Synthesis, Characterization and Kinetic Behavior of Supported Cobalt Catalysts for Oxidative after-Treatment of Methane Lean Mixtures

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Abstract: The present work addresses the influence of the support on the catalytic behavior of Co₂O₄-based catalysts in the combustion of lean methane present in the exhaust gases from natural gas vehicular engines. Three different supports were selected, namely γ-alumina, magnesia and ceria and the corresponding catalysts were loaded with a nominal cobalt content of 30 wt. %. The samples were characterized by N₂ physisorption, wavelength dispersive X-ray fluorescence (WDXRF), X-ray diffraction (XRD), Raman spectroscopy, X-ray photoelectron spectroscopy (XPS) and temperature-programmed reduction with hydrogen and methane. The performance was negatively influenced by a strong cobalt-support interaction, which in turn reduced the amount of active cobalt species as Co₂O₄. Hence, when alumina or magnesia supports were employed, the formation of CoAl₂O₄ or Co–Mg mixed oxides, respectively, with a low reducibility was evident, while ceria showed a lower affinity for deposited cobalt and this remained essentially as Co₂O₄. Furthermore, the observed partial insertion of Ce into the Co₂O₄ lattice played a beneficial role in promoting the oxygen mobility at low temperatures and consequently the catalytic activity. This catalyst also exhibited a good thermal stability while the presence of water vapor in the feedstream induced a partial inhibition, which was found to be completely reversible.

Keywords: supported cobalt catalysts; spinel cobalt oxide; methane oxidation; magnesia; ceria; γ-alumina; cobalt aluminate; Co-Mg mixed oxides; oxygen mobility: Mars–van Krevelen mechanism

1. Introduction

Compressed natural gas is regarded as a suitable alternative to substitute the traditional automotive fuels such as gasoline or diesel that are becoming more expensive and scarce with time. Natural gas vehicles have been demonstrated to produce less CO₂, NOₓ and soot emissions than gasoline or diesel vehicles and are safer in case of accident [1,2]. Nevertheless, the application of this technology is accompanied by the necessity of controlling the emissions of unburned methane from the engine, as this is a powerful greenhouse effect gas. The most commonly applied solution is the complete oxidation over a supported noble metal catalyst, such as platinum and/or palladium [3,4]. However, the price of these metals is extremely high and they are also prone to deactivation by sintering and the presence of water, and thus, this raises the cost of natural gas engines and limits their massive implementation. For this reason, the interest in developing noble-metal free catalysts for methane oxidation is increasing.
Cobalt oxide-based catalysts, and among them spinel-type cobalt oxide (Co$_3$O$_4$), are considered good alternative candidates to noble metals due to their already demonstrated high efficiency for the oxidation of hydrocarbons and a greater availability [5–7]. The reason for this high activity seems to lie on their good redox properties such as reducibility and mobility of oxygen species at low temperatures, which derives from the easiness that the constituent ions of these materials have to shift between oxidation states. Moreover, these catalysts are more thermally stable than noble-metal based catalysts and generally more resistant to water inhibition [8,9]. However, cobalt oxides tend to present poor structural and textural properties as well, especially when they are prepared by common synthesis methods such as precipitation, sol–gel or solution combustion [10,11]. For this reason, these materials are usually supported over porous materials as a way to improve their properties, and also as a way to facilitate their incorporation into the monolithic systems where they should eventually operate [12,13].

The selection of an appropriate support is not a trivial task and can have a significant effect on the properties of the final catalyst, due to the different nature of the cobalt-support interactions, to the point of even rendering the catalyst useless for the specific purpose under study. For cobalt oxide catalysts the most commonly used supports are alumina [14,15], magnesia [16,17], zirconia [18,19], silica [20,21] ceria [22,23], silicon carbide [24,25], zeolites [26,27] or cordierite [28,29]. The decision of using one or other support for a specific cobalt-based catalyst is generally made on the basis of the specific catalytic properties that regulate the activity for the reaction under study. Regarding this, many studies have dealt with the effect of specific supports on the performance of Co$_3$O$_4$ catalysts for different reactions. For instance, Grzybek et al. [30] found out that cobalt oxide showed very different behavior for N$_2$O abatement depending on which polymorph of alumina was used as the support, and concluded that it was better to sacrifice the textural properties of the final catalyst by using low-surface $\alpha$-Al$_2$O$_3$ instead of $\gamma$-Al$_2$O$_3$ with the objective of inhibiting the occurrence of cobalt-alumina interactions. This effect was also found by Solsona et al. [31], when studying the total oxidation of propane with cobalt oxide supported over alumina with low, medium and high surface area. On the other hand, Yung et al. [32] examined the oxidation of NO with cobalt catalysts supported over titania and zirconia, finding that the latter was a more suitable support for this purpose than the former. However, on a different study, Kim et al., [33] reported that high-surface ceria was a better support than titania or zirconia for the same reaction. Ceria was also found to be, as pointed out by Wyrwalski et al. [34], the most suitable support for cobalt oxide for the complete oxidation of propene. This type of studies has been also carried out for liquid phase reactions. For example, Zhang et al. [35] analyzed the influence of several supports for the degradation of organic dyes in solution. They concluded that MgO was the most suitable for this purpose due to the increase in the population of surface Co$^{2+}$ ions induced by cobalt–magnesia interactions.

