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Supported catalysts for induction-heated steam reforming of methane

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A B S T R A C T

Ni₆₀Co₄₀ nanoparticles supported on γ -Al₂O₃ capable of simultaneously catalysing the steam reforming reaction of methane and supplying *in-situ* the heat necessary to activate the reaction by induction heating, have been synthesized and characterized. Energy is remotely and promptly supplied by an alternating radiofrequency magnetic field (induction heating system) to supported nanoparticles that act as dissipating agents by virtue of their ferromagnetic properties. The temperature reached on the Ni–Co based catalyst surface is high enough to obtain good catalytic performances for the steam methane reforming (SMR). By varying synthesis conditions, samples with two different metal loading (17 wt% and 30 wt%) and different particle size distribution were prepared and characterized. Experimental results evidence that the temperature reached on the catalyst surface is related to the metal loading and to the particles size distribution that strongly affect the ability of ferromagnetic nanoparticles to convert the externally applied radio frequency field into heat. Catalyst pellets proved their effectiveness reaching the temperature of 720 °C during SMR reaction and 80% methane conversion.

Keywords:

Induction heating
Magnetic catalysis
Steam reforming
Ferromagnetic nanoparticles

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Introduction

Magnetic nanoparticles immersed in an alternating electromagnetic field dissipate energy, generating heat [1–4]. The exploitation of this heat for the purpose of energetically feeding chemical reactions is a novelty and has the potential to be a revolutionary technique in the construction of highly energy-efficient reactors [5–12]. In fact, the ability of radio-frequency induction to transfer energy directly to the material to be heated (catalytic bed), thanks to the selective absorption of energy by magnetic materials, allows to supply heat exclusively at the solid phase catalysing the reaction, leaving the feeding fluid mass and the reactor walls relatively cold. In contrast to conventional heating where heat is delivered from the outside to the inside of the reactor, induction heating generates heat right where is needed overcoming an important limit of the heating systems currently in use, where impressive thermal gradients between the reactor walls and the heart of the catalytic bed are observed. Furthermore, induction heating promotes process intensification since the non-catalysed parasitic reactions, however activated by temperature, are considerably reduced. With the instantaneous generation of localized heat directly or nearby the chemically active site, in principle it is possible to supply only the energy necessary to induce the catalytic reaction, considerably reducing heat dissipation and waste of material and energy with significant advantages in terms of efficiency of production processes.

Moreover, the heating rate can reach hundreds of degrees per minute allowing rapid reactor heating and shutdown. This has enormous advantages in terms of process security, allowing greater control in exothermic processes. An additional innovative aspect is that the high heating rate reached by magnetic induction offers the great opportunity to couple endothermic processes to discontinuous energy sources. The possibility of activating chemical reactions “on-demand” has recently attracted a great interest and several examples of induction heated, or microwave heated, chemical processes have been proposed [13–21].

While on the one hand the possibility of transferring contactless heat to the catalyst is very attractive because it immediately suggests energy savings, on the other hand the generation of an RF field is an expensive process that requires the use of electricity. Almind et al. have recently reported considerations on the energy transfer efficiency [22] for the steam methane reforming powered by induction. They developed a model that extrapolating lab scale experimental data to large systems predicted energy efficiency above 80%,

allowing the technology to be competitive with other electricity driven routes to hydrogen production.

In our previous research, a mixed metal catalyst with the Ni₆₀Co₄₀ (mole ratio) composition has been developed for the dry reforming of methane. The catalyst, consisting in nickel-cobalt alloy powders activated by milling, is able at the same time to catalyse the chemical process and deliver the heat necessary to run the endothermic reaction, reaching 900 °C when heated by induction [23,24]. For such material, considering the micrometric size of the particles, the heating mechanism was predominantly due to eddy currents induced on the surface of metallic powders. The catalyst proved to be active and initial stability tests were encouraging. On the other hand, the use of metal powders as such is uneconomical since, being the catalyst specific surface area very low, the overall catalytic activity is rather poor.

With the intention of preparing a more performing catalyst, we synthesized Ni₆₀Co₄₀ supported nanoparticles having a higher specific surface area of the active phase with respect to that of unsupported metal particles. However, unlike what has been observed for the nickel cobalt alloy micrometric powders [24], for ferromagnetic nanoparticles the dominant energy dissipation mechanism causing heat generation is unlikely to be attributable to eddy currents but will mainly be related to magnetic hysteresis and relaxation losses [2,25]. For this class of materials the particle size can significantly influence the heating properties [26,27]. Therefore, to achieve an appropriate temperature and an adequate energy load to feed the process, it is desirable to optimize the synthesis of the magnetic particles. For this purpose, an impregnation method was adopted using commercial γ -Al₂O₃ catalyst support and Ni₆₀Co₄₀ metallic nanoparticles were grown by changing synthesis parameters.

The steam methane reforming reaction [28] was selected as a model reaction to test the performances of the prepared samples because most of the industrial hydrogen demand is satisfied by the implementation of this highly endothermic process ($\Delta H^\circ = 206 \text{ kJ mol}^{-1}$).

The process is usually described by Eq. (1), often accompanied by competing or side reactions such as the water gas shift (2), the methane cracking (3) or carbon gasification (4), whose occurrence depends on process conditions:





The majority of industrial catalysts [29,30] are based on nickel, although many catalysts being researched now have better performance at laboratory scale [31,32].

In industrial reformers, heat transfer from the furnace to the catalyst bed is considered the source of principal energy waste [33–36] and an elaborate heat exchange network is utilized to increase the process efficiency. In this sense, induction heating could provide a useful solution, delivering heat in remote right where is needed and reducing losses.

In the following, we report the preparation of Ni₆₀Co₄₀ nanoparticles supported on commercial γ -Al₂O₃ pellets under different synthetic conditions (metal loading and calcination temperature) and their performance toward the steam methane reforming reaction powered by induction heating. The results obtained from the physical-chemical characterization on the metal alloy particles are combined with their heating performances to discuss the effect on the catalytic activity.

