Effect of pore geometry of mesoporous supports on catalytic performances in methane reforming

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Abstract—Catalysts prepared using three dimensional SBA-16 silica support (composed of micropores and cage-like mesopores) were tested in the reaction of methane dry reforming, in comparison with 2D hexagonal mesoporous SBA-15 support. The samples were evaluated by N\textsubscript{2} sorption and X-Ray diffraction (XRD) for the assessment of their textural and structural properties. The reducibility was characterized by temperature programmed reduction (TPR). The catalytic performances were evaluated in methane dry reforming and spent catalysts (after reaction) were characterized for the evaluation of sintering and coke formation by TPH/MS, XRD and HR-TEM.

Keywords—Mesoporous silica; methane dry reforming; pore geometry; sintering resistance

I. INTRODUCTION

Natural gas can be converted to liquid hydrocarbons through two major steps; the first one is its conversion into syngas, which is then transformed by Fischer-Tropsch synthesis (FTS) into liquid hydrocarbons. The first step can be accomplished by steam reforming, partial oxidation, autothermal reforming or other reactions. An alternative way for this conversion is the dry reforming of methane DRM (\text{CH}_4(g) + \text{CO}_2(g)\rightarrow 2\text{CO}(g) + 2\text{H}_2(g)) that has additional advantages compared to the previously cited reactions: consumption of the greenhouse gas CO2 and production of a ratio H\textsubscript{2}:CO close to one (good for increased selectivity in FTS).

DRM was performed over different types of catalysts especially noble metal catalysts like Pt, Ru, Rh, Pd and Ir that showed excellent activity and stability during reaction but these catalysts are expensive and rarely found. Nickel was as active as noble metals and thus became a promising alternative, especially that it is cheap and readily available [1-4]. However, Ni particles could still suffer from sintering or coke formation at high temperature, which drastically affects the catalytic performances. Consequently, researchers have intensified their efforts to solve these problems through several procedures such as incorporation of small amount of rare and alkaline earth metals over nickel based catalysts [5-7] or changing supports type and enhancing metal-support interaction [8,9].

Recent studies have shown that mesoporous SBA-15 silica support is promising for the confinement and stabilization of nickel in DRM reaction [10,11]. The addition of ceria to Ni/SBA-16 catalysts was studied by Zhang et al. [12] and the results showed that NiCe/SBA-16 was more stable, exhibiting higher conversion than Ni/SBA-16 in DRM. The studies in the literature were conducted under different experimental conditions (varying catalyst preparation and catalytic test conditions) and a direct comparison between SBA-15 and SBA-16 is not reported. Nevertheless, it is important to compare the two supports in order to check if the geometry of such materials can affect the performances of the resulting catalysts in the challenging DRM reaction.

II. EXPERIMENTAL

A. Catalysts preparation

Evaporation induced self-assembly [13,14] and sol-gel [15,16] methods are commonly used for the preparation of mesoporous materials. In this work, SBA-16(1) support was prepared using sol-gel technique following an established procedure of Cheng et al. [17]. In details, 5.66 g of the tri-block copolymer F-127 (PEO\textsubscript{106}PPO\textsubscript{70}PEO\textsubscript{106}) (Sigma-Aldrich, P2443) were added to 250 ml of a 2M HCl. The mixture was stirred for 2 hours at 30°C. Then 19.81 g of TEOS (Sigma-Aldrich, 86578) were used as the silica precursor and the resultant mixture was stirred for 20 hours before filtration, washing with distilled water and drying in air at 110°C overnight. The support was next calcined at 500°C for 9 hours (at 2°C.min\textsuperscript{-1}).

SBA-16(2) support was prepared as reported by Dos Santos et al. [18]. 4 g of F-127 were added to 0.4M HCl. After complete dissolution, co-surfactant butanol (Sigma-Aldrich, 33065) was added and stirred at 45°C for 1 hour. This was followed by the addition of 19.3 g of TEOS and vigorous stirring at 45°C for another 24 hours. The obtained white precipitate was then filtered, dried in air at 100°C for 24 hours and then calcined at 550°C for 2 hours (at 2°C.min\textsuperscript{-1}).