In the case of methane oxidation, it has been demonstrated that the population of Co$^{3+}$ species in the spinel lattice is the key parameter that provides the catalyst with the good reducibility and oxygen mobility involved in the Mars–van Krevelen mechanism [36,37]. Thus, an appropriate support for this reaction will be one that enables a high dispersion of the cobalt deposited, while at the same time allows the cobalt oxide species to maintain their good redox properties as intact as possible. However, this is not always achievable, as a high dispersion of the deposited cobalt is usually accompanied by a strong cobalt-support interaction that, more often than not, ends up being detrimental for the oxygen mobility of the final catalyst [38].

A wide number of works have investigated the oxidation of methane over supported cobalt catalysts, each one focusing on a specific support, under different reaction conditions, with different degrees of success [39–41]. However, there are no studies about the effect that supports with a varying physico-chemical nature have on the fundamental properties of cobalt oxide-based catalysts, and the comparison of their activity under the same conditions. For this reason, in the present work, three Co$_3$O$_4$ catalysts supported over $\gamma$-alumina, magnesia and ceria were prepared by the same synthesis route, characterized and examined for the oxidation of methane under lean conditions, with the
objective of determining the effect that the different supports have on the textural, structural and redox properties and the activity of the cobalt oxide active phase.

2. Materials and Methods

2.1. Synthesis of the Supports and Supported Cobalt Catalysts

Three different supports (γ-Al₂O₃, MgO and CeO₂) were used for preparing the cobalt catalysts. The employed alumina was kindly provided by Saint-Gobain (Paris, France). This was previously thermally stabilized at 850 °C for 4 h in static air. Both MgO and CeO₂ were prepared by precipitation with an aqueous solution of Na₂CO₃ (CAS 497-19-8) 1.2 M. This was slowly added to aqueous solutions of magnesium (II) nitrate hexahydrate (Mg(NO₃)₂ 6H₂O, CAS 13446-18-9) or cerium (III) nitrate (Ce(NO₃)₃ 6H₂O, CAS 10294-41-4), respectively, at a constant temperature of 80 °C, until the pH was 8.5 or 9.5. A similar synthesis route was followed for preparing the supported cobalt samples. Thus, for each support, 5 g of the selected support were mixed with 100 cm³ of a solution of Co(NO₃)₂·6H₂O (CAS 10026-22-9) with adjusted concentration and then a solution of Na₂CO₃ 1.2 M was added dropwise at 80 °C until the pH reached 8.5. The nominal Co content of the three metal oxide catalysts was 30 wt. %.

All samples were dried at 110 °C for 16 h and then calcined in static air to obtain the final supports (MgO and CeO₂) and cobalt catalysts. The calcination protocol consisted on three heating ramps separated by 30-min isothermal steps at 125 and 300 °C: An initial ramp at 5 °C min⁻¹ from room temperature to 125 °C, a second ramp at 1 °C min⁻¹ up to 300 °C, and a final ramp at 5 °C min⁻¹ up to 600 °C, temperature that was then kept constant for 4 h.

