Influence of synthesis parameters of mesocellular silica foams doped by nickel on methane reforming by CO$_2$

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Abstract— New catalysts based on Ni(0) dispersed onto mesocellular silica foams (MCF) were prepared for Dry Reforming of Methane (DRM). Different synthesis methods of MCF supports (with and without using NH$_4$F or n-butanol) were tested in order to study the influence of the textural properties of the support on the catalytic performances of the catalysts. In all cases, nickel was incorporated using the double solvents method. The resulting calcined materials were characterized by N$_2$ sorption, X-Ray diffraction (XRD) and transmission electron microscopy. Their reducibility was tested by temperature-programmed reduction (TPR). Interestingly, particularly large specific surfaces, pores diameters and pores volumes were observed when NH$_4$F was added to the synthesis gel (MCF(N) materials). The corresponding Ni-MCF(N) catalysts were shown to be the most attractive among other prepared samples with respect to their performances in DRM.

Keywords — Methane reforming ; Nickel ; Nanoparticles ; Mesocellular foams.

I. INTRODUCTION

In 1999, siliceous mesocellular foams (MCF) were prepared for the first time by combining self-assembly processes through a (S$^2$H$^+$)(X$^-$) pathway used for SBA-15 synthesis and pore swelling with organic molecules [1]. Preferentially solubilized in the core of micelles, these molecules lead, as expected, to an increase of the pore volume but also to a modification of the porosity organization [2]. Since, resulting MCF with large spherical cell sizes up to 40 nm, 3D interconnected pore system and large pore volumes [3] have shown great potential as supports for drug molecule release [1] or for the design of heterogeneous catalysts through the incorporation of enzymes/proteins [4], metal nanoparticles [5] to name only a few. Up to now, various synthesis methods of MCF have been reported in the literature. Among them, the challenge is to obtain complex mesophases using a four-component microemulsion (surfactant/co surfactant/oil/water) as a template [6]. Several examples of co-surfactant / oil combinations have been described in the literature. Among the molecules added, n-butanol (n-BuOH) is known to act both as a co-solvent and co-surfactant in block copolymer-water systems. In other terms, n-BuOH stabilizes the hydrophilic-hydrophobic interfaces acting on the micellar surface curvature and on the total volume of the hydrophobic chains of the block copolymer [7].

In the present contribution, MCF doped by 5 wt.% of nickel was used for the first time as a catalyst for Dry Reforming of Methane (DRM). Studies on DRM present both scientific and industrial interest. This reaction consists in transforming two abundantly available greenhouse gases (CH$_4$ and CO$_2$) to produce industrially important syngas (a mixture of CO and H$_2$). The latter is usable in the synthesis of a broad range of higher alkanes through Fischer–Tropsch synthesis [8]. However, DRM is accompanied by secondary reactions such as Methane Decomposition (MD), Reverse Water Gas Shift (RWGS) and Boudouard reaction (BD) [9]. To limit them, to favor the endothermic DRM reaction and to reach high equilibrium conversion of CH$_4$ and CO$_2$ to H$_2$ and CO, operating temperatures of 800-1000°C are usually required. This also minimizes side reactions [8]. For these reasons, inexpensive catalysts affording the expected CO/H$_2$ molar ratio are highly desired in order to lower the operating conditions.

In this work, four types of MCF materials with different textural properties were synthesized with the aim to select the most convenient supports for DRM catalysts preparation. The latter were obtained after deposition of 5 wt.% Ni using the double solvents method [10]. The porosity and the structure of the solids as well as the nickel dispersion were analyzed. The catalysts were tested in dry reforming of methane after reduction. Their catalytic performances in terms of conversions, products selectivity and stability were analyzed and compared to thermodynamic curves simulated by the HSC program considering or not carbon formation and similar reaction conditions.
II. EXPERIMENTAL PART

