STABILITY AND REGENERABILITY OF CATALYSTS FOR THE DESTRUCTION OF TARS FROM BIOMASS AND BLACK LIQUOR GASIFICATION

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ABSTRACT

The goal of this project was to develop catalytic materials and processes that would be effective in the destruction of tars formed during the gasification of black liquor and biomass. We report here the significant results obtained at the conclusion of this two year project. A one year no-cost extension of this project was requested and approved.

Task 1. Was completed. We identified Group VIII transition metal catalysts as active for the gasification of benzene (model tar compound) using steam. These catalysts, however, don’t have any sulfur tolerance. We have identified Group VI B sulfide materials such as tungsten sulfide or Cobalt-Molybdenum sulfides as active catalysts for the reaction, and with good tolerance towards sulfur.

Task 2. ThermoChem. Inc. and Kellogg Brown Root were identified as potential partners for the industrial site pilot reactor studies. However, no pilot scale work could be done because their gasification reactors didn’t come on stream which was a necessary step before we could go and perform the pilot scale reactor testing of the chosen catalysts. However, all the pieces and designs were completed to carry out such studies.

Task 3. Was completed. It involved characterization of aged catalysts. X-ray diffraction studies showed that Group VIII transition metal catalysts lost their activity in the presence of sulfur, due to the formation of sulfide and sulfate phases. Also complex phase transformations were observed in the presence of alkali compounds.

Task 4. Was completed. The goal was to develop effective regeneration methods. It was found that sulfur-poisoned Group VIII transition metal catalysts could not be regenerated using oxidation or steam treatments at elevated temperatures. Our efforts focused instead on the regeneration of carbon-deactivated catalysts. The efforts to regenerate Co-Mo catalysts using oxidation treatment were successful in removing carbon. The resulting material was subsequently sulfided. It was possible to achieve original catalytic activity even after three weeks of aging.

With the exception of pilot scale testing of the chosen catalysts, all other objectives for the project were completed.
BACKGROUND

Biomass and black liquor gasification, because of their higher energy efficiency and lower capital costs, have the promise to be attractive alternatives for traditional combustion processes. One of the major challenges facing the commercialization of integrated combined cycle gasification technology is the formation of tars. The formation of tars (semi-volatile and non-volatile organic matter) represents a potential loss of fuel and a contaminant that must be removed from the product gas before it is fired into a gas turbine. In black liquor gasification, tars are an unacceptable contaminant in green liquor with a potential safety hazard. Tars are formed in all gasification processes but they are a greater problem in biomass and black liquor gasification than in fossil fuel gasification. The amount and composition of the tar compounds depend greatly on the temperature and composition during the gasification process. In an earlier study (1,2), tars formed during gasification of kraft liquor consisted of mono-aromatic and poly-aromatic compounds. Benzene, toluene, and xylene were the primary tar species, but naphthalene, phenanthrene, acenaphthene, chrysene, and perylene were also observed.

Methods to eliminate tars include thermal cracking, steam reforming, and catalytic destruction. Of these, catalytic approach is the most promising alternative to destroy tar compounds since it can convert tar to gaseous fuels at moderate temperatures. Several different catalytic materials have been tested for biomass tar destruction, including dolomite, nickel and cobalt based catalysts (3-8). The rapid deactivation of catalysts in the fuel gas presents a major challenge. Catalytic destruction of tars involves breaking as well as the formation of carbon-carbon, carbon-oxygen, and carbon-hydrogen bonds. Since similar bond transformations are observed in petroleum processing (for example hydrotreating, catalytic reforming etc.), Group VIII transition metals would appear to be prime catalytic materials for tar destruction (9). Catalytic deactivation due to carbon and sulfur may be expected in both processes (10,11). However, there are two other potential sources of catalyst deactivation not encountered in the petroleum processing industry. The first is the presence of chlorine as HCl in the reaction environment (up to 1000 ppm). Secondly, high temperature operations (above 700 °C) may lead alkali metal (Na) to diffuse into the bulk of the catalytic metal, thus causing the catalytic phase to lose its structural strength.

The objectives of our study were to investigate various catalytic materials for tar destruction. The goal was to develop stable catalysts as well as to define approaches for regeneration of deactivated catalysts. We report below the results obtained from our sulfur poisoning studies as well as the potential of using sulfide phases as active catalysts for steam reforming of tars.