Materials and methods

Catalyst preparation and characterization

Catalysts are prepared using cylindrical (1/8") pellets of γ -Al₂O₃ (Alfa Aesar) as a support. Support specific surface area is 250 m² g⁻¹. Catalysts have been prepared by equilibrium adsorption [37], dipping alumina pellets (batches of 180 pellets) in a Ni(NO₃)₂*6H₂O (Merck-EMSURE) and Co(NO₃)₂*6H₂O (Merck-EMSURE) aqueous solution at room temperature and stirring the solution overnight. Two solutions having different total concentration, 4 M and 8 M respectively, with the same Ni:Co ratio were prepared. Excess solution was then removed and the samples were dried at 80 °C overnight. The as prepared samples, loaded with metal precursors, were subjected to a calcination treatment to remove the nitrates in air flow (50 mL min⁻¹) inside a horizontal tubular furnace (Carbolite) for 1 h. Three different calcination temperatures, i.e. 450 °C, 600 °C and 750 °C were used to evaluate their effect on the metal particle size, their distribution and ultimately on the catalytic activity. One hour was sufficient for the precursor complete elimination. Few samples were stored in glass desiccator before calcination (6 months) and in the manuscript are labeled as "aged". One sample was calcined in a static air atmosphere (see Table 1). Finally, the pellets were reduced by Temperature Programmed Reduction technique (TPR) in Ar–H₂ (3%vol.) flow (50 mL min⁻¹) with a heating rate of 10 °C min⁻¹ at 900 °C and maintaining the temperature for 5 h using a Micromeritics Autochem II 2920. Samples are labeled as NiCoxY, where x stands for the metal total loading and Y for the calcination temperature.

Atomic Absorption Spectroscopy (AAS) (Varian Spectra AA-220) was utilized to evaluate the amount of metal in the sample.

X-ray diffraction (XRD) analyses were carried on powders (pellets are easily grinded in a mortar) employing a diffractometer equipped with a 120 linear simultaneous detector

Table.1 List of prepared samples. Effect of synthesis conditions on particle size.

Sample	Me (wt %)	T calc (°C)	Cristallite size ^a (nm)	Particle size ^b (nm)
NiCo ₁₇ 450	17	450	7 ± 1	9 ± 2
NiCo ₁₇ 600	17	600	9 ± 2	10 ± 2
NiCo ₁₇ 600*	17	600	20 ± 2	23 ± 5
NiCo ₁₇ 750	17	750	24 ± 3	29 ± 5
NiCo ₁₇ 750*	17	750	28 ± 2	29 ± 5
NiCo ₃₀ 450*	30	450	32 ± 1	38 ± 9
NiCo ₃₀ 600	30	600	19 ± 2	18 ± 5
NiCo ₃₀ 600*	30	600	25 ± 2	23 ± 5
NiCo ₃₀ 600**	30	600	80 ± 2	52 ± 15
NiCo ₃₀ 750	30	750	45 ± 2	35 ± 12

^a By XRD Analysis applying the Debye-Scherrer equation.

^b By SEM Analysis, applying the Lognormal size distribution curve to size distribution histograms; statistical analysis from over 400 particles.

* Sample aged in glass desiccator before calcination (6 months).

** Calcination in static atmosphere (air).

from INEL and a monochromatized Fe K_{α1} source (Italstructure X-RED300).

The morphology of the catalyst pellets has been investigated by Scanning Electron Microscopy (SEM) using a high-resolution microscope HRSEM LEO130. Measurements were performed on both external and internal surface of the pellets, cleaving the sample into several pieces. To highlight the presence of the metallic phase dispersed on the alumina substrate, images were collected using a RBSD (backscatter) detector. To gain information on metal particles size and its distribution SEM images were processed by ImageJ [38] software. Size distribution histograms were fitted using a lognormal distribution function and evaluated Median particles diameters are reported. Specific surface area was evaluated by nitrogen adsorption at 77 K (BET method) [39] utilizing a Quantachrome NOVA 2200 apparatus.

Induction heating setup

Functional tests have been performed utilizing the catalysis flow line equipment schematized in Fig. 1. The core of the line is a tubular quartz reactor (internal diameter 2 cm) where the catalyst pellets are placed inside and held in place by a quartz frit. Typically, 18 catalyst pellets (about 0.5 g) positioned to form a single layer on the frit are used. An induction coil is positioned around the tube and the samples occupy the center of the coil to make the most of the magnetic field. An Ambrell EasyHeat 2.4 induction apparatus has been utilized to heat the catalysts. The heating module (work head) is connected by r. f. cables to the power supply, allowing an easy positioning of the coil around the reactor. The coil can be adjusted to have the samples positioned at its center. The apparatus can deliver a maximum power of 2 kW and operates in the 150–450 kHz frequency range. In this work, an 8 turns inductor coil of 2.5 cm diameter and 4.5 cm high working at 190 kHz is utilized. Power consumption versus generated magnetic field is re-reported in Ref. [23] for the empty coil, the linear response being

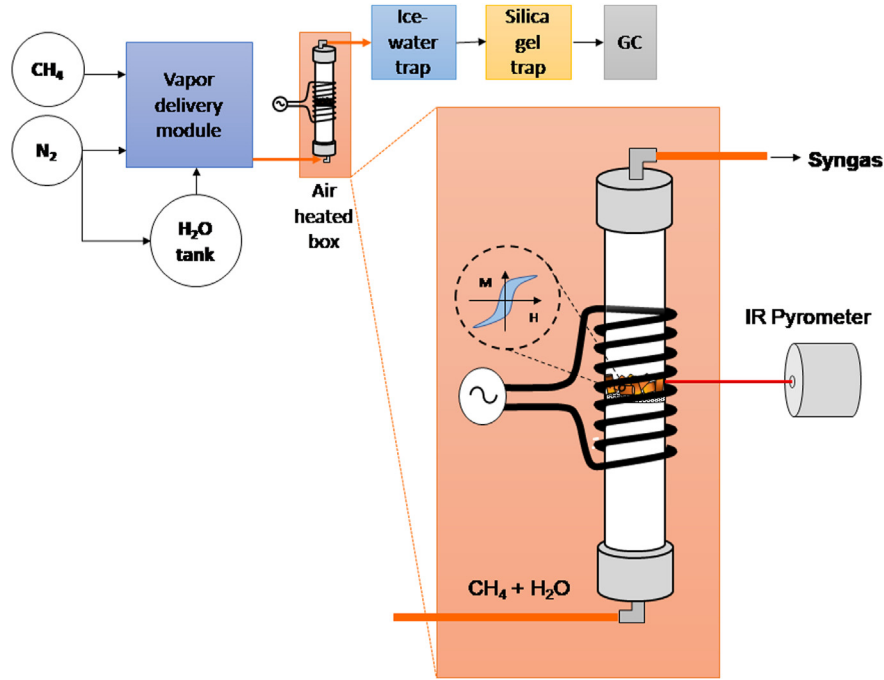


Fig.1 Outline of the catalyst testing line. Liquid water is vaporized inside a Vapour Delivery Module and conveyed to the reactor through heated lines. To prevent water condensation, the induction-heated reactor is placed inside an air-heated box. Typically, 18 catalyst pellets are utilized and placed in the same relative position within the coil. An IR pyrometer, positioned 15 cm from the catalyst, continuously measures the value of the average surface temperature of the catalyst (laser beam diameter 0.5 mm).

limited to 35 mT. The use of 18 pellets inside the inductor does not change the power consumed appreciably. Field values are calculated based on the current applied to the inductor. All catalytic activity measurements were performed at 49.1 mT. Although from an energy efficiency point of view it would have been more convenient to work within 35 mT range [23], it was a priority to clarify the limits of the material in terms of maximum obtainable temperatures and conversions rather than to optimize the energy transfer efficiency.

