. Preoxidation of polymer-derived carbons lowered catalyst activity whereas preoxidation of a carbon black composite support increased it. Similar mixed results have been reported in the literature. The modified catalytic activity cannot be unequivocally attributed to surface oxygen. Qualitatively, it is considered that these groups exert some influence, but it is the structure of the carbon which is ultimately the controlling factor. In some cases, oxidation may introduce sites which promote interaction with metal species and, in others, oxidation may modify or destroy favourable sites which are already extant.

Preoxidation was found to have a distinct influence in enhancing catalyst activity. The presence of nitrogen-containing surface groups may provide preferential sites for the adsorption of Mo species.

INTRODUCTION

In previously reported research [1], the hydrodesulfurization (HDS) activities and the propensities for carbon deposition were determined for a series of catalysts prepared on carbon and alumina supports. The results showed that the HDS activities were higher and the coking propensities were lower on the carbon than on the alumina supported catalysts and that, for the former, the activity was strongly dependent upon the nature of the carbon.

It is obviously of interest to identify those characteristics of carbon supports which are influential in determining catalyst properties. However, the carbons used in the earlier research were of different origin and varied in their chemical structure, textural properties and content of impurities. Further interpretation of the results was thus precluded in the absence of a sound basis for the comparison of data.

In the light of present knowledge, it seems that, ceteris paribus, catalyst
activity is strongly influenced by the interaction between the carbon surface and the deposited metal species. This interaction can be instrumental in controlling the dispersion of the active species and valence state of the metal.

For a given precursor compound, a review of the literature suggests that catalyst activity is related to active sites on the carbon surface or to the presence of surface functional groups.

Ehrburger and co-workers [2] observed that the dispersion of platinum on carbon black particles increased with the extent of gasification of the support. They proposed that the effect of gasification was to increase the heterogeneity of the carbon surface which raised the potential energy barrier for platinum diffusion. The surface oxygen functionality did not appear to influence metal dispersion: similar dispersions were measured for platinum loaded on a sample of carbon which had been burned-off to 21.4% and for a sample of the same material which was first reduced in hydrogen at 950°C to remove oxygen complexes before platinum addition. The catalytic activities of these materials were not determined. These results suggest that it is principally carbon active sites which influence metal dispersion. That is, sites which "exist on the carbon surface where the valency is not satisfied. In a "clean" carbon surface, these sites would be located on the edges of the exposed layer planes as well as at points of imperfections in the structure including vacancies, dislocations, and steps on the outer basal plane" [3].

Other results suggest that the type and concentration of surface functional groups can modify catalyst activity. It has been shown that the catalytic activity of unloaded carbons, for reactions such as SO₂ oxidation, can be greatly enhanced by treatment with NH₃ or HCN at elevated temperatures. The change in activity is attributed to the incorporation of nitrogen into the carbon surface [4]. No results have yet been reported for the catalytic behavior of metals supported on such carbons.

Ryan and Stacey [5] found that active carbons were selective catalysts for the reaction of phosgene and formaldehyde to produce dichloromethane. They attributed the catalytic activity to the presence of polar acid or base sites on the carbon surfaces.

Other work has shown that the treatment of active carbons with phosphoric acid induces catalytic activity for the conversion of methanol to C₁-C₄ paraffins and olefins [6].

Vissers et al. [7] applied a number of oxidative treatments to carbon supports, prior to loading with Mo or Co precursors, in an attempt to alter the dispersion of the active species. All of the treatments resulted in an increased concentration of oxygen-containing functional groups on the carbon surface. The effects on the activities of the catalysts for thiophene HDS were mixed; preoxidation enhanced activity in some cases and suppressed it in others.
The available evidence indicates that the nature of the carbon surface and the presence of functional groups can have an important influence upon the properties of carbon-supported metal catalysts.

In this paper we present the results of a program of research which was intended to investigate the importance of surface functional groups on the activity of carbon-supported MoS\(_2\) catalysts. For this purpose, a single metal salt precursor and a single parent carbon were selected. Samples of the carbon were subjected to a number of chemical treatments in order to introduce different oxygen and nitrogen-containing functional groups.