In order to destroy these tar compounds, catalytic conversion appears to be the most promising alternative since it can convert tars to fuel gas at moderate temperatures. A great deal of research has been conducted in search for superior catalysts and the optimal conditions for tar destruction processes. Extensive research has been performed with a variety of catalysts, both metallic and non-metallic. Mixing catalysts with the biomass (in-situ) gives unfavorable results for several reasons: coke formation because biomass is
in contact with the catalysts, breaking of catalyst particles, and agglomeration of catalysts and biomass particles. Use of a secondary reactor to destroy tar compounds is a more feasible approach. The catalyst deactivation by sulfur is a major consideration in the development of a successful catalytic process. During black liquor gasification, high levels of H₂S are expected to be present in the product gas stream along with other sulfur-containing species such as COS, CS₂, SO₂, and mercaptans. A large amount of sodium, chlorine, and potassium salts in the form of fine particles in the gas stream typically found in black liquor gasification would also affect the catalyst performance. The catalysts deactivate rapidly due to the poisoning by sulfur-, chlorine, and/or carbonaceous species in the feedstock. The amounts of alkali metals and sulfur obtained from black liquor gasification are different than those obtained from the biomass gasification, especially as it concerns the catalytic gasification of tar compounds. However, the pathways for the tar gasification as well as catalyst deactivation would be similar in these two processes. Use of guard beds can prolong the catalysts life against sulfur, but it is of little help in deactivation by carbonaceous species.

**APPROACH**

The protocols for increased catalyst life involve defining processing conditions and catalytic materials such that there are minimal transformations of the catalytically active solid-state process. In terms of processing conditions, the main variable is reducing vs. oxidizing environment at the catalyst surface, which can be altered by choosing between H₂, steam, and/or air. It should be kept in mind, however, that steam is a natural choice, since it is present in the fuel gas obtained from black liquor gasification. Use of air or oxygen to oxidize tar compounds may lead to lowered energy efficiency. The formation of stable catalytic phase might involve the use of spinels (e.g. nickel aluminate) copper chromite, nickel-chromite, or sulfided metals (e.g. CoS-MoS₂/Al₂O₃), which have demonstrated gasification activity and should be resistant to poisoning by sulfur. This research aims to focus on both classes of materials. The work to be accomplished has been divided into several tasks:

**Task 1. Bench Scale Reactor Studies**

This was the focus of our efforts during the first year. A variety of catalysts were examined for their activity and sulfur tolerance. Also, the effect of alkali compounds was examined. These studies were conducted using benzene as a model tar compound in a laboratory scale reactor operating in a fixed bed mode at atmospheric pressure.

**Task 2. Industrial Site Reactor Studies**

These studies were to be conducted using an industrial feed stream (at the site of the industry partner) with catalyst candidate(s) chosen based on the results from Task 1. We identified two industrial partners and visited one of the sites. We had designed the pilot scale reactor as well as had assembled the necessary components for the same. However, we never were able to do these runs because the gasification units didn’t come on stream.
as expected. There was no other feasible site within the US to conduct these tests during the period of this project.

**Task 3. Aged Catalyst Characterization**

Selected aged catalysts from bench scale reactor studies were characterized using x-ray diffraction (XRD) and scanning electron microscopy (SEM), and energy-dispersion analysis (EDAX). The purpose was to understand the underlying deactivation phenomena and to develop effective catalyst regeneration schemes.

**Task 4. Catalyst Regeneration Studies**

Task 4 involved regeneration of catalysts deactivated either by sulfur or carbon. The catalyst regeneration scheme(s) were developed based on the results from Task 3, and were tested in the bench scale reactor.

**EXPERIMENTAL APPARATUS & METHODS**

Figure 1 shows a simplified schematic diagram of the catalytic reactor system used for bench scale studies. The reactor was fabricated from alumina tubing (1/2” in diameter x 1 ft long). The catalyst was supported on a fritted disc located in the middle of the alumina tube. All the gas lines were heat-traced to prevent condensation of the reactant or any product species.

Figure 1. A schematic of the Reactor set-up
Briefly, a ½ inch diameter alumina tube served as the reactor tube that was placed inside a vertical tubular furnace. The feed gas system consisted of several gas streams that could be mixed in controlled flow rates to yield a desired feed composition. Benzene was introduced into the feed by sending helium through a benzene bubbler. Similarly, water (or steam) was generated by sending through a hot tube containing water at elevated temperature. All the feed lines leading to the reactor were heat-traced to prevent any condensation. Likewise, the reactor effluent lines were heat-traced for the same reasons. The reactor effluent was directly fed into an on-line FT-IR. A Bomen model MB-100 FT-IR was used for real-time analysis of feed and product composition. The Table I below shows the typical reaction conditions used in this study.