The temperature was measured continuously by focusing an IR pyrometer (IRTech Radiamatic IR20CF2150) on the surface of the pellets setting an emissivity value equal to 0.9. In this way the pyrometer provides the temperature of the catalyst positioned at the external reactor surface. It is reasonable to deduce that inner pellets may experience a higher temperature. A mixture of water vapour and methane (always in the ratio 2:1, balanced with N₂), are conveyed to the reactor by a Vapour Delivery Module (VDM) (Bronkhorst), setting different flows. Gas Hourly Space Velocity (GHSV) values reported in the text are calculated on the basis of the geometrical volume occupied by catalyst pellets.

To avoid water condensation, the lines connecting the VDM to the reactor are heated at 110 °C by Omegalux rope heaters. For the same reason, the quartz reactor is placed inside an air-heated box (110 °C) thus avoiding the reflux of water.

Two traps in series (a water/ice trap and a silica gel one) condense un-reacted water to prevent it from reaching the gas

chromatograph. The composition of reactant and product gas mixtures was analysed by a gas chromatograph Agilent GC490 equipped with two independent columns (MS5A and Plot-U).

The conversion of methane X_{CH_4} is calculated by Eq. (5):

$$X_{CH_4} = 100 \cdot \frac{F_{tot}^{in} \cdot CH_4^{in} - F_{tot}^{out} \cdot CH_4^{out}}{F_{tot}^{in} \cdot CH_4^{in}} \quad (5)$$

where F_{tot}^{in} is the total inlet flow, F_{tot}^{out} is the total outlet flow; CH_4^{in} and CH_4^{out} are the inlet or outlet volume fraction measured by the gas chromatograph.

The primary energy used by the inductor and transformed into chemical energy is reported in terms of energy transfer efficiency of the system. This value (equation (6)) has been calculated as the ratio between the energy consumed by the reaction (per unit time), i.e. the enthalpy content of the mixture leaving the reactor subtracted from the enthalpy content of the gas entering the reactor, and the power needed to generate the magnetic field [22]. Enthalpy content at the experimental temperature has been determined on the basis of thermodynamic data [40] and the experimental mixture molar fraction:

$$\eta = \frac{P_{reaction}}{P_{input}} \quad (6)$$

where $P_{reaction}$ has been evaluated according to

$$P_{reaction} = \sum_i H_i \cdot F_i \quad (7)$$

where H_i is enthalpy content of i 'th compound and F_i is its flow inside (negative) or outside (positive) the reactor. The induction apparatus displays the value of the input power (P_{input}) once the operator sets the current flowing through the coil.

Results and discussion

Catalyst characterization

In order to dissipate heat in the presence of an alternating magnetic field, catalyst nanoparticles must preserve their ferromagnetic properties at the reforming operating temperature ($T > 700$ °C). Nickel alone, although a perfect catalyst for reforming reactions, has a Curie temperature T_c (temperature at which a transition between the ferromagnetic and paramagnetic phases occurs) equal to 354 °C, way below the process requirements. Addition of cobalt ($T_c = 1121$ °C) to nickel in the selected composition ($Ni_{60}Co_{40}$) raises T_c value to ~780 °C [41], preserving at the same time catalytic activity toward the selected process [9,42,43].

Since the amount of generated heat is dependent, among other parameters, on the amount of dissipating agent, catalyst with considerable metal loading are required. In fact, for this class of materials the metal alloy particles have a dual task, acting as catalysts for the chosen reaction and as heat suppliers to reach the appropriate temperatures.

The list of produced samples is shown in Table 1. The results of AAS showed that two batches with a total metal loading of about 17 and 30 wt% respectively were obtained, with a metal ratio equal to the nominal composition (60:40) of the solution adopted for the catalyst preparation.

An example of nitrogen adsorption/desorption isotherm is reported in Fig. 2 for two samples, both calcined at 750 °C and subsequently reduced, having two different metal loadings.

Nitrogen adsorption isotherms of all prepared samples exhibit a type-IV (IUPAC) shape typical of mesoporous solids

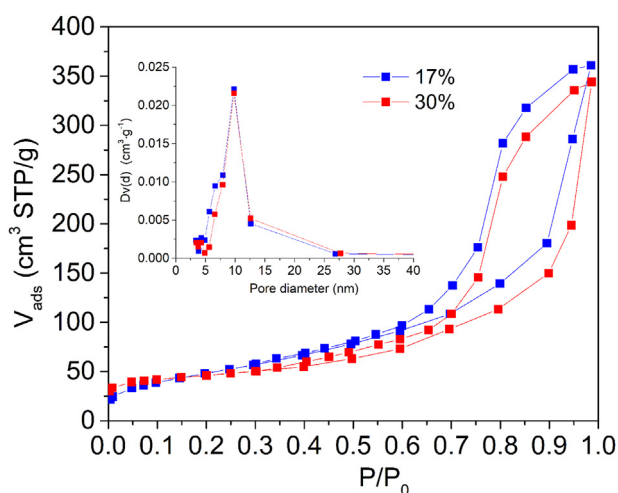


Fig.2 Nitrogen adsorption/desorption isotherms recorded at 77 K for two samples (calcination temperature 750 °C, reduction temperature 900 °C) with different metal loading, 17% and 30% ($NiCo_{17}750$, $NiCo_{30}750$).

characterized by a large uptake and a wide hysteresis at high partial pressure, explainable with the presence of a large mesoporous volume typical of the alumina substrate. Small differences between the two curves reported in Fig. 2 for the 17 wt% and 30 wt% $Ni_{60}Co_{40}$ supported catalysts are likely due to minor differences in the porosity distribution. The inset in Fig. 2 shows the pores size distribution curve in the mesoporous region evaluated by the BJH analysis [44]. Both samples show a relatively narrow distribution curve, a maximum in the porosity distribution centered on 10 nm and small differences for smaller diameter. The BET surface area of the γ - Al_2O_3 support was 250 $m^2 g^{-1}$ and drops to 190 $m^2 g^{-1}$ after the 5 h thermal treatment at 900 °C ($Ar-H_2(3\%vol.)$). The surface area of supported catalysts after the same treatment decreased further to about 180 $m^2 g^{-1}$ on $NiCo_{17}$ samples and to about 110 $m^2 g^{-1}$ on $NiCo_{30}$ ones, showing that samples surface morphology is measurably affected by the addition of increasing amount of nickel and cobalt. Differently, the calcination temperature of the catalyst oxidic precursor more mildly affects the catalyst surface area. The same effect is observed for pore volume that was 0.80 $cm^3 g^{-1}$ for thermally treated alumina and decreased by metal addition to about 0.58 $cm^3 g^{-1}$ on all $NiCo_{17}$ samples and 0.54 $cm^3 g^{-1}$ on $NiCo_{30}$. Introduction of metal content in the substrate resulted in the reduction of the measured adsorbed volume in the pores, indicating blockage of pores mouth. Such values are still considered appropriate for a catalyst.