A. Catalysts preparation

A typical synthesis of MCF, as described by Widyaningrum et al [11], was used first. It consisted in fully dissolving 4.0 g of Pluronic (P123) triblock copolymer (EO32PO70EO32) in a mixture of HCl (20 mL, 37%) and distilled water (120 mL) at 40°C, then adding dropwise 4.0 g of trimethylbenzene (TMB) as swelling agent, under vigorous stirring. To form a microemulsion, the suspension was stirred for 2 h then tetraethylorthosilicate (TEOS, 9 mL) was added dropwise and stirring was continued for 20 h at the same temperature. The solution was next transferred in an autoclave and submitted to a hydrothermal treatment at 100°C for 24 h. The resulting suspension was finally filtered and the solid was washed, dried and calcined under air flow at 550°C for 5 h (heating rate 1°C.min⁻¹). The obtained solid was named MCF.

Similar syntheses conditions were used to prepare sample MCF(N), except for the addition of 48 mg of NH₄F done just before the hydrothermal treatment. Also, two other MCF silica foam materials, i.e. MCF(B1) and MCF(B2), were prepared using n-butanol as a co-surfactant following the synthesis procedure proposed by X. Zhou et al [7]. The considered molar n-BuOH / TMB ratio was equal to 2, which corresponded to the addition of 2 g of TMB and 2.4 g of butanol for MCF(B1), and of 4 and 4.6 g, respectively, in the case of MCF(B2). Both, surfactant and co-surfactant, were added together.

Ni-MCF, Ni-MCF(N) and Ni-MCF(B1) materials were obtained by the impregnation of the supports with a Ni²⁺ containing solution using the double solvents method. The used nickel precursor was Ni(NO₃)₂.6H₂O and the theoretical loading of Ni added was 5 wt%. In details, the support (1.0 g) was suspended in 20 mL of n-hexane. After that, the appropriate amount of nickel precursor dissolved in a volume of water set equal to the silica pore volume (as determined by N₂ physisorption) was added dropwise. The impregnated solids were dried 3 days at room temperature then overnight at 60 °C then calcined at 550 °C under air for 2 h.

B. Catalysts characterisations

The catalysts were characterized in terms of textural, structural, reducibility and morphological properties. By recording N₂ adsorption-desorption isotherms on an ASAP 2020 (Micromeritics) apparatus the textural properties of solids were obtained. The samples were degassed, prior to experiments, under vacuum for 2 h at 250°C. The specific surface area values were obtained by using the BET equation. The pore diameters and specific pore volumes were calculated according to the BJH model. X-Ray Diffraction (XRD) data were recorded on a diffractometer XPert3 Powder (PANalytical, Netherlands) using CuKα radiation. The average particle sizes were calculated from the Scherer equation: D=Kλ/βcosθ, where K is a constant (K=0.9), λ = 1.5405 Å, β is the full width at half maximum (FWHM) of the diffraction peaks and θ is the peak position. Temperature-Programmed Reduction (TPR) done in a TPD/R/O instrument (Autochem 2920 Micromeritics apparatus) was used to inform on the nickel phase reducibility and on the interaction of the metal with the support. The catalysts (50 mg) were placed on a quartz wool in a U-shaped quartz sample tube and heated from room temperature to 800°C at a rate of 10°C.min⁻¹ under a gaseous mixture flow (25 mL.min⁻¹) of 5% H₂ diluted in Ar. Before reaching the thermal conductivity detector (TCD), the gas flow was passed through an ice bath to remove any water product formed during reduction. Transmission Electron Microscopy (TEM) analyses were performed on a JEOL-2010 (LaB) microscope operating at 200 kV. Ethanol was used to suspend the powdered samples. The resulting mixtures were deposited on a copper grid coated with a porous carbon film, then ethanol was left to evaporate by drying in room atmosphere.