**EXPERIMENTAL**

**Carbon supports**

The carbon used in this study, Ambersorb XE-340 (supplied by courtesy of Rohm and Haas Co.) is produced by the carbonization of a sulfonated styrene/divinylbenzene copolymer, and can be made in high purity with reproducible chemical and physical properties. It has a pore size distribution suitable for the hydrogenation of high-boiling liquids since a high proportion (ca. 70\%, by the method of Metcalfe et al. [8]) of the pore volume is contained in mesopores (2-50 nm diameter). Surface area measurements by BET using N\(_2\) as the adsorbate gas gave values of 399 m\(^2\) g\(^{-1}\), which is in good agreement with the manufacturer's data. Using CO\(_2\) as the adsorbate, the surface area was determined to be 635 m\(^2\) g\(^{-1}\).

A comparison of the two surface areas suggests that the carbon contains an appreciable micropore volume (0.8 - 2.0 nm diameter) [9]. The carbon particles are spherical in form (0.3 - 0.8 mm diameter) and hard and durable, with a crush strength greater than 3 kg/particle.

A few additional experiments were carried out using a carbon black composite support which was fabricated by the carbonization of a mixture of graphitized carbon black filler and polyfurfuryl alcohol binder. The carbon black particles were first graphitized by heat treatment to 2500°C for 2 h in flowing argon. The carbon black/binder mixture was subsequently carbonized by heating to 1200°C for 2 h in an atmosphere of flowing nitrogen. The preparation of carbon black composites has been described by Schmitt et al. [10].

**Support treatment**

Samples of Ambersorb XE-340 were subjected to a number of different treatments intended to introduce oxygen or nitrogen functionalities to the carbon surface. Following each treatment, the carbons were repeatedly washed with distilled, deionized water. The treatments and their effects on the elemental composition and surface areas of the carbons are presented in Table 1.

All of the oxidative treatments increased the oxygen content of the carbons. The HNO\(_3\) and boiling concentrated H\(_2\)SO\(_4\) treatments introduced the highest amounts of oxygen. The acid treatments also increased the nitrogen or the sulfur contents of the supports.
<table>
<thead>
<tr>
<th>Treatment</th>
<th>Elemental analysis/wt%</th>
<th>S.area /m² g⁻¹ (N₂ BET)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C</td>
<td>H</td>
</tr>
<tr>
<td>None</td>
<td>90.2</td>
<td>3.3</td>
</tr>
<tr>
<td>Boiling saturated</td>
<td>90.0</td>
<td>3.8</td>
</tr>
<tr>
<td>K₂Cr₂O₇ soln, 2 h</td>
<td>75.7</td>
<td>3.1</td>
</tr>
<tr>
<td>Boiling conc. KMnO₄ soln, 2 h</td>
<td>64.9</td>
<td>2.3</td>
</tr>
<tr>
<td>Conc. H₂SO₄, 2 h at room temp.</td>
<td>84.4</td>
<td>3.3</td>
</tr>
<tr>
<td>Boiling conc. H₂SO₄, 2 h</td>
<td>68.2</td>
<td>2.8</td>
</tr>
<tr>
<td>NH₃ at 400°C, 1.5 hᵃ</td>
<td>89.2</td>
<td>3.5</td>
</tr>
<tr>
<td>NH₃ at 600°C, 1.5 hᵃ</td>
<td>91.7</td>
<td>2.6</td>
</tr>
<tr>
<td>Analysis of coal liquids</td>
<td>89.8</td>
<td>7.1</td>
</tr>
</tbody>
</table>

ᵇ by difference  
ᶜ N.D. = not determined.
The treatments with potassium permanganate and potassium dichromate served principally to oxidize the carbon, the former reagent being the stronger oxidant of the two. In both cases, despite extensive post-treatment washing with distilled, deionized water, residual potassium was left on the carbon, as indicated by elemental analysis. Examination by X-ray diffraction confirmed that some potassium was present as potassium permanganate on the KMnO$_4$-treated carbon.