XRD patterns for the sample batch with metal loading 17 wt% calcined at 450 °C, 600 °C and 750 °C (Fig. 3) showed, irrespective of the calcination temperature, the presence of diffraction peaks of alumina, nickel and cobalt oxides, and no peaks from metal aluminates. The same applies for samples with metal loading 30%. The peaks of Co_3O_4 phase are well visible already at the lower calcination temperature, whereas those of NiO appeared well resolved only after calcination at 750 °C, despite the higher relative amount of Ni in the sample. This fact suggests that NiO segregation on Al_2O_3 occurs to a

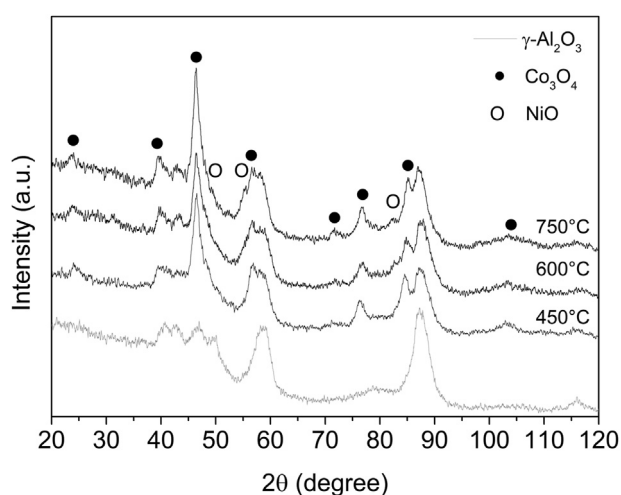


Fig.3 XRD patterns of samples calcined at three different temperatures: 450 °C, 600 °C and 750 °C, metal loading 17%, are reported together with pattern relative to bare alumina substrate.

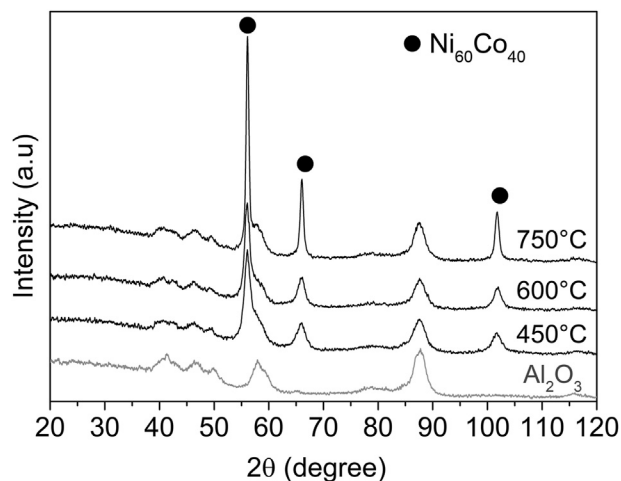


Fig. 4 XRD patterns of the bare alumina substrate thermally treated at 900 °C for 5 h and of the samples calcined at 450 °C, 600 °C and 750 °C, after reduction at 900 °C for 5 h ($\text{NiCo}_{17/450}$, $\text{NiCo}_{17/600}$, and $\text{NiCo}_{17/750}$). Sample metal loading 17%. NiCo alloy is determined by comparison with ICDD-JCPDS database cards (#4-850 and #15-806).

lower extent with respect to Co-oxide species, possibly due to an interaction of Ni with Al_2O_3 surface stronger than that of Co. XRD patterns for the corresponding reduced samples are reported in Fig. 4 together with the diffraction profile of the bare support, which underwent the same thermal treatment (900 °C, Ar-H_2 , 5 h). The three samples show a very similar pattern profile, where, together with the substrate, only the peaks belonging to a single fcc metallic phase are visible, suggesting that the reduction reaction is complete. The cell parameter for the metallic phase is the same ($3.532(1)$ Å) for all the samples, an evidence of the same stoichiometry. Its value is in a good agreement with the one expected for an alloy $\text{Ni}_{60}\text{Co}_{40}$, for which the cell parameter linearly changes with the composition, from 3.5238 Å for pure Ni fcc and 3.5447 Å for pure Co fcc [45]. The crystallite size of the metallic phase is calculated applying the Scherrer's equation on peak broad-ening and listed in Table 1. The crystallite size increases with the sample loading, aging time and calcination temperature, as expected [46]. At the same temperature, calcination in static atmosphere rather than flowing conditions produces larger particle size, possibly due to the presence of non-negligible water vapour pressure during the process [47,48].

The TPR output results for the NiCo_{17} and NiCo_{30} samples calcined at different temperatures showed a different hydrogen consumption profile as a function of temperature (Fig. 5). As the reduction was not complete after reaching 900 °C, an isotherm reduction for 5 h was required to obtain the complete reduction of oxides (total H_2 consumption evaluated from the peak integrated area, >99%) and the stabilization of the support morphology. The peak at lower temperature is assigned to Co_3O_4 reduction to CoO, as confirmed by XRD patterns performed on samples reduced at 350 °C (not shown). The reduction of nickel oxide phase cannot be easily separated from the reduction of cobalt (II) oxide phase and a broad peak is detectable starting above

