Characterization of HDS/HDN active sites in cluster-derived and conventionally-prepared sulfide catalysts

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Abstract

The catalytic and sorptive properties of a series of sulfide cluster-derived catalysts were compared to those of an analogous series of conventionally-prepared sulfides. The cluster-derived and conventionally-prepared catalysts had similar hydrodshydro treatment activities and chemisorbed similar amounts of \( \text{O}_2 \) and \( \text{NO} \). From infrared spectroscopy, it was observed that the clusters retained much of their character upon adsorption, and that the CO ligands were lost upon heating up to 350 K. Nitric oxide chemisorbed onto the bimetallic cluster-derived catalysts was initially associated with the Co sites then shifted to the Mo sites upon heating, indicating that the promoter and Mo were in close proximity. For the conventionally-prepared materials, NO adsorption suggested that the promoted and unpromoted sites behaved independently. Nitric oxide was more strongly held to the unpromoted sites in both the cluster-derived and conventionally-prepared materials. Thiophene adsorbed onto the cluster-derived and conventionally-prepared catalysts gave rise to nearly identical infrared spectra. Upon heating, the thiophene peaks disappeared and were replaced by peaks that we have attributed to a linear olefin, which is noteworthy since 1-butene was the primary product from thiophene HDS. Nitric oxide easily displaced adsorbed thiophene from the cluster-derived materials, indicating that NO and thiophene adsorbed onto the same sites.

1. Introduction

The promotional effect of late transition metals on early transition metal sulfide hydrotreatment catalysts has been extensively investigated [1–5]. Most commercial hydrotreatment catalysts consist of Mo or W sulfides promoted with Co or Ni. A clear understanding of hydrodesulfurization (HDS) and hydrodenitrogenation (HDN) over supported hydrotreatment catalysts remains elusive because the active
sites in conventionally-prepared materials are typically minority species. Organometallic clusters appear to be useful precursors to model heterogeneous catalysts because their uniformity makes them more amenable to characterization than conventionally-prepared catalysts. Previous thiophene HDS [6] and pyridine HDN [7] activity studies in our group have shown that catalysts derived from the sulfide clusters, \((\text{MeCp})_2\text{Mo}_2(\mu-\text{SH})_2(\mu-S)_2\) \([\text{MoS}]\), \((\text{MeCp})_2\text{Mo}_2\text{Fe}_2(\mu_3-S)_2(\text{CO})_8\) \([\text{MoFeS}]\), and \((\text{MeCp})_2\text{Mo}_2\text{Co}_2(\mu_3-S)_2(\mu_4-S)_2(\text{CO})_2\) \([\text{MoCoS}]\), where \text{MeCp} = methylcyclopentadienyl, had activities and product distributions that were similar to \text{MoS}_2 and sulfided Co-Mo/\gamma-\text{Al}_2\text{O}_3 catalysts. In this paper, we describe the use of infrared spectroscopy to characterize the active sites in the cluster-derived and conventionally-prepared materials.

2. Experimental

2.1. Preparation

The \text{MoS} cluster was synthesized according to procedures described by Rakowski DuBois et al. [8] and the \text{MoCoS} and \text{MoFeS} were prepared using procedures outlined by Curtis and coworkers [9, 10]. Details concerning the dispersion of these clusters onto high surface area supports have been reported elsewhere [11]. Care was taken to minimize or eliminate air exposure of the supported clusters both during preparation and characterization.