These chemical treatments were intended to introduce oxygen functionalities while producing only minimum change to the textural properties. (Oxidation through partial gasification can result in substantial carbon loss and can cause alterations to surface area and pore size distribution). As can be seen from Table 1, this aim was unsuccessful for those treatments with nitric and sulfuric acids which effected drastic reductions in surface area. Evidently, the attack by these acids was too severe and must have resulted in a collapse of the pore structure. Surprisingly, however, the structural integrity of the carbon particles was maintained.

The general procedure for introducing nitrogen functionality was adopted from Boehm et al. [4]. Reaction with ammonia at both 400 and 600°C increased the nitrogen content of the carbon but otherwise had little effect upon the elemental composition. There was some weight loss by gasification during these reactions (1% at 400°C and 4.3% at 600°C) with corresponding reductions in surface area.

**Catalyst preparation and characterization**

The carbons were impregnated with 3% wt Mo from solution using the incipient wetness technique. A solution of ammonium heptamolybdate was prepared by dissolution of the salt in a mixture of 90% H$_2$O/10% EtOH. The presence of alcohol was found necessary to assist in wetting the carbon surface which is somewhat hydrophobic. The solution was then saturated with H$_2$S at room temperature to form the tetrathiaammonium salt, (NH$_4$)$_2$MoS$_4$. Thus, the active catalyst form can be produced in situ, thereby eliminating the need for catalyst presulphiding. To achieve the required loading, repeated impregnations were required with the preparation being dried in vacuum at 100°C between each step. All of the prepared catalysts were used without further heat treatment.

The modified carbons, before and after loading, were analyzed by Fourier Transform Infrared Spectroscopy (FTIR) in diffuse reflectance (Digilab, Model FTS-15E). The samples were prepared by first grinding to less than 0.3 mm dia. A solid mixture of 5% wt of the sample with KBr was intimately mixed by grinding in a mortar and pestle and the resulting mixture was placed in a Barnes diffuse reflectance unit.
### TABLE 2
Principal changes in carbon structure observed by FTIR (Carbon: Ambersorb XE-340)

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Unloaded</th>
<th>Loaded (3% Mo)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) None</td>
<td>(a1) Predominantly aromatic, =CH groups in evidence in the 1600-1300 cm(^{-1}) regions.</td>
<td>(a2) Intensity changes in 1600-1100 cm(^{-1}) and 900-600 cm(^{-1}) regions</td>
</tr>
<tr>
<td>h) (\text{K}_2\text{Cr}_2\text{O}_7)</td>
<td>(b1) Similar to (a2)</td>
<td>(b2) No significant changes on Mo addition</td>
</tr>
<tr>
<td>c) (\text{KMnO}_4)</td>
<td>(c1) Similar to (a2)</td>
<td>(c2) Intensity increases in 900-600 cm(^{-1}) region (C-H out-of-plane bending)</td>
</tr>
<tr>
<td>d) (\text{HNO}_3)</td>
<td>(d1) Major reduction in aromatic character. Substantial increase in C=O band (1720 cm(^{-1})). Introduction of intense bands at 1530 cm(^{-1}) and 1340 cm(^{-1}) indicative of presence of NO(_2) groups. Major changes in 900-600 cm(^{-1}) region.</td>
<td>(d2) No significant changes on loading</td>
</tr>
<tr>
<td>e) cold (\text{H}_2\text{SO}_4)</td>
<td>(e1) Some loss of aromatic character. Increase in C=O band. Major spectral changes in regions 1600-1100 cm(^{-1}) and 900-600 cm(^{-1}). Introduction of band envelope 1360-1300 cm(^{-1}) and band at 1170 cm(^{-1}) probably attributable to SO(_2) groups.</td>
<td>(e2) Increase in intensity of 1360-1100 cm(^{-1}) envelope and bands in 900-600 cm(^{-1}) region</td>
</tr>
<tr>
<td>f) hot (\text{H}_2\text{SO}_4)</td>
<td>(f1) Generally similar to (e1) but much more dramatic changes in intensity of bands described</td>
<td>(f2) Change in character of 1360-1300 cm(^{-1}) and 900-600 cm(^{-1}) regions</td>
</tr>
<tr>
<td>g) (\text{NH}_3), 400(^\circ)C</td>
<td>(g1) Very similar to (a2)</td>
<td>(g1) No significant change on loading</td>
</tr>
<tr>
<td>h) (\text{NH}_3), 600(^\circ)C</td>
<td>(h1) Some loss of aromatic character. Increase in intensity and spectral changes in 1600-1100 cm(^{-1}) and 900-600 cm(^{-1}) regions</td>
<td>(h2) Some modification of intensity between 1400-1200 cm(^{-1}). No other significant changes.</td>
</tr>
</tbody>
</table>
The prepared catalysts were examined for their relative activities in the hydrogenative upgrading of coal liquids. The feed was a 454+°C recycle solvent fraction from a coal liquefaction pilot plant (Lummus Integrated Two-stage Liquefaction unit processing subbituminous coal) and contained 38.2% asphaltenes (hexane-insoluble). The elemental analysis of this material is shown at the foot of Table 1. The total ash content was determined to be 0.05 wt%, although the concentrations of specific metals have not been determined.