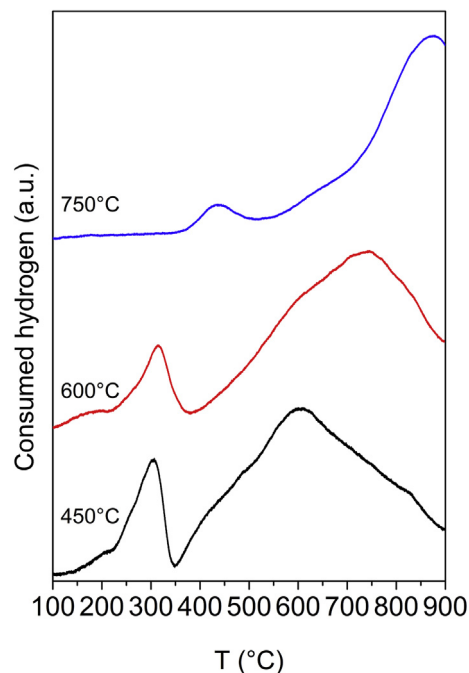
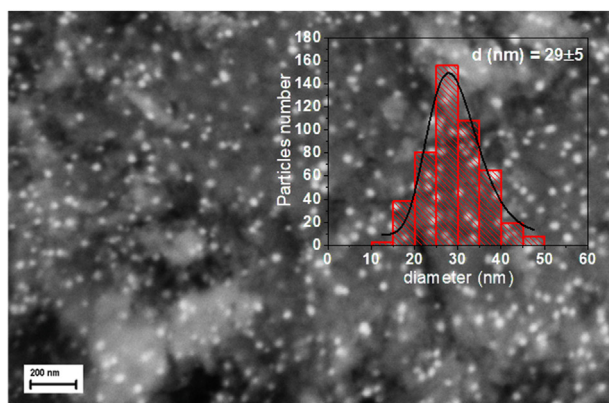


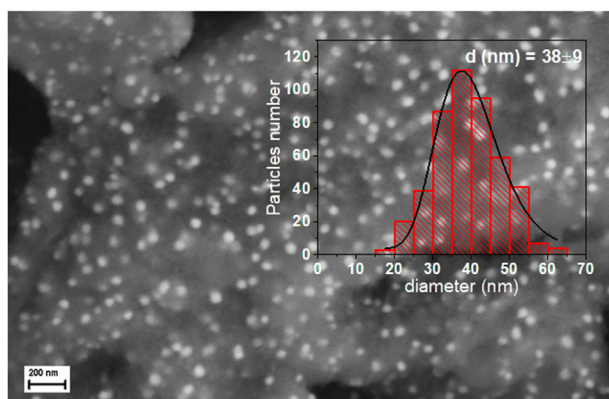
Fig.5 TPR profiles of samples calcined at 450 °C, 600 °C and 750 °C measured in in ArH_2 (3%vol.). Sample metal loading 17%. Temperature scan rate 10 °C m i n^{-1} .

350 °C for samples calcined at 450 °C and 600 °C and above 500 °C for the sample calcined at 750 °C. The position of such broad reduction peak does not change significantly for the samples calcined at 600 °C and 750 °C. We assume that the broad reaction peak represents the reduction of Ni(II) and Co(II) oxides to form a $\text{Ni}_{60}\text{Co}_{40}$ alloy. The T_{max} of the broad peaks shifted towards higher temperature with increasing the calcination temperature of the sample, possibly due to a greater interaction of catalyst precursors with the support. The same trend was found for the NiCo_{30} samples at different calcination temperature.

The SEM images of all samples showed that the metal nanoparticles (light gray dots in the image due to their elemental composition different from the substrate) are evenly distributed within the porous alumina structure. In Fig. 6 the SEM images showing the morphology of two reduced samples $\text{NiCo}_{17/750}$ (a) and $\text{NiCo}_{30/450}$ * (b) are re-reported. Particles size distribution histogram, included as an inset in the same figure, yielded the particle average diameter (Table 1) that are in line with crystallite size calculated from XRD data, an indication that metallic particles are mostly single crystalline domain. The particles diameter and the corresponding spreading increase with metal loadings. Depending on the synthesis conditions, samples loaded at 17% show median diameters in the range 9–29 nm while for samples loaded at 30% the size distribution curve shifts to higher values, 18–52 nm. Correspondingly, the width of the size distribution in-crases from 5 to 15 nm, indicating that lower metal loading favors also a narrower size distribution.



(a)



(b)

Fig.6 SEM images and size distribution histograms (top inset) of $\text{Ni}_{60}\text{Co}_{40}$ nanoparticles supported on $\text{g-Al}_2\text{O}_3$: (a) sample NiCo_{17750} and (b) sample NiCo_{30450}^* .

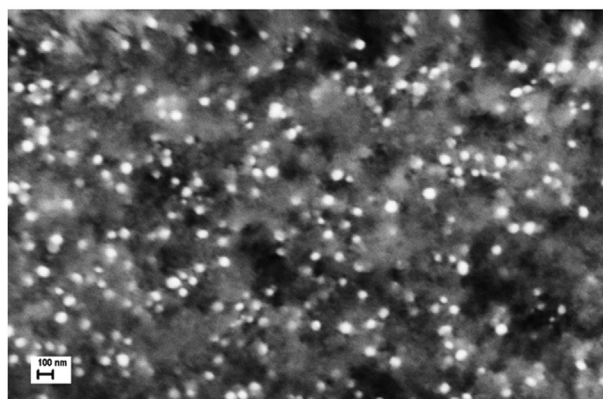
Two additional SEM images are shown in Fig. 7. In Fig. 7a is reported an image at low magnification to highlight the homogeneity of metal particle dispersion on the substrate, even for samples with high metal loading (30%). In Fig. 7b is shown an image relative to NiCo_{30600}^{**} whose particles have the biggest mean diameter obtained (Table 1). For this sample a static calcination (instead of the one in flow) was adopted during the preparation procedure. Even in this condition, that favored the growth of oxide particle size, it is possible to see that the metallic phase is well dispersed in isolated spherical particles, indicating a certain interaction alloy-support.

Heating performances and catalytic activity

The temperature reached in the catalytic bed under nitrogen flow (90 mL min^{-1}) for different values of applied magnetic field is presented in Fig. 8 for NiCo_{30600} . Such trend is typical for all the samples, whereas the maximum temperature reached by each sample is different (Table 2) and depends on nanoparticles features. Indeed, as it will be later discussed, the heat generation rate can be governed by the amount of magnetic material loaded on the support and by the nanoparticles size. The temperature increases as the applied field increases up to a "saturation value", such that a further raise of the external field causes a negligible increase in



(a)



(b)

Fig.7 SEM images of two samples loaded at 30%, calcined under different conditions. Section (a) NiCo_{30600} . Section (b) NiCo_{30600}^{**} (static calcination).

temperature, at the expense of a considerable rise in power required to generate the field [23]. In the inset of Fig. 8 the variation of temperature as a function of time is recorded for a fixed input power. A steep increase in temperature is observed as soon as the field is switched on. After few minutes the observed temperature is the balance between heat dissipated by the nanoparticles and radiative and convective losses from the surface of the pellets. The heating system let to reach about 95% of the maximum temperature value in 2 min. When the external field is removed the temperature rapidly drops down.