Commercial SBA-15 mesostructured silica was used as the third support in this study (Sigma Aldrich, 777242).
Impregnation of nickel (5 wt%) on the three different supports was performed following “two solvents” loading method described elsewhere [19] and the calcination based on the optimized conditions reported in previous work [11]. The resulting samples were labeled as Ni/SBA-16(1), Ni/SBA-16(2), and Ni/SBA-15.

### B. Catalysts characterization

The ASAP 2020 apparatus from Micromeritics was used to obtain the N2-sorption isotherms at -196°C after proper degas under vacuum at 250°C for 2 hours.

X-Ray diffraction peaks and results were performed on a PANalytical X’Pert³ diffractometer using Cu Kα radiation (λ=1.5405 nm) at 20 between 35 and 85°.

Temperature programmed reduction (TPR) was tested using an AutoChem II, 2920 Micromeritics apparatus.

Temperature programmed hydrogenation (TPH) coupled to mass spectrometry measurements were done on spent catalysts while heating from 100 to around 900°C (at 10°C.min⁻¹) at atmospheric pressure under 5 vol% H2/Ar (30 ml.min⁻¹). Experiments were performed by connecting the AutoChem 2920 to an online quadrupole mass spectrometer (MKS, Cirrus 2). This technique is helpful in determining the type of deposited carbon in the spent catalyst according to its gasification temperature to CH4: Cα <500°C and Cβ above 500°C [20].

A JEOl JEM-200 electron microscope operating at 200 keV (LaB6 gun) was used for Transmission electron microscopy (TEM) images.

### C. Calcined testing

Calcined catalysts were tested in a Hastelloy-X reactor (Microreactivity, PID Eng and Tech, internal diameter 9 mm). 200 mg of each catalyst were first reduced in-situ using 5 vol% H2/Ar (30 ml.min⁻¹) at 650°C for 2 hours to ensure complete reduction of nickel oxide into Ni⁰ metallic nickel. After reduction, the temperature was lowered to 200°C and then increased from 200°C to about 800°C (at 5°C.min⁻¹) again under an equimolar mixture of CH4 and CO2 using a gas hourly space velocity (GHSV) of 18 L.hr⁻¹.g⁻¹. The catalytic stability was measured by keeping the reactor at 650°C for 12 hours. Effluent gases from the reactor were analyzed online using an Inficon micro-GC equipped with a TCD and two types of columns (molecular sieves and plot-U) to detect H2, CH4, CO and CO2 gases. The reactants conversions were calculated using the following equations:

\[ \text{CH}_4 \text{ conversion (\%) } = 100 \times \frac{[\text{CH}_4(\text{in}) - \text{CH}_4(\text{out})]}{\text{CH}_4(\text{in})} \]

\[ \text{CO}_2 \text{ conversion (\%) } = 100 \times \frac{[\text{CO}_2(\text{in}) - \text{CO}_2(\text{out})]}{\text{CO}_2(\text{in})} \]

### III. RESULTS AND DISCUSSION

#### A. Structure and porosity of catalysts

N₂-sorption isotherms show a type IV isotherm for all catalysts, typical of mesoporous silica materials (data not shown). Ni/SBA-15 catalyst has H1-type hysteresis loop typical of 2-D hexagonal structure whereas both Ni/SBA-16(1) and Ni/SBA-16(2) show an H₂ hysteresis loop reflecting a cubic structure of the material with cage-like pores typical for SBA-16 parent [21].