2.2. Characterization Techniques

Textural properties of the supports and catalysts were examined by N₂ physisorption in a Micromeritics TriStar II apparatus (Micromeritics Instrument Corp. Norcross, GA, USA). The Brunauer-Emmett-Teller BET (Brunauer-Emmett-Teller) method was used to determine the specific surface area of the samples while the BJH (Barrett, Joyner and Halenda) method was applied for the estimation of the average pore size. Degassing of the samples prior to analysis was performed on a Micromeritics SmartPrep apparatus (Micromeritics Instrument Corp. Norcross, GA, USA). at 300 °C for 10 h with a N₂ flow. The elemental composition of the cobalt catalysts was determined by wavelength dispersive X-ray fluorescence (WDXRF). Each sample was mixed with a flux agent (Spectromelt A12, Merck 111802, Darmstadt, Germany) in an approximate proportion of 20:1 and placed in an induction micro-furnace at 1200 °C to form a boron glass pearl. The pearls were analyzed under vacuum in a PANalytical AXIOS sequential WDXRF spectrometer (Malvern Panalytical Ltd, Royston, UK), equipped with a Rh tube and three different detectors (gas flow, scintillation and Xe sealed).

Structural properties of the catalysts were determined by X-Ray diffraction and Raman spectroscopy. XRD analysis was performed on a X’PERT-PRO X-Ray diffractometer (Malvern Panalytical Ltd, Royston, UK), using Cu Kα radiation (λ = 1.5406 Å) and a Ni filter. The X-Ray source was operated at 40 kV and 40 mA of current. The diffractograms were obtained in the 2θ range of 5–80° with a step size of 0.026° and a counting time of 2.0 s. Phase identification was carried out by comparing the diffraction patterns with JCPDS (Joint Committee on Powder Diffraction Standards) database cards. Additionally, in order to perform a detailed XRD analysis over the supported cobalt catalysts a longer counting time (26.8 s) was applied. The cell size of the cobalt spinel phase was estimated by profile matching of the detailed XRD patterns using FullProf.2k software (version 6.30, Institut Laue-Langevin, Grenoble, France).

The analysis by Raman spectroscopy was carried out by using a Renishaw InVia Raman spectrometer (Renishaw, Wotton-under-Edge, Gloucestershire, UK), coupled to a Leica DMLM microscope (Wetzlar, Germany) with a spatial resolution of 2 microns. For each spectrum, 20 s were
employed and five scans were accumulated with the 10% of the maximum power of a 514 nm laser (ion-argon laser, Modu-Laser, Centerville, UT, USA) in a spectral window of 150–1500 cm$^{-1}$. X-ray photoelectron spectroscopy (XPS) analysis was performed using a SPECS system (SPECS GmbH, Berlin, Germany) equipped with a Phoibos 150 1D analyzer (SPECS GmbH, Berlin, Germany) and a DLD-monochromatic radiation source. The obtained spectra were calibrated by fixing the signal of adventitious carbon at 284.6 eV.

Temperature-programmed reduction with hydrogen (H$_2$-TPR) analysis were carried out on a Micromeritics Autochem 2920 apparatus (Micromeritics Instrument Corp. Norcross, GA, USA), using a 5%H$_2$/Ar mixture as the reducing gas. Each sample was subjected to a pre-treatment with a 5% O$_2$/He mixture at 300 °C for 30 min prior to the analysis. All TPR experiments were performed up to 950 °C with an isothermal step of 10 min at that temperature. The water produced throughout each experiment was removed from the outlet stream using a cold trap, to avoid interference with the thermal conductivity detector. Additional information regarding the activation of methane was obtained by means of temperature programmed reaction with a 5% CH$_4$/He mixture in the absence of oxygen (CH$_4$-TPR) coupled to mass spectrometry (MKS Cirrus Quadrupole Mass Spectrometer, Andover, MA, USA). The experiments were conducted up to 600 °C with a heating ramp of 10 °C min$^{-1}$ followed by an isothermal step at 600 °C for 30 min.