The NiCo catalysts heated by induction were active for the methane steam reforming reaction, reaching temperature in the range $490\text{--}620 \text{ }^\circ\text{C}$ and correspondent CH_4 conversion in the range about 20 and 60% (Table 2). Measured values are in line with conversion data reported in the literature in this temperature range [49]. The effect of the flow rate over the methane conversion on the catalyst NiCo_{30600} is reported in Fig. 9 keeping the applied input power constant, where essentially it appears that reactions are carried out in the kinetic regime.

Fig. 10a shows the composition of the outlet gas mixture for NiCo_{17600} sample on a dry basis at the temperature reached by induction heating under reactant flow ($525 \text{ }^\circ\text{C}$). In all cases, the reaction temperature was slightly lower than

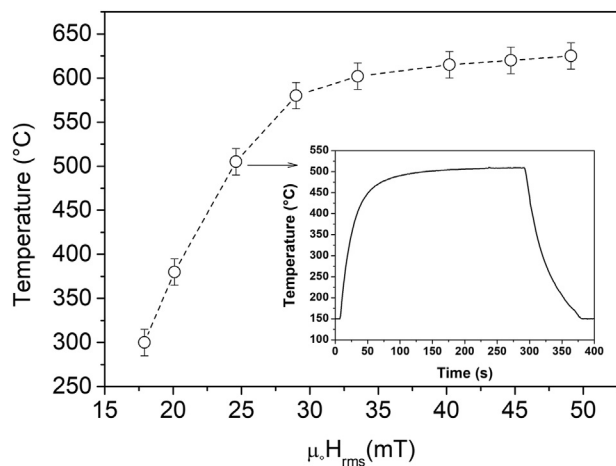


Fig.8 Temperature measured by the pyrometer focused on the catalytic bed external surface (18 pellets of NiCo₃₀600*) as a function of the magnetic field amplitude. The inset shows the heating profile at I $\frac{1}{4}$ 110 A, B $\frac{1}{4}$ 24.6 mT, H $\frac{1}{4}$ 19.59 kA m⁻¹, N₂ flow $\frac{1}{4}$ 90 mL min⁻¹. Once the current passing through the inductor is switched off, the temperature rapidly drops demonstrating the advantage of a rapid on/off switching associated with induction heating.

that measured under N₂ flow. In the present experiment, when the reactive mixture reaches the catalyst, the temperature decreases from 560 °C to about 525 °C due to contact with gas at a higher specific heat (water and methane compared to nitrogen) and also due to the absorbed heat of reaction (endothermic). As soon as methane and steam reach the catalyst, hydrogen and carbon monoxide are produced and their values rapidly reach a plateau. On the contrary, carbon dioxide is observed only after few minutes and increases with time-on-stream, reaching a plateau in 20 min. The conversion of methane was around 35%, in line with the equilibrium value corresponding at the measured temperature indicating the good activity of the catalyst [39]. The considerable amount of carbon dioxide observed is also in line with the measured temperature values. Actually, at this temperature, the water gas shift reaction is favored, and carbon dioxide and hydrogen are produced at expenses of carbon monoxide, whose quantity in the mixture is limited to 4% by volume.

For NiCo₃₀ catalysts (high metal content) it is immediately evident that methane conversion and carbon monoxide amount are higher compared to those on samples with lower metal loading. This is certainly due to the higher values of temperature that catalysts reach compared to those of the NiCo₁₇ series (see Table 2). Fig. 10b shows the conversion of methane and product distribution for the most performing sample (NiCo₃₀450*), for which the measured temperature is 665 °C under nitrogen flow, and 618 °C during the process. The measured methane conversion is the one expected at such temperature, indicating also for this high metal loading catalyst the good catalytic performance. These data showed

Table 2 Temperature on samples surface when heated by induction T_{N₂}, and during the reforming reaction, T_{reac}; methane conversion, X_{CH₄}, and catalyst reaction rate. Data measured on 18 pellets of catalyst.

Sample	T _{N₂} (°C) ^a	T _{reac} (°C) ^{a,b}	X _{CH₄} (%)	Reaction rate ^c (Mol _{CH₄} g _{Me} ⁻¹ h ⁻¹)
NiCo ₁₇ 450	520	490	19	0.40
NiCo ₁₇ 600	560	525	36	0.74
NiCo ₁₇ 600*	575	550	44	0.93
NiCo ₁₇ 750	570	535	38	0.80
NiCo ₁₇ 750*	536	510	33	0.69
NiCo ₃₀ 450*	665	618	60	0.58
NiCo ₃₀ 600	570	530	45	0.45
NiCo ₃₀ 600*	630	600	56	0.55
NiCo ₃₀ 600**	415	400	15	0.15
NiCo ₃₀ 750	625	605	48	0.50

^a Heating parameters: I = 220 A, B = 49.1 mT, H = 39.19 kAm⁻¹, P = 0.9 kW in nitrogen flow (N₂ = 90 mL min⁻¹). The estimated error was ± 10 °C due to difficulties in focusing the IR pyrometer on the pellets external surface.

^b Reactant flow: CH₄ = 50 mL min⁻¹, H₂O = 100 mL min⁻¹.

^c Calculated from the amount (moles) of methane converted.

that, under the same magnetic field condition, a higher metal loading favors the attainment of higher temperatures, as the number of dissipating agents increases.

The capability of magnetic nanoparticles to dissipate energy in an alternating magnetic field, usually reported as Specific Power Loss (SPL i.e. the amount of heat generated per unit gram of magnetic material and per unit time) depends on several parameters [3,27,50–53]. Essentially, the ability of magnetic nanoparticles to act as heat mediators is affected by their composition, structural, morphological and magnetic properties (intrinsic parameters directly dependent on the sample preparation method) and the amplitudes and frequency of the applied field (extrinsic parameters), not straightforward predictable because of several mechanisms

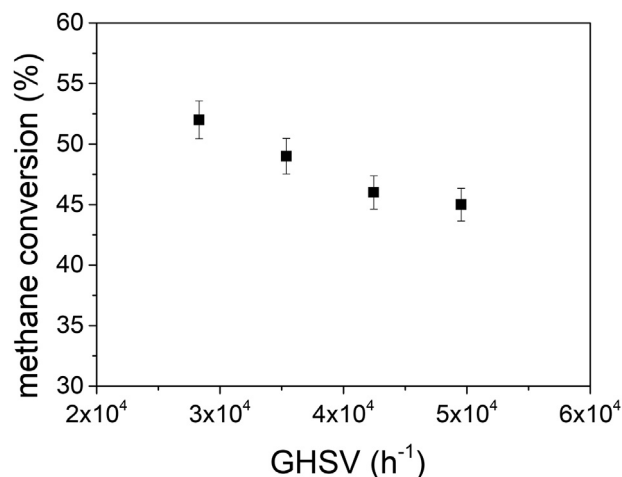


Fig.9 Percent methane conversion as a function of the gas hourly space velocity (h⁻¹) defined as the ratio of the volumetric feed flow rate (normal condition) to the volume of the catalyst bed, geometrically determined. Sample NiCo₃₀600.