Table I. Experimental Conditions for Catalytic Testing

<table>
<thead>
<tr>
<th>Variable</th>
<th>Normal Values</th>
<th>Operating Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Catalyst loading (gm)</td>
<td>0.4</td>
<td>0.4 - 2</td>
</tr>
<tr>
<td>Sand (gm)</td>
<td>2</td>
<td>0 - 2</td>
</tr>
<tr>
<td>Temperature (°C)</td>
<td>800</td>
<td>600 - 900</td>
</tr>
<tr>
<td>Feed Flow Rate (ml/min)</td>
<td>192</td>
<td>190 - 192</td>
</tr>
<tr>
<td>Benzene concentration (%)</td>
<td>0.23</td>
<td>0.15 - 0.30</td>
</tr>
<tr>
<td>Water concentration (%)</td>
<td>1.3</td>
<td>0.6 - 2.3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Impurities when Tested</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>H2S concentration (ppm)</td>
<td>225</td>
<td>100 - 400</td>
</tr>
<tr>
<td>Alkali loading (gm)</td>
<td>0.2</td>
<td>0.05 - 1</td>
</tr>
</tbody>
</table>

A number of catalysts were tested- some prepared in the laboratory and some off-the-shelf from manufacturers. These catalysts were pretreated (the exact pretreatment will be noted when appropriate) before being loaded into the reactor. Reaction-aged and fresh catalysts were stored in glass vials for XRD and SEM/EDS analyses. XRD studies were carried out using a Phillips PW1800 diffractometer. The SEM was a LEO 1530 model and EDS was an INCA Energy (model 7426) from Oxford Instruments. All other details are given elsewhere (12).

The model tar compound (benzene) is added by bubbling helium through a benzene container placed in a constant temperature water bath. A water vapor generator was used to feed water into the reactor. The reactor was operated at temperatures between 500°C
and 800°C. The reactor effluent stream was analyzed using on-line FTIR for CO, CO₂, CH₄, and other hydrocarbons (including the reactant benzene), as well as sulfur-containing compounds.

In initial runs, the catalyst weight was 0-2 gm, with an additional 2 gm of inert material (purified sand), except where noted. The feed stream consisted of 2400 ppm benzene, while the water vapor concentration was adjusted to one and two times the stoichiometric need (1.44% and 2.88% respectively) based on assuming total conversion of tar species to CO and H₂. Balance of the feed stream was helium. In experiments where the effect of sulfur on catalyst activity was examined, 225-350 ppm H₂S was added to the feed stream; all else remained unchanged. Fine Na₂S and Na₂CO₃ particles were added to the catalyst bed in order to study the effect of alkali salt on catalyst activity. Reactor-aged catalyst samples were stored under He in glass vials for further analyses using XRD, SEM etc.

RESULTS SUMMARY

Task 1.1 Activity Measurements

Table I lists the catalysts tested in Bench Scale Reactor studies. Group VIII transition metal catalysts (Pt, Pd, Rh, Ru, Ni) supported on Al₂O₃, SiO₂, or SiO₂–Al₂O₃ were found to be active for benzene gasification using steam. Figure 2 shows the transient behavior for product gas concentrations for the Pt/Al₂O₃ catalyst at 700°C. The major reaction products are CO and CO₂, with small quantities of methane. Furthermore, the activity for CO formation decays slowly due to the build-up of carbonaceous species. Among all Group VIII transition metals, Pt/Al₂O₃ was the most active catalyst (at least on a per unit weight basis).

![Figure 2. Product Gas Concentrations from a Catalyst Screening Test with 5% Pt/Al₂O₃ At 700 ⁰C.](image-url)
In contrast, mixed metal oxide catalysts showed significantly lower activity even at temperatures as high as 800°C. The same was true of sulfided catalysts, e.g. CoS-MoS2/A12O3 etc. The only exception appeared to be tungsten sulfide (WS2) and nickel – tungsten sulfide catalysts. At higher temperatures, much less CO2 is observed, and CO is the major reaction product. This is presumably due to the water gas shift reaction.