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>BET surface area from N₂ sorption isotherms (m².g⁻¹)</th>
<th>Crystallite size by XRD (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fresh catalyst (NiO)</td>
<td>Spent catalyst (NP)</td>
</tr>
<tr>
<td>SBA-16(1)</td>
<td>408</td>
<td>7.7</td>
</tr>
<tr>
<td>Ni/SBA-16(1)</td>
<td>295</td>
<td>336</td>
</tr>
<tr>
<td>SBA-16(2)</td>
<td>138</td>
<td>7.5</td>
</tr>
<tr>
<td>Ni/SBA-15</td>
<td>701</td>
<td>-</td>
</tr>
<tr>
<td>SBA-15</td>
<td>528</td>
<td>3.3</td>
</tr>
</tbody>
</table>

BET surface areas of the three different supports with their impregnated active phase (5wt %) are grouped in Table I. Concerning SBA-16 supports, their surface area was in the range of 300-410 m².g⁻¹ whereas SBA-15 support had much higher surface area (701 m².g⁻¹). For all SBA based samples, the surface area was decreased after nickel impregnation which is attributed to partially filled pores or to some collapse of the silica walls. But since the shape of isotherms for the impregnated SBA stayed similar to the parent SBA, a well-organized framework was preserved and it can be concluded that there was no destruction of support. Such destruction can take place under the preparation procedure used in this study, but the TEM images (Fig. 4(a)) prove that SBA-16 framework and pore structures remained intact, similar to the study done on DRM in [12].

XRD for calcined samples at wide angle (35 < 2θ < 85°) are represented in Fig.1A. Fresh Ni/SBA-15/16 catalysts exhibit three main peaks at 2θ = 37, 43 and 63° (identified by ●) corresponding to NiO particles with face centered cubic (FCC) unit cell and reticular planes respectively indexed (111), (200), (220) (JCPDS 71-4750). Based on the Scherrer equation, the mean NiO particle sizes of each catalyst are found to be 7.7, 7.5 and 3.3 nm for Ni/SBA-16(1), Ni/SBA-16(2) and Ni/SBA-15, respectively (Table I). Since all SBA based catalysts are impregnated with the same amount of nickel (5 wt%), the difference in crystallite size means that nickel particles are better dispersed over SBA-15 support than on SBA-16.

#### B. Reducibility of Ni/SBA-16(1), Ni/SBA-16(2), and Ni/SBA-15

TPR profiles of reduced Ni/SBA catalysts (Fig. 2) show two hydrogen consumption peaks: a first peak between 300 and 450°C corresponding to free NiO particles easily reduced and a second peak attributable to Ni oxide species in strong interaction with the support and reduced at higher temperature, greater than 450°C [23]. With Ni/SBA-15, the main NiO species appear to be in enhanced interaction with the support. Whereas for Ni/SBA-16(1) and Ni/SBA-16(2), most of NiO where reduced at the lower temperature range.
This behavior could be due to the different availabilities of SiO$_2$ surface sites especially that Ni/SBA-15 has larger surface area than both Ni/SBA-16 catalysts and thus the smaller surfaces accessible on SBA-16 might limit the formation of mixed phases upon the addition of the nickel precursor [24].

C. Activity and stability in DRM

Catalytic stability experiments (Fig. 3) were carried out at 650°C for 12 hours under stream. Ni/SBA-15 catalyst appears to be fully stable with preservation of CH$_4$ and CO$_2$ conversions (75 and 80% respectively). On the other hand, conversions on both Ni/SBA-16(1) and Ni/SBA-16(2) start at lower values than Ni/SBA-15 and are deactivation appears through time with an average conversion loss of about 20% and 30% respectively at the end of reaction. This reveals that SBA-15 structure could prevent deactivation and maintain catalyst stability, not to mention that CO$_2$ conversions remain higher than those of CH$_4$ due to the occurrence of RWGS. Reasons behind such behaviors resulting in deactivation will be discussed in the following section.

D. Physicochemical properties of the spent catalysts

To understand the reasons behind Ni/SBA-16 catalysts deactivation, XRD of spent catalysts, TEM and TPH/MS experiments were performed to check the presence of coke and/or sintering.