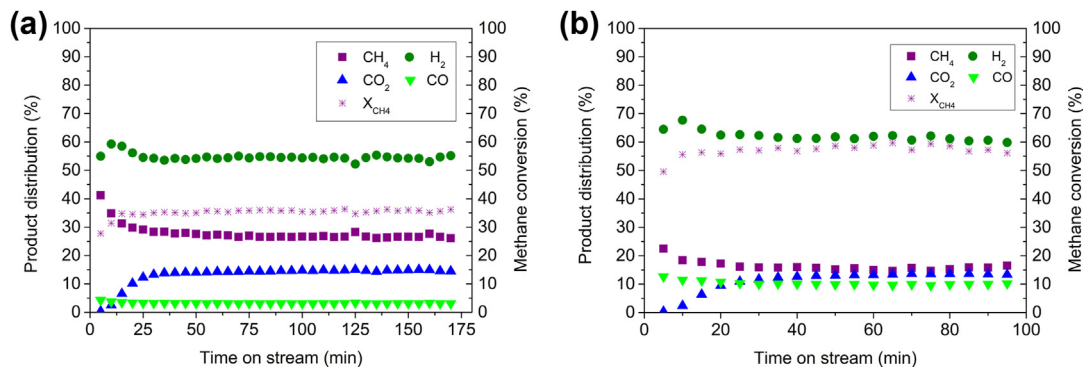


Fig.10 Composition of the outlet mixture and methane conversion at the reaction temperature (T_{reac}) reached upon radiofrequency heating. Section (a) sample NiCo₁₇/600 (T_{reac} ¼ 525 °C); section (b) NiCo₃₀/450* (T_{reac} ¼ 618 °C). Applied magnetic field 49.1 mT, I ¼ 220 A. Volumetric inlet flow: N₂ ¼ 90 mL min⁻¹, C H₄ ¼ 50 mL min⁻¹, H₂O ¼ 100 mL min⁻¹, (GHSV ¼ 28,310 h⁻¹).

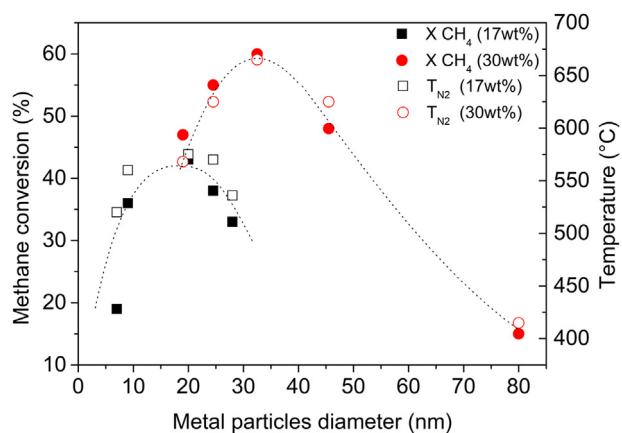


Fig.11 The measured methane conversion as a function of Ni₆₀Co₄₀ alloy particle diameter. In the same picture the temperature measured on the sample surface (T_{N_2}), which accounts for the sample ability to dissipate energy under nitrogen flow (90 mL min⁻¹) is reported. The dotted line is a guide for the eye.

contributing to magnetic losses. Experimental work is therefore often required in view of materials optimization. An attempt to rationalize the results is shown in Fig. 11 reporting for all the samples the methane conversion and the temperature reached on the sample surface in nitrogen flow, T_{N_2} , as a function of the Ni₆₀Co₄₀ particle diameter. All measurements were carried out keeping constant the number of catalyst pellets in the reactor, the gas flow and the applied magnetic field (frequency and amplitude). The data were grouped according to the sample metal loading. It is evident a strong dependence of T_{N_2} (and therefore of the conversion of methane which directly depends on it) on the size of the nanoparticles. Considering the samples with lower loading, it can be observed how for particles smaller than 20 nm T_{N_2} increases as the size increases and then descends as the latter increases. The same profile can be observed for samples with higher loading, with the difference that the maximum of the curve is shifted to 30–35 nm. These temperature profiles follow the dependence of the Specific Power Loss on particles

size, which has been modeled by Hergt [48] for magnetic iron oxide nanoparticles for hyperthermia. Such profile is the result of different magnetic loss mechanisms intervening in the heat dissipation. It is known [2,4,5,48,54] that hysteresis loss and relaxation loss are the two major power loss mechanisms of magnetic induction heating and that exists a critical particle size below which relaxation loss dominates, or else the hysteresis loss prevails. Then, in induction heated catalytic processes, in order to enhance the heat dissipation efficiency of the sample, it is of great importance the control of synthesis process to obtain particles with optimal diameter. As can be seen in Table 2, for samples loaded with the same amount of metal (30%), a very wide range of the T_{N_2} reached by induction heating (~200 °C) was measured, indicating very different dissipation ability for materials with the same nominal composition. For application purposes, the use of a sample such as NiCo₃₀/600** would be highly inefficient.

What is unexpected is that the trend curve drawn for the set of samples loaded at 30% has its maximum moved to slightly larger size when instead we would have expected, given the same chemical composition of the particles, that the two curves showed a parallel trend. It is probable that the effect of a wider distribution of particle size for samples with 30% metal content, compared to samples with 17% is such that moves the curve toward larger dimensions [2]. Magnetic interaction among nanoparticles can also be taken in consideration as a possible cause. In order to uniquely assess the cause of such shift a detailed magnetic characterization, beyond the scope of the present work, is needed.