In order to allow a comparison between different catalysts, and to normalize against amount of catalysts used in the reactor, the catalyst activity was defined as reaction rate in moles (CO + CO2) per gm catalyst per minute. In the following pages, the rate results will be compared with and without H2S.

Table I. List of Catalysts Tested in Bench Scale Reactor

<table>
<thead>
<tr>
<th>Catalyst Description</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>5% Pt/A12O3</td>
<td>Aldrich</td>
</tr>
<tr>
<td>5% Pd/A12O3</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>5% Rh/A12O3</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>5% Ru/A12O3</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>64% Ni/SiO2-Al2O3</td>
<td>Acros</td>
</tr>
<tr>
<td>24.4% Ni/SiO2</td>
<td>Laboratory Prep.</td>
</tr>
<tr>
<td>43% CuO- 39% Cr2O3/Al2O3</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>3.5% CoO+ 14% MoO3/Al2O3</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>10% MoO3/Al2O3</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>3% FeCl3/SiO2 – Al2O3</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>4% CoO – 12% MoO3/Al2O3</td>
<td>Laboratory Prep.</td>
</tr>
<tr>
<td>Sulfided CoO-MoO3/Al2O3</td>
<td>Laboratory Prep.</td>
</tr>
<tr>
<td>Sulfided NiO – MoO3/Al2O3</td>
<td>Laboratory Prep.</td>
</tr>
<tr>
<td>Tungsten Sulfide (WS2)</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>Tungsten carbide (94%), Cobolt (6%)</td>
<td>Alfa Aesar</td>
</tr>
<tr>
<td>Sulfided WO3/Al2O3</td>
<td>Laboratory Prep.</td>
</tr>
<tr>
<td>WS2 – NiS</td>
<td>Alfa Aesar</td>
</tr>
</tbody>
</table>
Task 1.2 Deactivation Rates

Sulfur Deactivation

Addition of 225-250 ppm H$_2$S to the feed stream resulted in rapid loss of catalytic activity for most catalysts. Figure 3 shows the transient activity behavior for 5% Pt/A1$_2$O$_3$ Catalyst at 600°C. For the initial 150 minutes, there was no sulfur present in the feed gas. The catalyst activity decayed slowly due to carbon deactivation and lost almost half of its initial activity due to carbon build-up. After 150 minutes, 225 ppm H$_2$S was introduced into the feed stream, the catalyst lost its activity very quickly. Calculations showed that 30-40 min. would be needed to provide just enough H$_2$S to poison all surface sites. In the case of Group VIII transition metal catalysts, the catalyst activity continued to decay in the presence of 225 ppm H$_2$S. In most cases, there was no visible activity left. Removal of sulfur from the feed stream after 320 minutes resulted in negligible recovery of the lost catalytic activity. XRD spectra of deactivated catalyst showed the formation of PtS$_2$ bulk phase (Figure 4).

Figure 3. Effect of Sulfur Poisoning on the Benzene Reforming Activity of 5% Pt/alumina Catalyst at 600 °C

Efforts to regenerate poisoned catalyst using air/O$_2$ were unsuccessful. Group VIII transition metal catalysts that had been deactivated in the presence of H$_2$S were examined by x-ray diffraction. In all cases, formation of sulfide and/or sulfate phases was observed. Figure 5 shows typical XRD spectra of an aged Nickel catalyst that was exposed to reaction conditions at 800°C in the presence of 225 ppm sulfur. Similar observations were made for nickel catalysts deactivated at other temperatures as well as for other Group VIII transition metals.
A Ni/Al₂O₃ catalyst was calcined at 1000°C for 12 hours to yield nickel aluminate (spinel phase). However, this material showed no measurable catalytic activity for steam-benzene reaction. Same was true for nickel sulfide, nickel chromite, and tungsten carbide species.

The only exception in terms of sulfur-tolerance has been tungsten sulfide (WS₂), Co-Mo sulfides and Ni-W sulfide species. The activity at 600°C and 700°C is rather low. Hence the catalytic activity of these materials has been measured at 800°C. There is a bit of caking involved with these materials, as evidenced by the difficulty in removing aged
catalysts from the alumina reactor tube. This has affected the reproducibility of rate measurements. At least 20 different experiments have been performed with tungsten sulfide. Table II shows a summary of reaction rate (in the absence and in the presence of H₂S). It is clear that the activity of 5% Pt/A₁₂O₃ goes to zero in the presence of 225-250 ppm H₂S, whereas that of WS₂ is largely unchanged in the presence of sulfur. This is the first material tested under mildly oxidizing conditions that shows sulfur tolerance. It is because of this unusual sulfur tolerance, in light of the behavior of other sulfided materials, that we have attempted to repeat the results multiple times; sulfur-tolerance has been confirmed. In contrast, cobalt-molybdenum sulfide catalysts (3.5% CoO –14% MoO₃/Alumina, Surface Area = 244 m²/gm, Alfa Aesar) showed no poisoning effect of sulfur. Figure 6 shows the results from one such run.