2.3. Evaluation of the Catalytic Performance

Catalytic activity was examined in a bench-scale fixed bed reactor (PID Eng&Tech S.L. Madrid, Spain) in the 300–600 °C temperature range at atmospheric pressure. In each reaction experiment, 1 g of catalyst was used (particle size of 0.25–0.3 mm). The catalyst was diluted with the same mass of inert quartz (particle size 0.5–0.8 mm) to ensure a good distribution of heat and reactants along the catalytic bed. The feedstream (1%CH$_4$, 10%O$_2$ and N$_2$ as the balance gas) was fed to the reactor with a total flow of 500 cm$^3$ min$^{-1}$, which corresponded to a space velocity of 300 cm$^3$ CH$_4$ g$^{-1}$ h$^{-1}$ (60,000 h$^{-1}$ approximately for an estimated catalyst density of 2 g cm$^{-3}$). The temperature of the reactor was increased in a stepwise progression, with heating ramps of 1 °C min$^{-1}$ followed by 15-min isothermal periods each 25 °C, where methane conversion and product profiles were determined. Each chromatographic analysis was performed in triplicate. Methane conversion was calculated by the difference between inlet and outlet CH$_4$ concentrations. Inlet and outlet streams were analyzed using an on-line Agilent Technologies 7890N gas chromatograph (Agilent Technologies, Santa Clara, CA, USA) equipped with a thermal conductivity detector (TCD) and two columns: for the analysis of CH$_4$, O$_2$, N$_2$ and CO, a PLOT 5A molecular sieve column was used. For CO$_2$ analysis, a PLOT U column was used. To ensure that mass or heat transfer limitations were not affecting the obtained kinetic results, the criteria for intra and extra-particle mass diffusion, heat transfer and temperature gradients were checked to be above the limits, according to the Eurokin procedure [42,43].

3. Results

3.1. Characterization of the Supports

The textural properties of the commercial γ-alumina support and the as-prepared magnesia and ceria supports in terms of BET surface area, pore volume and mean pore diameter are shown in Table 1. Notable differences were noticed among the investigated supports. Hence, the commercial alumina showed the largest specific surface area (136 m$^2$ g$^{-1}$), followed by MgO (80 m$^2$ g$^{-1}$) and CeO$_2$ (8 m$^2$ g$^{-1}$). This decreasing order was also consistent for the estimated pore volume of the samples. It varied from 0.55 cm$^3$ g$^{-1}$ (γ-alumina) to 0.08 cm$^3$ g$^{-1}$ (ceria). The samples showed type IV isotherms with H2 hysteresis loops.
Table 1. Textural properties of the supports and the supported cobalt catalysts.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Surface Area, m² g⁻¹</th>
<th>Pore Volume, cm³ g⁻¹</th>
<th>Mean Pore Diameter, Å</th>
<th>Cobalt Content, wt.%</th>
<th>Support Crystallite Size, nm</th>
<th>Co₃O₄ Crystallite Size, nm</th>
<th>Co/M Molar Ratio ¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al₂O₃</td>
<td>136</td>
<td>0.55</td>
<td>123</td>
<td>-</td>
<td>5</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>MgO</td>
<td>80</td>
<td>0.19</td>
<td>106</td>
<td>-</td>
<td>21</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CeO₂</td>
<td>8</td>
<td>0.03</td>
<td>230</td>
<td>-</td>
<td>32</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Co/Al₂O₃</td>
<td>108</td>
<td>0.29</td>
<td>89</td>
<td>27.9</td>
<td>6</td>
<td>29²</td>
<td>0.43 (0.39)</td>
</tr>
<tr>
<td>Co/MgO</td>
<td>47</td>
<td>0.16</td>
<td>204</td>
<td>31.9</td>
<td>16</td>
<td>17</td>
<td>0.12 (0.38)</td>
</tr>
<tr>
<td>Co/CeO₂</td>
<td>18</td>
<td>0.07</td>
<td>225</td>
<td>28.9</td>
<td>33</td>
<td>44</td>
<td>3.15 (1.39)</td>
</tr>
</tbody>
</table>

¹ The values in brackets correspond to the molar ratio determined by X-ray fluorescence (XRF). M stands for metal of the oxide support (Al, Mg and Ce, respectively). ² This value corresponds to the size of the spinel Co₃O₄-Co₃Al₂O₄ phases present in this sample, which are indistinguishable by XRD.

On the other hand, XRD patterns of the as-prepared supports well matched with those expected for the pure materials (Figure 1). Hence, the observed diffraction signals could be indexed as γ-alumina (2θ at 37.7, 45.8 and 67.3°, JCPDS 01-074-2206), magnesium oxide (2θ at 43.0, 62.3, 74.7 and 78.6°, JCPDS 00-004-0829) and cerium oxide (2θ at 28.5, 33.3, 47.5, 56.4 and 76.7°, JCPDS 00-004-0593). Moreover, the crystallinity of both magnesia and ceria was higher than that of alumina in view of their noticeably more intense and sharper signals. The formation of ceria was further corroborated by Raman spectroscopy, which revealed a strong peak assigned to the F₃ Raman-active mode characteristic of the fluorite-like lattice of CeO₂. Note that the vibrational modes of MgO and γ-Al₂O₃ are essentially Raman inactive.