A variety of methods have been used to prepare unpromoted and promoted supported molybdate catalysts. We compared results for the cluster-derived materials with those for catalysts made by incipient wetness and equilibrium adsorption techniques. The molybdates were supported onto \gamma-\text{Al}_2\text{O}_3 and promoted with Co, Ni, or Fe to produce loadings of 1-10 wt.-% total metal with a 1:1 atomic ratio of Mo to promoter. Ammonium heptamolybdate and Co, Ni, or Fe nitrate salts were impregnated to the point of incipient wetness either sequentially with Mo first (SIW) or simultaneously (CIW). Additional materials were prepared using the equilibrium adsorption (EA) protocol for Mo [12] followed by incipient wetness impregnation of Co, Ni, or Fe. For future reference, the code \text{M}_1\text{Mo/A} will denote a sulfided material containing promoter metal \text{M}_1 and Mo supported on \gamma-\text{Al}_2\text{O}_3. Unless otherwise specified, the total metal loadings of the catalysts were 1 wt.-% metal.

After each impregnation or adsorption step, the material was calcined in dry air at 773 K for 5 hours. The metal loadings were confirmed by neutron activation analysis. Finally, the calcined catalysts were sulfided in a flowing 2% \text{H}_2\text{S}/\text{H}_2 mixture (100 cm³/min) at 673 K for 4 hours and stored in a dessicator. Unsupported \text{MoS}_2, Crosfield 477 (2-5% CoO, 12-16% \text{MoO}_3), and Crosfield 504 (2-5% NiO, 12-16% \text{MoO}_3) catalysts were also examined. The Crosfield catalysts were sulfided in 2% \text{H}_2\text{S}/\text{H}_2 at 673 K for 5 hours.

2.2. Volumetric chemisorption

We normalized the activities by volumetric chemisorption in order to make rough comparisons between the cluster-derived and conventionally-prepared catalysts. It
has been reported that oxygen and nitric oxide chemisorb on both promoted and unpromoted sites in late transition metal promoted Mo sulfides [13–19]. Thus, the chemisorption uptake-based activities should be viewed as an average activity for the promoted Mo sulfide catalysts and a lower limit of the intrinsic activity of the promoted sites.

Oxygen uptake experiments were carried out at the dry ice/acetone temperature (195 K) using a Quantasorb sorption analyzer. The pretreatment protocol consisted of purging the catalyst bed with He, followed by reduction in H₂ at 673 K for at least 4 hours, then purging with He for 1 hour. Pulses of 9.98% O₂/He were dosed over the catalyst until the eluted peak area was constant for at least three consecutive injections. No significant differences were observed between the initial chemisorption measurement and one taken after rereduction. Nitric oxide (99.9%) chemisorption was performed on some of the catalysts using the same pretreatment protocol, and the adsorption was carried out at room temperature.

2.3. Pyridine HDN activity measurements

The catalytic activities were measured under differential conditions (conversion < 5%) at 573–673 K and atmospheric pressure using a reactant mixture containing 0.6% pyridine in H₂. Reaction rates were normalized based on the Mo loading or chemisorption uptake. The cluster-derived materials were heated in He up to 673 K at 2 K/min, cooled to 573 K, heated back up to 673 K in H₂ at 2 K/min, then reduced overnight at 673 K prior to use. The conventionally-prepared catalysts were heated in He at 2 K/min up to 673 K, sulfided for 4 hours in a 2% H₂S/H₂ mixture at 673 K, then reduced overnight at 673 K. Reactants and products were separated using a 1.8 m Hayesep D column (Alltech) and analyzed using a Hewlett-Packard 5890 gas chromatograph. Peak areas and retention times were determined using a SpectraPhysics integrator and calibrated using standard alkane and alkene gas mixtures (Scott, ± 2%).

2.4. Infrared spectroscopy

Materials for infrared spectroscopic characterization were prepared inside a nitrogen-filled glovebox. The cluster of interest was dissolved in dichloromethane and then pipetted onto wafers of γ-Al₂O₃ (10 mg/cm²), which had been calcined at 773 K, so as to produce a catalyst with a total metal loading of 1 wt.-%. The cluster-derived materials were then inserted into an in situ cell also located inside the glovebox, and spectra were collected without exposure to ambient conditions. Pretreatment of the cluster-derived catalysts consisted of heating the materials to 673 K at 2 K/min in He and holding at that temperature for 4 hours. After cooling and spectral collection, the wafers were heated in He back up to 673 K and reduced in H₂ at 673 K for 4 hours. Unless otherwise noted, each treatment was followed by an hour-long He purge at 673 K. For comparison, pressed wafers of blank γ-Al₂O₃ and alumina-supported catalysts prepared via the CIW, SIW, or EA-IW
methods with the same total metal loading and atomic ratio were also examined after a similar treatment.