C. Catalytic measurements

The catalysts were tested in a PID ENG & TECH Microactivity Effi Reactor. They were loaded into the reactor and reduced in situ at 650°C for 2 h (rate 10°C.min⁻¹) in a 5% H₂/Ar flow (30 mL.min⁻¹) to form reduced nickel known to be the active phase in DRM. After this activation treatment, the temperature was cooled down to 200°C and dry reforming of methane was conducted under atmospheric pressure with CH₄/CO₂= 1:1 (diluted with Ar) and a total GHSV=36 L.g⁻¹.h⁻¹ (100 mg of catalyst were used for each test). Good reproducibility of catalytic results under these conditions was obtained. For activity measurements, the reaction temperature was increased from 200°C (5°C.min⁻¹) up to 750°C followed by a decrease of temperature till 650°C which was maintained for 4 h. The gaseous products were analyzed online by a micro GC Agilent 490 mGC. The conversions of methane (1) and of carbon dioxide (2), as well as the H₂/CO ratios (3) were calculated as follows:

\[
\text{CH}_4 \text{ conversion} \% = \frac{\text{CH}_4(\text{in}) - \text{CH}_4(\text{out})}{\text{CH}_4(\text{in})} \times 100 \quad (1)
\]

\[
\text{CO}_2 \text{ conversion} \% = \frac{\text{CO}_2(\text{in}) - \text{CO}_2(\text{out})}{\text{CO}_2(\text{in})} \times 100 \quad (2)
\]

\[
\frac{\text{H}_2}{\text{CO}} = \frac{\text{H}_2(\text{out})}{\text{CO}(\text{out})} \quad (3)
\]

where CH₄(in) and CH₄(out) (resp. CO₂(in) and CO₂(out)) are the initial and measured CH₄ (resp. CO₂) concentrations, respectively, whereas H₂(out) and CO(out) are the measured products concentrations.

III. RESULTS AND DISCUSSIONS

A. N₂ physisorption

MCF, MCF(N), MCF(B1) and MCF(B2) supports present type IV isotherms (Fig. 1). The observed hysteresis loops (IUPAC) are typical of mesoporous materials.
Most of them are of type H2 or H1, which is in agreement with the expected presence, in mesocellular foams, of connected cells and of pores with different shapes.

Addition of ammonium fluoride or n-butanol during synthesis led to a modification of the pores volumes and of the dimensions of the cells/pores, but not as much as for the specific surface area values (Table I). On one hand, the influence of NH4F has been already described as being related to the mineralizing effect of fluoride ions, giving improved structural organization and higher pore volume [3]. On the other hand, n-butanol used as a cosurfactant in MCF(B1) and MCF(B2) led to quite low volumes values which turned to be even smaller than those obtained with SBA-15.

**TABLE I. TEXTURAL PROPERTIES OF SOLIDS BEFORE AND AFTER NI IMPREGNATION**

<table>
<thead>
<tr>
<th>Samples</th>
<th>Textural properties</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>S_{BET} (m²/g)</td>
</tr>
<tr>
<td>MCF</td>
<td>722</td>
</tr>
<tr>
<td>MCF(N)</td>
<td>755</td>
</tr>
<tr>
<td>MCF(B1)</td>
<td>696</td>
</tr>
<tr>
<td>MCF(B2)</td>
<td>687</td>
</tr>
<tr>
<td>Ni-MCF</td>
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<tr>
<td>Ni-MCF(N)</td>
<td>490</td>
</tr>
<tr>
<td>Ni-MCF(B1)</td>
<td>529</td>
</tr>
</tbody>
</table>

Due to the similar textural features of MCF(B1) and MCF(B2), further work in this paper will be focused only on MCF(B1) amongst these two samples. The shapes of the N₂ adsorption and desorption isotherms of the materials after impregnation of these supports with 5 wt.% Ni were similar to those before Ni addition, with type IV isotherms and H2 or H1 hysteresis loop (IUPAC) (Fig. 2). Nevertheless, the pores volumes and specific surface area decreased by 20 to 30% upon Ni insertion, the effect being the most pronounced in the case of Ni-MCF(N) (Table I). The diameter of the cells was retained in the case of Ni-MCF(N) and Ni-MCF(B1) samples but decreased a little bit, from 21 to 18 nm, for Ni-MCF. On the other hand, windows dimensions were reduced only for MCF(N). However, further characterization is needed in order to correlate the decrease of the dimensions of the cells or windows with the preferential location of the NiO particles.