![Figure 1. XRD patterns of the bare supports and the supported cobalt catalysts.](image-url)

3.2. Characterization of the Supported Cobalt Catalysts

Table 1 lists the cobalt loading of the synthesized cobalt catalysts as determined by WDXRF and ICP-AES (Inductively Coupled Plasma-Atomic Emission Spectroscopy) in the case of the Co/CeO₂ sample. It was verified that this content was relatively close to the nominal value (30% wt. Co). BET measurements revealed that cobalt species markedly blocked the pores of the alumina and magnesia supports, as evidenced by the notable decrease in the surface area of the Co/Al₂O₃ and Co/MgO catalysts, from 136 to 108 m² g⁻¹ (26%) and 80 to 47 m² g⁻¹ (42%), respectively. As with the supports,
the catalysts showed type IV isotherms with H2 hysteresis loops as well. Both samples presented a lower pore volume with respect to their corresponding support (0.29 and 0.16 cm³ g⁻¹, respectively). By contrast, the impact on the textural properties on the Co/CeO₂ was less noticeable. Thus, a slight increase in both surface area and pore volume was found (18 m² g⁻¹ and 0.07 cm³ g⁻¹, respectively).

The comparative analysis of the pore size distributions of the supports and the cobalt catalysts (Figure 2) evidenced that the deposition process of cobalt particles was highly dependent on the pore accessibility and interconnectivity. Hence, when cobalt was deposited over γ-alumina, which exhibited a bimodal distribution centered at 90 and 150 Å, the cobalt preferentially deposited over its largest pores. Conversely, when using magnesia as a support, characterized by pores with a markedly different size (35–50 and 325 Å), the cobalt favorably located over the smaller pores. Finally, in the case of ceria, both support and catalyst possessed a unimodal distribution centered around 225 Å, but with an increased width for the cobalt catalyst. This could be due to the fact that cobalt species did not find enough space to deposit on the pores of ceria, and subsequently located on its external surface as well. In addition, since the amount of pores of 335 Å (the prevalent pore size of bulk Co₃O₄) was larger in the Co/CeO₂ catalyst than in the ceria support, it could be assumed that this catalyst contained segregated cobalt oxide to some extent.

Figure 2. Pore size distributions of the bare supports and the supported cobalt catalysts.

Figure 1 includes the diffractograms of the cobalt catalysts. Their patterns were characterized by the presence of Co₃O₄ (2θ = 31.3, 37.0, 45.1, 59.4 and 65.3°, JCPDS 00-042-1467) along with some weak signals corresponding to the respective support (2θ = 43.0 and 62.3° and 2θ at 28.5, 47.5 and 56.4°, respectively). On the other hand, it is worth pointing out that the formation of CoAl₂O₄ is very frequent in Co/alumina systems due to the strong interaction between Co₃O₄ and the support at mild temperatures (>450 °C) [44,45]. However, from the XRD analysis the extent of the formation of this undesired phase was not possible. Note that both spinel-like cobalt phases (Co₃O₄ and CoAl₂O₄, JCPDS 00-044-0160) crystallize in the cubic structure with comparable cell parameters, thereby showing very close 2θ values in their diffraction patterns. In addition to that, the crystallinity of the support phases did not noticeably change since their crystallite size was similar before and after cobalt deposition (Table 1). However, the crystallite size of the cobalt spinel was highly dependent on the employed
The smallest crystallite size was obtained over the magnesia (17 nm) while the largest size was found over the ceria (44 nm). This finding was consistent with the poorer textural properties of the synthesized ceria, which resulted in a preferential location of Co3O4 in its external surface. Nevertheless, in all cases the crystallite size was smaller than that of the bulk Co3O4 (63 nm), which evidenced a good dispersion of the cobalt spinel over the surface of the studied supports.