Following reduction in H2 and a He purge at 673 K, nitric oxide was adsorbed onto the cluster-derived materials at room temperature (RT). Spectra were collected after purging the catalysts in He at RT for one hour so as to remove any physisorbed species. Nitric oxide desorption experiments consisted of heating the wafers at 2 K/min up to a given temperature, holding at that temperature for one hour, cooling to RT, then collecting a spectrum.

The amount of pyridine or thiophene adsorbed onto the cluster-derived ensembles and not the alumina at temperatures lower than 673 K was insufficient to be detected by infrared spectroscopy. Therefore, the adsorption was carried out at elevated temperatures. The treatments prior to the adsorption measurement consisted of heating the reduced catalyst wafers in He up to 673 K, exposing the wafers to a 0.6% pyridine or 2.7% thiophene in He mixture for 4 hours at 673 K, cooling in the flowing adsorbate to room temperature, and then purging in He for 1 hour.

3. Results and discussion

3.1. Volumetric chemisorption

Neither O2 nor NO adsorbed onto the blank γ-Al2O3. Oxygen and NO chemisorptive uptakes on the bimetallic cluster-derived catalysts (MoCoS/A and MoFeS/A) were within experimental error (±15%) of each other. The oxygen uptakes for materials which had been used for HDN were typically 5–10% lower than those for materials which had not been used, but this was also within the error of the measurements. Table 1 summarizes the chemisorption results. Notice that the uptakes on the cluster-derived materials were similar to those on the conventionally-prepared materials. In fact, with the exception of the catalysts prepared by the coimpregnation (CIW) method, the O/Mo ratio was similar for all the catalysts. The fact that the CIW-prepared, Fe- and Co-containing materials adsorbed less oxygen than the others may be a consequence of interactions of Mo and Fe or Co in the impregnating solution. When we prepared impregnating solutions for the 5

<table>
<thead>
<tr>
<th>Total metal (wt.-%)</th>
<th>Preparation</th>
<th>Moles adsorbed oxygen per mole molybdenum</th>
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<tr>
<td></td>
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<td>Mo/A</td>
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<tr>
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<td>CLUSTER</td>
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<tr>
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<td>CIW</td>
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<td>EA-IW</td>
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<tr>
<td>10</td>
<td>CIW</td>
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<td>SIW</td>
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<tr>
<td>10</td>
<td>EA-IW</td>
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</table>

*aCould not be prepared.
Table 2
Oxygen chemisorption normalized pyridine HDN activity

<table>
<thead>
<tr>
<th>Total metal (wt.-%)</th>
<th>Preparation</th>
<th>Pyridine HDN activity(^a) ((10^7 \text{ moles pyridine/mole O}_2/\text{s}))</th>
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</thead>
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<tr>
<td></td>
<td></td>
<td>Mo/A</td>
</tr>
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<tr>
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<td>CIW</td>
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<tr>
<td>10</td>
<td>EA-IW</td>
<td>37</td>
</tr>
</tbody>
</table>

\(^a\)Pyridine HDN at 633 K, oxygen chemisorption at 195 K.

\(^b\)Could not be prepared.

Fig. 1. Infrared spectra: (A) after adsorption of MoCoS cluster onto alumina; (B) after heating in He to 673 K; and (C) after reduction at 673 K.

wt.-% Fe–Mo/A catalyst, a gel formed within minutes. Bubbles were formed when Co was added to Mo-containing impregnating solutions for the high-loaded CoMo/A catalysts. No visual evidence was observed for interactions of the Mo and Ni
salts in solution. All of the 10 wt.-% promoted catalysts adsorbed less oxygen than the unpromoted catalysts. This was due to the formation of Co and Ni sulfides, which have lower chemisorptive capacities than do Mo sulfides. The amount of oxygen chemisorbed after pyridine HDN was within experimental error of that after sulfiding.