![Figure 6. Effect of 225 ppm H₂S on normalized benzene reforming activity of sulfided Co-Mo catalyst at 800 °C.](image)

It is hard to distinguish between the two curves shown above (Figure 6). The slow deactivation is again due to the carbon build-up. The catalytic activity of the Co-Mo catalysts is lower than that of Pt catalysts (on a per gram basis) and we needed to use higher temperatures (800 °C) to measure the activity. It is helpful to remember that poisoned Pt catalyst had little or no activity even at 800 °C. Another item to note is that the intrinsic activity of Co-Mo remains unchanged in the presence of sulfur. It should also be noted that no attempt has been made yet to increase or optimize the activity of this sulfur-tolerant catalyst. These results were verified and confirmed in many reactor runs.
Table II. Catalyst Deactivation by Sulfur

<table>
<thead>
<tr>
<th>CATALYST</th>
<th>Reaction Rate</th>
<th>(10^{-6} \text{ mole} (\text{CO}_2+\text{CO}) \text{ gm cat-min} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>5% Pt/Al₂O₃</td>
<td>185</td>
<td>225-250 ppm H₂S</td>
</tr>
<tr>
<td></td>
<td>154</td>
<td>0</td>
</tr>
<tr>
<td>WS₂(Alfa)</td>
<td>3.48</td>
<td>3.43</td>
</tr>
<tr>
<td></td>
<td>1.40</td>
<td>3.77</td>
</tr>
<tr>
<td></td>
<td>3.67</td>
<td>5.26</td>
</tr>
<tr>
<td></td>
<td>4.46</td>
<td>6.67</td>
</tr>
</tbody>
</table>
| Blanks indicate where no sulfur poisoning experiment was carried out.

1.3 Alkali Deactivation

To check the effect of alkali deactivation, Na₂CO₃ and Na₂S were added to the catalyst bed (in separate runs). In the case of Group VIII transition metals, addition of Na₂CO₃ has little or no effect on the observed reaction rate (in the absence of H₂S). With H₂S present in the feed, Na₂CO₃ appears to have negligible effect on the sulfur-tolerance (or a lack thereof) of Group VIII transition metals. There were weak indications that the addition of Na₂CO₃ might act as a scavenger for H₂S and thus delay the sulfur-poisoning effects. No significant effect of Na₂CO₃ addition was observed to warrant a more detailed investigation.

In contrast, when Na₂S was added to the catalyst bed, no activity was observed. Presumably, enough H₂S was generated during the initial start-up (heating phase) that the Group VIII transition metals were completely deactivated. Table III presents a summary of the role of alkali species in sulfur poisoning of Group VIII transition metals.

<table>
<thead>
<tr>
<th>Table III POISONING BY SULFUR COMPOUNDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Catalyst</td>
</tr>
<tr>
<td>----------</td>
</tr>
<tr>
<td>64%Ni</td>
</tr>
<tr>
<td>64%Ni</td>
</tr>
<tr>
<td>64%Ni</td>
</tr>
<tr>
<td>64%Ni</td>
</tr>
<tr>
<td>5%Pt</td>
</tr>
<tr>
<td>5%Ru</td>
</tr>
</tbody>
</table>
We also tested the effect of alkali metals (both Na₂CO₃ and Na₂S) on the catalytic activity of tungsten sulfide (WS₂). In every case, we have found a total loss of catalytic activity. This result is rather surprising and unexpected, especially since Na₂CO₃ has no effect on Group VIII transition metals. XRD studies of aged WS₂ catalyst identified the formation of inactive phases in the presence of alkali compounds. We examined the role of four different alkali salts: Na₂SO₄, Na₂S, Na₂CO₃, and NaCl. Interestingly, the effect of each salt was markedly different in terms of the transient behavior. Figure 7 shows the effect of Na₂SO₄. This run was begun with 225 ppm H₂S in the feed. The activity stabilizes after the first 100 minutes and there is very little deactivation after that. After 500 minutes of reactor operation with sulfur in the feed, H₂S was removed. Interestingly, removal of sulfur caused a loss of catalytic activity. After ~ 1000 minutes of sulfur-free operation, the catalytic activity had been reduced by half. When sulfur was added back to the feed stream, there was a slow recovery in the catalytic activity. The slow loss and recovery of catalytic activity point to solid-state or bulk-phase transformations occurring in the sulfided Co-Mo catalysts.