Reactions were carried out in a batch tubing bombs of about 25 cc capacity. The following conditions were used: 5.0 g feed, 0.5 g catalyst, 7 MPa H₂ (cold). Approximately 0.02 µ of carbon disulfide were added to the reaction mixture to maintain the catalyst in sulfided form. The sealed reactor was immersed in a fluidized sandbath heater, at 200°C and held at this temperature, under agitation, for 30 min to convert the CS₂ into CH₄ and H₂S. The presence of H₂S is desirable for maintaining the catalyst in the sulfided form. The temperature was then raised to 450°C over a period of 30 min and held for a further 60 min at this temperature before removing the reactor and quenching in cold water.

The gases were vented from the bomb and were analyzed by gas chromatography. The contents of the bomb were washed out using tetrahydrofuran (THF) and filtered to recover the remaining solids. The THF was removed from the filtrate by vacuum evaporation and the material was then refluxed in hexane for 24 h to obtain a partition into hexane-soluble (oils) and hexane-insoluble (asphaltene) fractions. The asphaltene conversion was calculated from the formula 100 x (percent asphaltenes in feed - percent asphaltenes in product)/(percent asphaltenes in feed).

RESULTS AND DISCUSSION

FTIR analysis of supports and catalysts

The main features observed in the FTIR spectra of the carbons and of the prepared catalysts are summarized in Table 2. After impregnation of the untreated carbon there was evident modification of the absorption bands at 812, 1160, 1240, 1320, 1420 and 1440 cm⁻¹, Figure 1. Those at 810, 1160 and 1240 cm⁻¹ may correspond to CH substituents in the para position on a benzene ring. Similarly, the bands at 1320 cm⁻¹ and 1420 cm⁻¹ may represent =CH, and =CH₂, respectively. It may be implied that the impregnated metal salt or its decomposition products become chemically associated with aromatic ring substituents or to conjugated double bonds on the carbon surface.

After treatment with both K₂Cr₂O₇, and K₂MnO₄, the carbon FTIR spectra were both similar to that of the loaded parent carbon. Upon loading these oxidized carbons, there was no apparent change in the spectrum of the carbon reacted with potassium dichromate, and only minor changes for that treated with potassium permanganate. There was therefore little, if any, evidence from the FTIR spectra of any significant interaction between the carbon and the Mo surface species on these oxidized supports.
As would be anticipated, the HNO₃ and H₂SO₄ treatments caused more pronounced changes to the carbon structure. Notably there was a loss of aromaticity, as evidenced by a reduction in the intensity of the absorption band at 3070 cm⁻¹ (aromatic C–H stretch), and an increase in the intensity of the C=O band, at 1720 cm⁻¹. Further, the HNO₃ treatment introduced NO₂ groups while both the cold and hot H₂SO₄ introduced SO₄²⁻ groups to the carbon surface. The hot H₂SO₄ also produced absorption bands which may be due to S=O. The extensive structural changes in the Ambersorb carbon, which were brought about by these acid treatments and which are evident from the FTIR spectra, are consistent with the drastic losses in surface area discussed above. Following loading, there was little change in the spectrum of the HNO₃ treated carbon in contrast to those treated with H₂SO₄ in which distinct spectral changes were observed upon impregnation, consistent with there being some interaction between the carbon surface and the impregnated metal species.