To further clarify the role of particle size in determining the observed catalytic behavior we compare for all catalysts the rate of CH₄ converted *per* gram of metal in the catalyst (mol CH₄ g⁻¹ Me⁻¹ h⁻¹) whose values for dilute NiCo₁₇ samples are almost all higher than those for concentrated NiCo₃₀ samples (Table 2). Particles with similar size (~20 nm) NiCo₁₇/600* and NiCo₃₀/600 yield the same T_{N_2} but different rate, higher on dilute catalyst where they are in a lower amount. This suggests a different nature of the exposed surface in concentrated samples. However, samples with the same metal content but having particles with different mean diameter (i.e. different exposed metal surface values) yield similar T_{N_2} and similar CH₄

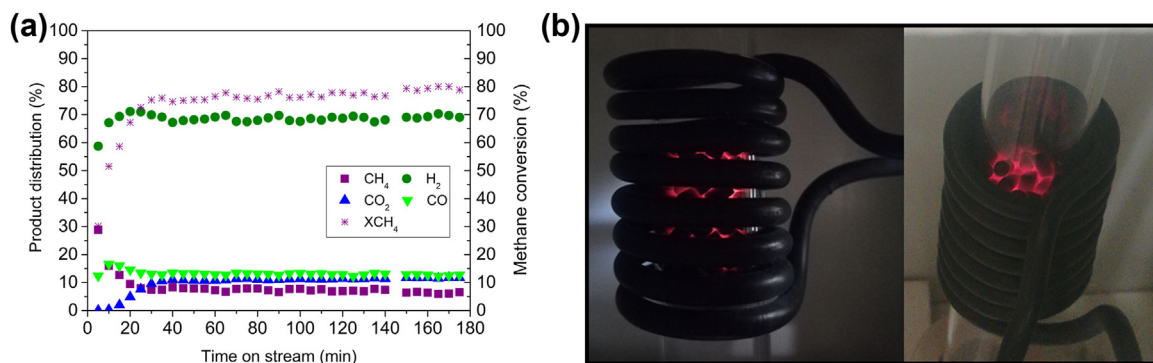


Fig. 12 (a) Experimental product distribution and methane conversion values for sample NiCo₃₀600. Volumetric inlet flow: N₂ ¼ 90 mL min⁻¹, C H₄ ¼ 50 mL min⁻¹, H₂O ¼ 100 mL min⁻¹. GHSV ¼ 2800 h⁻¹. T_{reac} ¼ 720 °C. Section (b): a digital photograph of the NiCo₃₀600 catalysts heated by induction at work (180 pellets housed in the quartz reactor placed inside the coil). The current applied to the coil was 1 ¼ 220 A.

conversions and rates (see Table 2 and Fig. 11, samples NiCo₁₇600 and NiCo₁₇750*). These results suggest that the dependence of the reaction rate on the exposed metal surface value affects the reaction rate less than temperature values.

In the experiments reported so far, approximately 0.5 g of catalysts (18 pellets) have been used which occupy a geometric volume equal to 0.5 mL. Such volume it is only a small fraction of inductor volume (≈ 20 mL) and indeed the power delivered to the inductor does not change appreciably when the magnetic catalyst is placed inside. In this configuration, the best result in terms of methane conversion ($X_{\text{CH}_4} = 60\%$, when T_{N_2} was 665 °C, see Table 2) has been obtained with the NiCo₃₀450* sample that shows a catalyst productivity in the range 0.17–0.25 mol_{CH₄} g⁻¹ cat⁻¹ h⁻¹ depending on the methane flowrate conveyed to the reactor. The reaction rate was calculated considering the moles per hour of methane converted during the process over the total mass of the supported catalyst (Al₂O₃–Ni₆₀Co₄₀). The reaction rate rises to 0.58–0.84 mol h⁻¹ g⁻¹ Me⁻¹ considering the amount of the metal only, as reported in Table 2.

In order to better exploit the magnetic field generated inside the inductor, the reactor was loaded with a greater amount of catalyst (180 pellets of NiCo₃₀600, volume occupied in reactor ≈ 10 mL, central part of the coil). Results are reported in Fig. 12 together with images of the reactor in operation. T_{N_2} value rises from 630 °C, obtained using 18 pellets of catalyst, to 740 °C, obtained with 180 pellets, because a greater number of pellets participate in the dissipative process. It is therefore expected that at this thermal level the conversion of methane to carbon monoxide and hydrogen is higher than what has been seen so far. Indeed, the measured methane conversion reaches 80% and the amount of carbon dioxide further decreases, since the reaction of water gas shift is less favored at this temperature (H₂/CO ratio ~ 6). At the same time, the input power needed to generate the magnetic field (49.1 mT) increases from 900 W (empty coil) to 970 W. In the reported experimental conditions, the energy transfer efficiency is around 1.75%. The main factors influencing this value are related to our specific experimental set-up, which is not optimized. As previously discussed, when the quantity of catalyst has been increased ten times, the temperature on the

catalyst surface increases considerably due to minimization of thermal losses, leading to an increase of the conversion value. At the same time, due to instrumental set-up limitations, it was not possible to increase the flow of reagent sent the catalyst and the catalytic activity remains low, the process being limited by diffusional reasons. By a simple approximation, using the same GHSV of the previous experiments the efficiency would raise to 16%. An optimization of the catalyst test plant from an engineering point of view will certainly provide higher efficiency values [22].

Finally, initial stability tests have shown that the catalyst is stable in several successive work cycles showing conversion values equal to the initial one, an encouraging detail in view of long-term stability tests.

Conclusion

The progress in the design and development of materials for magnetic catalysis has the potential to allow the implementation of highly energy-intensive processes on small-scale, compact reactors, thanks to the rapid and targeted response of ferromagnetic nanoparticles to an applied magnetic field. Ferromagnetic Ni₆₀Co₄₀ nanoparticles supported on commercial γ -Al₂O₃ pellets proved to be active for the steam methane reforming reaction, when inductively heated by a radiofrequency alternating magnetic field, demonstrating their dual function as heat dissipating agent and active catalyst for the chemical process. Despite the high metal loading of the synthesized catalysts (17% and 30%), Ni₆₀Co₄₀ nanoparticles are well dispersed on the support, suggesting that the selected synthetic methodology, easily scalable to mass production, is appropriate. In concentrated catalysts the higher concentration of dissipating agent maximizes the productivity per mass of catalyst; in dilute catalysts the presence of small nanoparticles maximizes activity per metal site. The measured methane conversion values give indication of an important and peculiar dependence on the metal particle size distribution that reflects the ability of the nanoparticles to dissipate energy into heat upon the application of the magnetic field. Indeed, the size of the metal particles is a critical

key factor to obtain a higher heating rate and an efficient conversion of magnetic energy into heat, essential to maximize the overall catalyst performance. This finding is particularly relevant for high temperature endothermic reactions in which the effectiveness of heat transfer is a fundamental parameter in defining the efficiency of the process. Optimization of synthesis parameters is therefore of utmost importance to obtain nanoparticles with fine-tuned features in terms of dissipating agents and active surface. Some of the materials designed and produced achieve methane conversions close the target value; efficiency must be improved optimizing operative parameters of reforming reaction functional tests.

Declaration of competing interest

The authors declare that they have no known competing financial interest or personal relationships that could have appeared to influence the work reported in this paper.

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