Figure 7. Benzene Reforming Activity of Sulfided Co-Mo + Na₂SO₄ at 800 °C

The effect of NaCl on the transient activity behavior was even more complex and slow. As Figure 8 shows, the catalytic activity continued to increase over the first 800 minutes of reactor operation. When sulfur was removed from the feed stream, there was a gradual loss of catalytic activity over the next 600 min. Figure 9 shows the XRD spectrum of unused, heated sulfided Co-Mo +NaCl catalyst. There is no trace of sodium molybdate or sodium aluminate, but there is a fair amount of molybdenum sulfide (MoS₂). Figure 10
shows the XRD spectrum of the same catalyst after being aged in the reactor. Now we see the presence of significant amounts of sodium molybdate. We generated similar information for other alkali compounds as well. It is possible to identify phases that are active for the steam reforming of benzene from the ones that are not active at all or have very low activity.

We have identified the following active phases for steam reforming of benzene:

MoS$_2$, MoO$_2$, and Na$_2$-O-S

We also find that the following phases are inactive for the reaction:

Na$_2$MoO$_4$ and NaAlO$_2$

Figure 8. Benzene Steam Reforming Activity of Sulfided Co-Mo + NaCl at 800 °C
Figure 9. XRD Spectrum of Unused, Heated Sulfided Co-Mo + NaCl Catalyst

Figure 10. XRD Spectrum of Spent Sulfided Co-Mo + NaCl Catalyst
In general, there are significant phase transformations occurring in the presence of alkali compounds. In the presence of alkali, it helps to have sulfur present in the feed to maintain the active phase. In the absence of sulfur, active phases of Co-Mo sulfides are degraded to inactive phases. Table IV presents a summary of the catalytic activity at various operating conditions.

Table IV. Steady State Activities of Sulfided Co-Mo for Various Process Conditions

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Reaction Rate $10^{-6}$ mole (CO+CO2+COS)/gm-cat. minute</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particle Size</td>
<td>No H2S</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Sulfided Co-Mo</td>
<td></td>
</tr>
<tr>
<td>PS: random</td>
<td>45.9</td>
</tr>
<tr>
<td>PS: 175-250 µm</td>
<td>29</td>
</tr>
<tr>
<td>PS: &lt; 175, random</td>
<td>31.4</td>
</tr>
<tr>
<td>PS: 100-150 µm</td>
<td>34.4</td>
</tr>
<tr>
<td>With Alkali Metals</td>
<td></td>
</tr>
<tr>
<td>0.2 gm Na₂CO₃</td>
<td>8.8</td>
</tr>
<tr>
<td>0.1 gm Na₂SO₄</td>
<td>11.3</td>
</tr>
<tr>
<td>0.1 gm NaCl</td>
<td>13.2</td>
</tr>
<tr>
<td>0.2 gm Na₂S</td>
<td>45.9</td>
</tr>
</tbody>
</table>

**CONCLUSIONS**

Group VIII Transition metal catalysts are active for the steam reforming of benzene, but they deactivate rapidly in the presence of trace sulfur quantities. Sulfided phases of Group VIB-VIII show promise as sulfur-tolerant catalysts. Slow deactivation is caused by the carbon build-up, but deactivated catalysts can be regenerated by oxidation treatment. The presence of alkali species results in the formation of various phases and structures depending on the alkali species. These slow solid state transformations cause complex transient activity behavior. The presence of 225 ppm H₂S leads to the formation of sulfide and sulfate phases of Group VIII transition metals (e.g., Pt, Ni etc.), and this results in an almost complete and irreversible loss of catalytic activity for tar reforming. However, Group VIB-VIII sulfided catalysts maintain a stable catalytic activity in the presence of H₂S and show promise as potential catalytic candidates.
ACKNOWLEDGEMENT

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REFERENCES