3.2. HDN activity measurements

Thiophene HDS [6] and pyridine HDN [7] activities and product distributions for the cluster-derived materials have been reported previously. In this paper, we report pyridine HDN activities for the conventionally-prepared catalysts and compared them with those of the cluster-derived materials (Table 2). The pyridine HDN activities of the cluster-derived materials were somewhat lower than those of the conventionally-prepared catalysts. The oxygen-uptake normalized activities for the conventionally-prepared materials decreased with increasing loading, although the per-gram activities were about the same. We believe that this was due to the formation of promoter metal sulfide domains that were less active than MoS$_2$ or promoted-MoS$_2$ domains. For the low-loaded catalysts, the trend with respect to the promotional effect for HDN was similar to that observed for HDS (Co–Mo > Ni–Mo > Fe–Mo > Mo). Ho et al. reported that Ni–Mo and Fe–Mo catalysts were superior to Co–Mo for HDN at high loadings [20]. Zdražil and coworkers did not observe a synergistic effect for Co–Mo HDN catalysts [21].

![Infrared spectra of NO adsorbed on catalysts](image)

Fig. 2. Infrared spectra of NO adsorbed on (A) 1% Co/A IW, (B) 1% Mo/A IW, (C) 1% Co–Mo/A SIW, and (D) 1% MoCoS/A.
3.3. Infrared characterization during pretreatment of MoCoS/A

Fig. 1A displays the spectrum of the untreated MoCoS/A catalyst. The three peaks at 2009, 1989, and 1958 cm\(^{-1}\) were due to terminal carbonyl stretch bands characteristic of the precursor cluster [11,22]. The only difference between the spectrum of the pure cluster and that of the alumina-supported cluster in this region was a slight increase in the relative intensity of the high wavenumber peak. The peaks observed between 1200 and 1800 cm\(^{-1}\) (Fig. 1B) were due to aluminum carbonates and bicarbonates, some of which remained after reduction (Fig. 1C). Since the clusters were the only source of CO, the presence of carbonates suggested that the MoCoS cluster binded to the alumina by removal of carbonyl ligands from one of the Co atoms.

In situ temperature-programmed decomposition experiments revealed that the terminal carbonyl bands were lost abruptly and completely over the range of 330–360 K. The narrow temperature range for decarbonylation suggested that the clusters produced highly uniform sites. This is consistent with our transmission electron microscopy results, which revealed a narrow size distribution for the cluster-derived scattering centers (ca. 20 Å).

Fig. 3. Infrared spectra detailing NO adsorption/desorption onto MoCoS/A after (A) reduction and (B) NO exposure. Spectra C–H were collected at room temperature after heating the catalyst in He at 323, 373, 423, 473, 573, and 673 K, respectively.
3.4. Infrared spectroscopy of adsorbed nitric oxide

The infrared spectra of NO adsorbed onto the 1 wt.-% cluster-derived MoCoS/A have been compared with those for conventionally-prepared Co–Mo/A, Mo/A, and Co/A sulfide catalysts (Fig. 2). The peak positions and relative intensities for NO adsorbed onto the conventionally-prepared catalysts were consistent with the wide body of literature on the subject [23–33]. For the unpromoted Mo/A catalyst, two peaks typical of NO adsorbed on Mo were observed (Fig. 2B). Nitric oxide adsorbed onto Co/A also gave rise to two peaks, one of which overlapped the 1805 cm⁻¹ peak for Mo (Fig. 2A). Nitric oxide adsorbed onto the three conventionally-prepared Co–Mo/A catalysts (Fig. 2C) produced three peaks which were roughly but not exactly the weighted sum of the spectra of NO adsorbed onto the Co and Mo sulfides. For the cluster-derived MoCoS/A catalyst, NO adsorption only produced bands consistent in position with NO adsorbed onto Co (Fig. 2D). We, therefore, concluded that the character of sites on MoCoS/A was predominantly, that of the ‘promoted’ sites. From this point, we will refer to sites consistent with NO on Co as ‘promoted’ sites and those with NO on Mo as ‘unpromoted’ sites.