**B. X-Ray Diffraction (XRD)**

Wide angle XRD patterns of the calcined Ni-supports are shown in Fig. 3. In all cases, a broad peak at about $2\theta = 22^\circ$ is observed corresponding to diffusion phenomena in amorphous silica.
Fig. 3. XRD patterns of (a): Ni-MCF(N), (b): Ni-MCF, (c): Ni-MCF(B1) samples.

Five diffractograms peaks at $2\theta = 37$, 43, 62, 76 and 80° are also systematically seen and attributable to (111), (200), (220), (311) and (222) lattice planes of the face centered cubic structure of NiO oxide nanoparticles (JCPDS: 01-089-7130). The Scherrer formula was used to estimate the mean size of the NiO crystallites, assuming a spherical shape. The results are reported in Table II. With a 8.7 nm mean diameter, Ni-MCF(N) is characterized by the smaller crystallites.

This size is inferior to the critical maximum value (9 nm) reported by Kawi et al [9] as providing good performances in DRM reaction. Hence, the nanoparticles in Ni-MCF(N) could be used advantageously in the studied reaction. Conversely, it is commonly admitted that Ni metal particles having bigger size than the critical one are known to favor carbon deposition and hence enhance catalyst deactivation [9].

C. Morphological properties

Fig. 4 shows typical TEM images of Ni-MCF(N) which was characterized by the best textural properties both before and after Ni addition. The images are consistent with an homogeneous organization of the support and reveal rather homogenous dispersion of the NiO nanoparticles. With low magnification, rather big Ni deposits seem to be present (Fig. 4A) but they consist in aggregates of small Ni nanoparticles with mean diameters below 10 nm (Fig. 4B, B'). Very small isolated nanoparticles are also seen everywhere in the material, as is exemplified in Fig. 4C (wight circles) and in the associated zoomed image (Fig. 4C') which also highlights the pores. The particle sizes estimated from TEM are thus in good agreement with those deduced from XRD data (Table II).

| TABLE II. NiO CRYSTALLITES SIZE DETERMINED FROM XRD DATA ANALYSIS |
|----------------------|-----------------|-----------------|-----------------|
| Samples              | Ni-MCF          | Ni-MCF(N)       | Ni-MCF(B1)      |
| Mean NiO particle sizes (nm) | 11.2            | 8.7             | 11.7            |

D. Reducibility

All calcined Ni-supports were reduced using $\text{H}_2$ under 650°C to convert NiO to Ni$^0$. TPR measurements were conducted in order to study the reducibility of deposited nickel in the calcined nickel mesocellular foams materials and indirectly its interaction with the support. Fig. 5 shows only one reduction peak per sample, centered at 405°C and 410°C for respectively Ni-MCF and Ni-MCF(B1), instead of ~ 470°C for Ni-MCF(N). Hence, both former samples behave similarly, whereas the occurrence of an $\text{H}_2$ consumption at higher temperature would suggest a stronger interaction between NiO and the support in the case of Ni-MCF(N). Nevertheless, peak surfaces are comparable for the three samples, in line with the similar Ni contents in all catalysts (5 wt %). Note also that the reduction is achieved for all samples before 600°C, confirming that the temperature of 650°C applied for in situ activation of the catalysts prior catalytic tests is appropriate for full metal reduction.