The Raman spectra in the 150–900 cm−1 region of the cobalt catalysts supported on alumina, magnesia and ceria are shown in Figure 3. As a reference, the spectrum of pure Co3O4 is shown as well. Apart from a relatively intense band at 462 cm−1 (F2g mode of CoO2) for the Co/ CeO2 catalyst, all supported catalysts displayed the five Raman actives modes attributable to Co3O4, namely three F2g modes located at 194, 519 and 617 cm−1, and the Eg and A1g modes at 479 cm−1 and 687 cm−1, respectively [46]. In addition, two shoulders at 705 and 725 cm−1 attached to the A1g vibration mode were also visible in the case of the Co/ Al2O3 catalyst. These two signals evidenced the presence of cobalt aluminate in this sample [47,48]. As far as the Co/MgO catalyst was concerned, it should be remarked that small bands at about 1250 and 1350 cm−1 were detected (not shown). These signals could be associated with a certain increase in the disorder of the structure of MgO owing to the insertion of cobalt ions leading to the formation of a Co–Mg solid solution [49,50]. Note that the formation of this mixed oxide was difficult to verify by XRD since no significant changes in the 2θ diffraction angles of the Co/MgO catalysts were noted with respect to those of pure MgO.

![Figure 3. Raman spectra of the supported cobalt catalysts.](image)

A closer inspection of the A1g mode as function of the type of used support could be helpful in determining the properties of the lattice of deposited Co3O4. This influence was analyzed in terms of the shift and the full width at half maximum (FWHM) of this signal. The Raman spectra of a bulk Co3O4 were used as a reference. Thus, this Raman mode was located at 687 cm−1 and its FWHM was 11 cm−1. While no significant shift in the position of the band was found for the Co/Al2O3 (688 cm−1) and Co/MgO (688 cm−1) catalysts, a notable shift to a lower frequency (681 cm−1) was noted over the Co/CeO2 sample. This redshift of the signal could be assigned to the distortion of the spinel lattice, probably owing to insertion of Ce ions [51]. On the other hand, the largest FWHM value for the Co/MgO catalyst was assigned to the presence of Co–Mg mixed oxides. The lattice distortion in the cobalt spinel generated by the insertion of Ce ions and/or formation of Co–Mg mixed oxides was
further evidenced by a slight increase in the cell parameter of the Co$_3$O$_4$ phase for both Co/MgO (8.083 Å) and Co/CeO$_2$ (8.082 Å) catalysts with respect to the bulk Co$_3$O$_4$ sample (8.052 Å), where the distortion of the cobalt spinel was minimal.

The surface composition of the samples was investigated by XPS. The Co2p spectra of the supported cobalt catalysts are shown in Figure 4, along with the spectrum of bulk Co$_3$O$_4$ for the sake of comparison. All samples showed broad signals suggesting the presence of various different cobalt species on the surface of the catalysts. More specifically, all spectra showed the main Co2p$_{3/2}$ signal in the position range 781.4–779.9 eV, along with two satellite signals centered around 785.9–786.7 eV and 789.5–790.2 eV, that were attributed to the presence of Co$^{2+}$ and Co$^{3+}$ ions, respectively [52].

The position of the main signal exhibited a shift with respect to the position for bulk Co$_3$O$_4$ (779.9 eV), which depended upon the chosen support. For the Co/CeO$_2$ catalyst, the position of the main signal and the intensity of the satellite signals were comparable to that of the bulk sample. This pointed out that the nature of the cobalt oxide for this catalyst was similar to that of the bulk sample. The main signal of the Co/MgO samples was shifted towards higher binding energy values while the intensity of the Co$^{2+}$ satellite signal was notably stronger with respect to the bulk oxide. Both features were compatible with a higher presence of Co$^{2+}$ ions in the surface of the Co/MgO catalyst [53], as a result of the Co–Mg interaction and the subsequent formation of the Co–Mg solid solution. Lastly, for the Co/Al$_2$O$_3$ catalyst the main signal was located at 781.4 eV, which was a position indicative of the presence of cobalt aluminate [54].

Besides, the surface composition was determined from the integration of the XPS spectra. The respective Co/M (M = Al, Mg and Ce) surface molar ratio could be then calculated and compared with the bulk ratio calculated from the XRF analysis. For both Co/Al$_2$O$_3$ and Co/CeO$_2$ catalysts their surface ratio (0.43 and 3.15, respectively) was higher than the bulk ratio (0.39 and 1.39, respectively), which indicated a more pronounced presence of cobalt on the surface of these samples. For the Co/MgO, however, the surface ratio (0.12) was notably lower than the bulk ratio (0.38). This could be due to the strong Co–Mg interaction and the partial insertion or dissolution of Co ions in the MgO lattice to form a Co–Mg solid solution, which in turn decreased the amount of cobalt present on the surface of this catalyst.