XRD for the 3 spent catalysts (Fig. 1(b)) show three peaks at 20 = 44, 52 and 76° for planes (111), (200) and (220) respectively, corresponding to Ni metallic particles (identified by ♦, JCPDS 01-1258). This means that no re-oxidation of Ni$_0$ due to water formation through RWGS occurred during reaction [25,26] and therefore deactivation of Ni/SBA-16 is independent of this phenomenon. However, the peaks are much wider and almost invisible for Ni/SBA-15 catalyst than those for both Ni/SBA-16 catalysts that appear very intense. Table I shows that Ni particles over SBA-15 support remained small (3.6 nm) stressing on their high dispersion that is well retained even after test. Ni crystallite sizes for Ni/SBA-16(1) and Ni/SBA-16(2) are 11 and 13.2 nm respectively which highlights the occurrence of sintering, thus destabilizing and deactivating the catalysts as seen during catalytic test.

Fig. 4(b) of spent Ni/SBA-16(1) displays extremely large Ni nanoparticles with some carbon deposition. The weak interaction of the Ni species with SBA-16 could have facilitated their migration outside the mesopores and their aggregation at the surface at high temperature (compared to
fresh ones, data not shown), thus reducing stability and catalytic performance [27]. However, on Ni/SBA-15 (Fig. 4(c)), Ni nanoparticles are still confined inside the pores with no migration to the external surface even after 12 hours under stream. In addition to this, Ni/SBA-15 shows high level of Ni dispersion that is restrained also after test which is in agreement with the small size of Ni particles (3.6 nm) obtained by XRD. Based on these findings and on TPR experiments, Ni species strongly interacting with the support can influence the catalytic stability of Ni/SBA-15. Therefore these catalysts were more stable and showed high resistance towards particles sintering, regardless of the coke formation that will be further elaborated in TPH experiments.

In addition to metal sintering effect, coke formation might also have certain impact on catalysts behavior. TPH/MS experiments applied on the different spent catalysts determine the types of carbon deposits (Fig. 5). These experiments rely on the temperature at which methane (m/z = 16) is formed by hydrogenation of carbon species H₂ consumption.

From the three types of carbon discussed earlier, Cα is a reactive intermediate that does not have influence on catalyst stability; Cβ represents nanotubes and Cγ, a graphitic carbon, which is the most detrimental type responsible of catalyst deactivation [20,28,29]. Fig. 5 shows that Ni/SBA-15 catalyst (Fig. 5 (a)) contains the highest amount of carbon among others, but it is the only catalyst carrying the least amount of Cγ and the highest amount of Cα carbon. Cα is expected to be produced from thermal decomposition of methane and can lead after to the formation of CO by hydrogenation at low temperature [30]. Therefore, even if present at high concentrations, its effect is benign on catalyst performance and activity and this is emphasized by the high stability of Ni/SBA-15 during time on stream. Such behavior is also elaborated in a study done by Igor Luissetto et al. [31] where they show that Ni/CoO₂-Al₂O₃ prepared by co-precipitation method was highly active and stable in spite of the greater carbon deposition on its surface and related this behavior to the fact that Cα does not encapsulate Ni⁰ sites which remain accessible to the reactants.

Thus, the main cause behind catalysts deactivation of Ni/SBA-16 is attributed to sintering, weak of metal-support interaction and low nickel dispersion as shown in [32]. It is important to mention that even if carbon formation does not alter catalyst performance, its mitigation and removal is a must because it can accumulate with time leading to reactor plugging and reaction perturbation.

IV. CONCLUSION

In this study, 5 wt% Ni were impregnated on commercial SBA-15 and synthesized SBA-16 supports and the properties of the resulting samples in addition to their catalytic performances were compared. While the SBA-15 2D hexagonal structure permits the stabilization of nickel inside the pores, SBA-16 cubic structure was not ideal for dispersion of nickel inside the pores nor for their stabilization under
methane dry reforming. Ni/SBA-15 (without any promoter) are promising for DRM in opposition to Ni/SBA-16 catalysts where sintering is leading to rapid deactivation of the catalysts under stream.

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REFERENCES