The spectrum of the carbon treated in ammonia at 400°C was similar to those obtained after oxidation by the potassium salts. After treatment with NH₃ at 600°C, there were more distinct alterations to the carbon structure as evidenced by a loss of aromaticity and marked changes in the 600-900 and 1100-1600 cm⁻¹ regions. The intensities of the bands in these regions increased after loading. In previous work by Boehm et al. [4], it was suggested that the reaction of carbons with NH₃ or HCN at elevated temperatures can lead to the incorporation of nitrogen into the aromatic layers as a pyridine-like species. The lone pairs of electrons of such hetero-atomic nitrogen species could be favorable sites for the interaction of Mo species with the support. If such species were produced in these experiments, they would not be easily observed by FTIR because of the low nitrogen concentration and the coupling of C=N vibrations with C=C aromatic vibrations.

To pursue this aspect further, a sample of poly-4-vinylpyridine (containing
12.3% N) was loaded with 3% Mo and examined by FTIR. As shown in Figure 2, upon impregnation there was extensive broadening of the absorption bands which represent the vibrations of the C=N and C=C bonds in the pyridine ring. There was, in addition, a shift in the absorption frequency to lower wave numbers. Both of these features are indicative of there being a strong chemical interaction between Mo species and delocalized electrons (due to C=C and C=N bands) and/or the lone pairs of electrons on the nitrogen atoms in the polymer.

The implication from the foregoing is that, if similar nitrogen functionalities were present on the carbon, they could provide preferred sites for the adsorption of Mo species. The specific nature of the nitrogen-containing functionalities is not, however, known. Furthermore, the type and concentration of nitrogen (or other) groups which can be introduced by chemical reaction will evidently be dependent upon the carbon structure and will be affected by factors such as the degree of structural order, crystallite size, the concentration of dislocations and vacancies and the proportion of edge to basal planes which are exposed at the surface. Consequently, subjecting different carbons to the same chemical treatments will not necessarily lead to similar results.

**Effects of preoxidation on catalytic activity**

The activities of the catalysts prepared on the parent and the preoxidized carbons, as well as that of an equivalently loaded γ-Al₂O₃ catalyst, are presented in Figure 3. The asphaltene conversions and C₁-C₄ yields obtained in a blank run using the unloaded and untreated support were 8.5% and 7.5%, respectively. Also shown in the figure are the percentage increases in catalyst weight due to the deposition of carbonaceous (THF insoluble) reaction products. The nature of these
deposits has not yet been determined and is the subject of continuing studies.

The reported activities, having been determined under similar reaction conditions, are related to the unit masses of catalyst and of molybdenum. Of course, there are other ways of expressing the catalyst activities, the activity per unit area of the support surface being the most obvious. In this case, there would be a change in the ranking order. However, it is probable that there are significant differences in the dispersion of the active molybdenum species on the variously treated supports. Since the dispersion cannot be quantified, it is considered that a more realistic practical means of comparing activities is in relation to the mass of catalyst and this is the basis used in the following discussion.

The principal feature of the data shown in Figure 3 is that preoxidation of the carbon supports reduced the activities of all of the catalysts for asphaltene-oil conversion. For those catalysts prepared on the acid treated supports, the low catalyst activities can be explained by the greatly reduced surface areas of the carbons (Table 1).

For the carbons preoxidized with $K_2Cr_2O_7$ and $KMnO_4$, the reductions in catalyst activity cannot be attributed to changes in surface area alone. One possible explanation is that the effects of oxidation were to remove or modify the surface groups present on the untreated Ambersorb, such as aromatic ring substituents and conjugated double bonds, which are preferred sites for interaction with Mo species. The reduced interaction would correspond to the observations made from the FTIR spectra of the carbons and catalysts. In turn, it would be anticipated that a weaker interaction between the surface and metal species would result in a lower dispersion, consistent with the decrease in activity.