A series of spectra detailing the consequence of heating the MoCoS/A catalyst following NO adsorption are given Fig. 3. Upon heating to 323 and 373 K (Spectra 3C and 3D), the peak at 1865 cm⁻¹ due to NO on Co decreased progressively with

![Fig. 4. No significant differences were observed in the infrared spectra of thiophene on the MoCoS/A and the conventionally-prepared Co–Mo/A catalysts. Illustrated are thiophene adsorbed on (A) MoCoS/A, (B) Co–Mo/A CIW, (C) Co–Mo/A SIW, and (D) Co–Mo/A EA-IW.](image-url)
respect to the peak at 1805 cm$^{-1}$. Concomitantly, the intensity of the peak at 1692 cm$^{-1}$, consistent with NO associated with unpromoted sites, increased. This result was reproduced several times. Therefore, we concluded that Mo and Co were in close proximity for the cluster-derived catalysts. This contradicts EXAFS data which suggested no Mo–Co interaction after the material had been heated above 400 K during the initial pretreatment [22].

The desorption behavior revealed no evidence for an interaction between the NO on promoted and unpromoted sites on the conventionally-prepared Co–Mo/A materials. However, for both the cluster-derived and conventionally-prepared catalysts, the NO desorption temperatures were 373 K and 673 K for the promoted and unpromoted sites, respectively. Rosen et al. suggested that NO adsorbs effectively as a dimer [34]. Our results were in agreement with this.

Nitric oxide could not be readorsed onto the cluster-derived materials without reduction. After reduction, however, the materials readorsed NO. This suggested that NO dissociated on heating, leaving atomic nitrogen or oxygen behind. Presumably, the nitrogen and oxygen were removed during reduction.

3.5. Infrared spectroscopy of adsorbed pyridine and thiophene

The spectra of pyridine adsorbed onto the cluster-derived catalysts and blank γ-Al$_2$O$_3$ were indistinguishable. This suggested that most, if not all, of the pyridine adsorbed onto the γ-Al$_2$O$_3$ surface; therefore, pyridine adsorption results will not be discussed further.

![Infrared spectra detailing adsorption/desorption of thiophene onto MoCoS/A following (A) reduction at 673 K, and (B) exposure to thiophene. Spectra C–G were collected at room temperature after heating the catalyst in He at 323, 373, 473, 573, and 673 K, respectively. Spectrum H is of the rereduced material.](image-url)
Fig. 6. Infrared spectra of the MoCoS/A catalyst following (A) reduction at 673 K, (B) exposure to thiophene, (C) exposure to NO after thiophene, (D) rereduction at 673 K, and (E) exposure again to thiophene.

The character of thiophene adsorbed onto the cluster-derived MoCoS/A and conventionally-prepared Co–Mo/A catalysts were strikingly similar (Fig. 4). We attributed the bands at 1576, 1471, and 1253 cm\(^{-1}\) to aromatic C=C stretches from adsorbed thiophene based on a comparison of the positions with liquid-phase thiophene (1592, 1435–1400, and 1255–1210 cm\(^{-1}\), [35]) and gaseous thiophene (1601, 1461, and 1258 cm\(^{-1}\)).