Fig. 4. TEM image of Ni-MCF(N) showing (A) a full grain and zones with (B) Ni aggregates or (C) isolated Ni nanoparticles. B' and C' are zoomed images of selected zones in B and C, respectively.
E. Catalytic performance in DRM

The Ni-MCF, Ni-MCF(N) and Ni-MCF(B1) samples were tested in DRM from 200 to 800°C. At low temperature, all catalysts are inactive, then the activity rises while increasing the temperature, in line with the endothermic nature of the reaction. Conversion of CH₄ and CO₂ vs. temperature are shown in Fig. 6, revealing significant differences in terms of catalytic performances between Ni-MCF, on one hand, and Ni-MCF(N) and Ni-MCF(B1), on the other hand.

By comparing the activity of the solids at 650°C, Ni-MCF showed the lower performances (42% CH₄ and 40% CO₂ conv.), while Ni-MCF(B1) (67% CH₄ and 75% CO₂ conv.) and Ni-MCF(N) (67% CH₄ and 68% CO₂ conv.) showed very good results. For the best catalysts, the conversions are very close to the expected thermodynamic values (maximum possible conversions) estimated with carbon formation. Not only the reactants conversions but also the selectivity of the catalysts in given products, especially their ability to give the desired H₂/CO ratio, is a very important criterion. Targeting the production of liquid fuels [12] for Fisher-Tropsch operations, a H₂/CO=1 is highly desirable. The values effectively obtained with the three selected materials are presented in Fig. 7.

Hence, the less active Ni-MCF catalyst never gave the desired ratio, even at high temperature (770°C). In contrast, H₂/CO values very close to 1 were obtained with Ni-MCF(B1) at 650°C (0.98), 750°C (1.03) and 770°C (1.04) as well as with Ni-MCF(N) at 750°C (1) and 770°C (1.01). As a whole, the Ni-MCF(N) sample turned out to gather a lot of advantages such as the best textural properties, the smallest Ni crystallites size, the highest reduction temperature and good activity and selectivity.

For these reasons, this material was selected for a study of the stability at 650°C during 4 h in flowing reactant gases as above (Fig. 8). shows that Ni-MCF(N) is characterized by a really good stability vs. time at 650°C, working very closely to the thermodynamic conditions (with carbon formation), i.e. with conversion rates of CH₄ and CO₂ equal to 60 and 71%, respectively.

![Fig. 5. Temperature programed reduction of: (a): Ni-MCF(N), (b): Ni-MCF, (c): Ni-MCF(B1) samples.](image)

![Fig. 6. Activity of reduced (•): Ni-MCF, (□): Ni-MCF(N) and (△): Ni-MCF(B1) in DRM. Thermodynamic curves with (—) and without (—-) carbon formation are also mentioned.](image)

![Fig. 7. H₂/CO ratio vs temperature for Ni-MCF, Ni-MCF(N) and Ni-MCF(B1) samples.](image)
IV. CONCLUSION

Silica mesocellular foams (MCF) with different textural properties were synthesized and used as nickel supports for preparing dry methane reforming catalysts. Unlike what was expected, silica samples synthesized using butanol as a co-surfactant (MCF(B1) and MCF(B2) materials) did not display the optimal textural properties. The largest windows apertures, pores volumes and surface area were obtained with the MCF(N) support which synthesis involved NH$_4$F. After nickel addition using the double-solvents method (post-synthesis) then activation, Ni-MCF(N) also led to the best nickel support-interaction and best nickel dispersion. It also led to the best catalytic performances in terms of CO$_2$ and CH$_4$ conversions, close to maximum thermodynamic limitations and slightly above those obtained on the samples with MCF support synthesized in presence of n-butanol. Also, Ni-MCF(N) was slightly more selective towards the production of syngas with a H$_2$/CO ratio of 1 convenient for Fischer Tropsch. Compared to bibliography, those catalyst showed satisfying activity even slightly below that previously obtained on a Ni-SiO$_2$ catalyst with the same Ni content [12]. Characterization data obtained on the spent catalyst agreed with very little carbon formation during the tests and the absence of metal active phase sintering.

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