However, the possible influence of the residual potassium on catalyst activity cannot be ignored. Bridgewater et al. [13] found that small quantities of potassium
acted as a poison to carbon-supported molybdenum catalysts for synthesis gas reactions. The authors noted that this finding was in contrast to other published observations.

In their study of thiophene hydrodesulfurization (HDS) over MoS₂/C catalysts, Vissers et al. [7] found that nitric acid pretreatment increased the activities of some carbon supported molybdenum catalysts and reduced them in other cases. They suggested that when the activity was enhanced, the high concentration of oxygen functional groups had a promotional effect on metal dispersion; for the cases where low activity was observed, the molybdate ions were weakly bonded to the carbon surface due to low concentrations of oxygen functional groups. This, they suggested, favored the mobility of the Mo species and resulted in sintering under reaction conditions. The sintering caused poor dispersion and hence low HDS activity.

However, these workers did not present information on the dispersion of the surface metal species. Neither was it apparent that the support surface areas had been determined after reaction with HNO₃, the conditions for which were at least as severe as those employed in the research reported here. It is therefore possible that changes in support textural properties may account, at least in part, for the apparently unpredictable effects of preoxidation upon catalyst activity. During carbon oxidation, there is usually an increase in surface area due to the generation of new porosity. This is balanced by the loss of carbon and the enlargement of existing pores. Eventually the surface area passes through a maximum and begins to decrease [12]. Depending upon the susceptibility of the different carbons to HNO₃ oxidation it is quite possible that, in some cases, the final surface area was greater than that of the starting material and, in others, it was lower.

From the foregoing, the thesis that the introduction of oxygen functionality can promote catalytic activity, possibly through providing preferred sites for the attachment of metal species and, thus, enhancing dispersion, cannot be unequivocally established since there are other possible explanations for the changes in activity caused by support oxidation.

The results obtained with catalysts prepared on the carbon black composite (CBC) support are presented in Table 3 and appear to show more directly that oxygen surface functionality can be instrumental in increasing catalytic activity. The catalyst prepared on the parent CBC support possessed virtually no activity for asphaltene conversion. Following reaction in boiling conc. HNO₃ for 2 h, the oxygen content was increased from < 0.3 to 2.3% with an increase in surface area of around 30%. The support pretreatment also effected a substantial increase in the activity of the prepared catalyst and the full extent of this increase cannot be simply attributed to the increase in surface area.

From the results shown in Table 1, it is likely that the acid treatment also introduced nitrogen functionality. The FTIR spectrum of the similarly treated Amberisorb carbon indicated that much of this nitrogen was in the form of nitrogen-
<table>
<thead>
<tr>
<th>Support</th>
<th>Oxygen content /wt%</th>
<th>% Asphaltene conversion</th>
<th>% C&lt;sub&gt;1&lt;/sub&gt;-C&lt;sub&gt;4&lt;/sub&gt;</th>
<th>% Catalyst weight gain</th>
<th>S. area/m² g⁻¹ (N₂ BET)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) Graphitized CBC</td>
<td>&lt; 0.3</td>
<td>0.0</td>
<td>2.9</td>
<td>5.0</td>
<td>120</td>
</tr>
<tr>
<td>(b) After oxidation of (a) for 2 h in boiling conc. HNO₃</td>
<td>2.3</td>
<td>47.6</td>
<td>3.5</td>
<td>4.0</td>
<td>160</td>
</tr>
</tbody>
</table>

*aReaction conditions are described in the text.*
oxygen complexes. At this point, it cannot be stated whether the increase in catalyst activity is related to the presence of these particular functionalities.

Analysis of the carbons and catalysts by FTIR did not provide any further information. All of the spectra were featureless, which is believed to be due to the low concentration of functional groups, the graphitic nature of the carbon and the high absorption of the infrared beam.