A series of spectra tracking the fate of thiophene adsorbed on the MoCoS/A catalyst is illustrated in Fig. 5. Heating to 323 K in He caused a gradual decrease in the aromatic C=C features at 1576 and 1471 cm\(^{-1}\) (Fig. 5C). The weak but sharp peak observed at ca. 1253 cm\(^{-1}\) increased in intensity. We also observed the emergence of a broad band at 1710 cm\(^{-1}\), consistent with alkene C=C bonding. The strong bands at ca. 1400 cm\(^{-1}\) were characteristic of an in-plane scissoring, terminal alkene C–H bond. Weaker bands at 1451–1461 cm\(^{-1}\) indicative of methylene groups were also observed. All these features suggested that the thiophenic surface species was replaced by a linear olefin. This olefinic group may be a precursor to 1-butene, the most common product of thiophene HDS over both the cluster-derived [6] and conventionally-prepared materials [36]. A small, as-yet unassigned peak was also observed at 1307 cm\(^{-1}\). After a one hour purge in He at 373 K, the peaks at 1307 and 1248 cm\(^{-1}\) were no longer discernable, and the peaks attributed to the linear olefin increased in intensity for all the catalysts. After the materials were heated to 473 K for one hour, the band at 1710 cm\(^{-1}\) was almost completely removed (Fig. 5E), indicating that the olefinic species had desorbed.
Further heating in He did not significantly influence the spectra (Fig. 5F and 5G). Finally, spectra taken after reduction at 673 K (Fig. 5H) were nearly identical to spectra prior to the thiophene exposure (Fig. 5A), indicating that the reduction sufficiently regenerated the catalysts.

It appeared that thiophene was weakly adsorbed on both the MoCoS/A and conventionally-prepared CoMo/A catalysts and desulfurized once adsorbed. From the infrared spectra of thiophene alone, we were unable to distinguish between the promoted and unpromoted sites. However, NO easily displaced thiophene from sites in the MoCoS/A catalyst as indicated in Fig. 6. Note that the NO spectrum following thiophene displacement was similar to the NO spectrum for the freshly reduced material. This behavior was reproducible as indicated in Figs. 6D and 6E. Based on this evidence, we concluded that NO and thiophene adsorbed onto the same sites.

4. Conclusions

The chemisorption results suggested similar site densities for the cluster-derived and conventionally-prepared catalysts. The ratio of adsorbed \( \text{O}_2 \) or NO to Mo was generally the same for the cluster-derived and conventionally-prepared catalysts. The only exceptions were several high-loaded catalysts prepared by coimpregnation, for which interactions occurred between Mo and the Fe or Co species in the impregnating solution.

The pyridine HDN activities of the cluster-derived materials were somewhat less than those of the conventionally-prepared catalysts. The trends with respect to the promotional effect for HDN for both the 1 wt.-% metal-loaded cluster-derived and conventionally-prepared materials were the same as those observed for HDS (CoMo/A > Ni-Mo/A > FeMo/A > Mo/A). For the 10 wt.-% metal loadings, however, the HDN activity decreased upon the addition of Co and Ni, which we believe to be due to the formation of bulk promoter sulfides.

Infrared spectroscopy indicated that NO adsorbed predominantly onto Co sites in the MoCoS cluster-derived catalysts. As the materials were heated, the intensity of the peak attributed to NO on Co sites decreased concomitantly with the appearance of a band consistent with NO on Mo, indicating that in these catalysts, the chemistries of the promoted and unpromoted sites were connected. One would also deduce that Mo and Co were in close proximity. In contrast, NO desorbed from the Co and Mo sites independently for the conventionally-prepared materials.

Thiophene adsorbed onto the cluster-derived and conventionally-prepared Co-Mo catalysts gave rise to nearly identical infrared spectra. Upon heating, the thiophene peaks were replaced by peaks that we have attributed to a linear olefin, which might be a precursor to 1-butene, the primary product from thiophene HDS. Adsorbed thiophene was easily displaced by NO at room temperature. The resulting NO spectrum was nearly identical to that of NO adsorbed on the freshly reduced catalyst. This suggested that thiophene and NO adsorbed onto the same site.
Acknowledgments

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References