In synopsis, there is some qualitative evidence to advocate that oxygen functionality might advantageously alter catalyst activity. The results presented and discussed also indicate that carbon surface oxidation can, in some cases, reduce catalyst activity. It is suggested that the effect of surface treatment is dependent upon the structure of the parent carbon and that this, essentially, is the controlling factor. For example, it appears that certain surface sites, extant on the parent Ambersorb carbon, provided a more favorable interaction with the deposited metal species than those resulting from oxidative treatment.

**Effects of ammonia pretreatment on catalytic activity**

In Figure 4, the activities of catalysts prepared on the ammonia pretreated carbon supports are compared with those of the untreated support and of the γ-Al₂O₃ supported catalyst. The figure also shows the extent of coke formation on these catalysts.

In contrast to the effects of oxidative pretreatment, nitriding by reaction with NH₃ promoted catalytic activity, which increased with the nitrogen content of the supports. The surface areas (N₂, BET at 77 K) of the original and the nitrided (NH₃, 600°C) carbons are 399 and 365 m² g⁻¹, respectively. Hence the high activity of the catalyst prepared on the nitrided carbon cannot be explained by the change in surface area. It is considered that the increased activity is due to the interaction between the incorporated nitrogen and the metal species. The FTIR spectrum for the catalyst prepared on the support treated in NH₃ at 600°C indicated that there may be some chemical interaction between the support and the metal species. This postulate is supported by the changes in the FTIR spectrum of poly-4-vinylpyridine upon metal loading (Figure 2).

The increase in activity of carbon-supported catalysts due to prenitriding was accompanied by only small increases in coke formation. In both instances, the extent of coke formation was much lower than obtained on the γ-Al₂O₃ supported catalyst. Presumably this can be related to the much lower concentration of acid sites which would be present on carbon compared to alumina surfaces.

**SUMMARY**

In this work, the activities of catalysts, prepared on preoxidized and pre-nitrided Ambersorb XE-340 carbon and on a preoxidized carbon black composite (CBC) support have been investigated for asphaltene conversion.

Preoxidation of the Ambersorb supports reduced catalyst activity. In some cases
FIGURE 4 Relative activities of Mo/C catalysts prepared on pre-nitrided supports.

the decrease in activity was clearly due to loss of support surface area. Pre-oxidation of the CBC support enhanced activity. While it has not been unambiguously demonstrated, it is considered that the effect of oxygen surface functionality on catalyst activity is dependent upon the structure of the carbon itself. In some cases, oxidation may provide preferred sites for the interaction of metal species, thus promoting dispersion. For other carbons, oxidative treatment may modify or destroy existing sites would could otherwise have provided a more favorable interaction.

In contrast to the effects of preoxidation, prenitriding was found to have a distinct effect in enhancing catalyst activity. At the same time, there was no commensurate increase in coking propensity. It is hypothesized that the increased activity may be related to the presence of nitrogen-containing surface groups which provide preferential sites for the adsorption of Mo species.

The evidence from this and other studies indicates that the properties of carbon-supported catalysts can be significantly improved by the introduction of surface functional groups. However, the type and concentration of such groups will be controlled by the chemical structure of the carbon. The almost infinite variation which is possible in the structural composition of carbons may well provide the principal reason why different researchers working with their own selected carbons have produced conflicting results.

The importance of using carbons which have been well-characterized in terms of their physical and chemical structure does not appear to have been generally recognized in this field of research. Nonetheless, such information is essential to deriving any fundamental understanding of the chemistry of the catalytic processes. In view of the technical difficulties in achieving this objective, there
is cause to consider the advantages which could be derived by different research groups adopting "standard" carbon supports.

It is also important to note that, in these studies, only one precursor compound was used and all of the foregoing discussion is concerned with the interaction of various carbon surfaces with this compound. Obviously, the affinity between a particular carbon surface and the precursor will depend upon the compatibility of the two chemical structures and it is quite possible that the activities of the oxidized Ambersorb samples could be improved by employing a different precursor. Some of our recent results, yet to be published, illustrate that the HDS activity of certain carbon-supported molybdenum catalysts can be greatly increased by loading the carbon with an organometallic molybdenum compound.

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