![Figure 4](image_url)

**Figure 4.** X-ray photoelectron spectroscopy (XPS) spectra of the supported cobalt catalysts.

The redox properties of the catalysts were studied by temperature-programmed reduction with hydrogen in the 50–950 °C temperature range. The corresponding profiles are displayed in Figure 5 while the quantitative results of the analysis are listed in Table 2. A noticeably different redox behavior
was found among the three different cobalt catalysts. Hence, the largest H₂ uptake was exhibited by the sample supported on ceria (7.6 mmol g⁻¹) followed by the catalysts supported on alumina (5.6 mmol g⁻¹) and magnesia (4.6 mmol g⁻¹). First, the redox properties of the latter two samples will be comparatively discussed since both γ-Al₂O₃ and MgO could be considered non-reducible in the studied temperature window. Hence, the observed H₂ consumption could be exclusively assigned to the reduction of the present cobalt species. In this sense, and taking as a reference the ideal specific H₂ uptake for the reduction of Co₃O₄ as the only cobalt phase (22.6 mmol H₂ g⁻¹Co), both Co/Al₂O₃ and Co/MgO catalysts revealed a significantly lower consumption, namely 18.7 and 14.5 mmol H₂ g⁻¹Co, respectively. Therefore, the estimated degrees of Co reduction were 88 and 64%. In line with the results given by Raman spectroscopy, these findings suggested that a fraction of deposited cobalt species strongly interacted with these supports, thereby negatively influencing their redox properties.

Figure 5. Temperature-programmed reduction with hydrogen (H₂-TPR) profiles of the supported cobalt catalysts.

Table 2. Redox properties of the supported cobalt catalysts derived from TPR analysis with H₂ and CH₄.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Total H₂ Uptake (^1) (mmol g⁻¹Co)</th>
<th>H₂ Uptake at Low Temperature (^1) (mmol g⁻¹Co)</th>
<th>Degree of Co Reduction (^2) (%)</th>
<th>O₂ Consumption at Low Temperature (CH₄-TPR) (^1) (mmol g⁻¹Co)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co/Al₂O₃</td>
<td>18.7 (5.6)</td>
<td>9.8 (2.7)</td>
<td>88</td>
<td>0.28</td>
</tr>
<tr>
<td>Co/MgO</td>
<td>14.5 (4.6)</td>
<td>9.9 (3.2)</td>
<td>64</td>
<td>0.51</td>
</tr>
<tr>
<td>Co/CoO₂</td>
<td>26.2 (7.6)</td>
<td>23.4 (6.8)</td>
<td>100</td>
<td>0.70</td>
</tr>
<tr>
<td>CeO₂</td>
<td>- (1.5)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

\(^1\) The values in brackets correspond to the H₂ uptake on a catalyst weight basis. \(^2\) This degree of reduction was estimated based on 22.6 mmol H₂ g⁻¹Co for full reduction of cobalt as Co₃O₄.

The reduction process of the Co/Al₂O₃ catalyst consisted on two main reduction events. The first event, centered at about 250–550 °C, could be attributed to the reduction of free Co₃O₄. This reduction process could be subsequently divided into other two features with peak reduction temperatures at 310 and 400 °C, following the same reduction process as for the bulk Co₃O₄ catalysts. This consisted of the sequential reduction to CoO and metallic Co, respectively [55,56]. An additional peak between 550–750 °C was clearly ascertained. This was attributed to the presence of significant amounts of
CoAl$_2$O$_4$ derived from the strong interaction between Co$_3$O$_4$ and Al$_2$O$_3$. It was quantitatively deduced that the total amount of cobalt of the sample was equally distributed as Co$_3$O$_4$ (49%) and CoAl$_2$O$_4$ (51%). It must be noted that all cobalt species present in the catalysts were completely reduced to metallic Co. This was verified by XRD analysis of the samples recovered after the TPR run where only metallic Co and Al$_2$O$_3$ were detected.

The H$_2$-TPR profile of the Co/MgO sample also revealed two distinct reduction regions although the temperatures windows were markedly different when compared with the alumina-supported counterpart. Hence, the observed consumption at low-temperature (200–350 °C) was ascribed to easily reducible free Co$_3$O$_4$ species as well. The band located at higher temperatures (350–650 °C) was assigned to the reduction of cobalt-MgO species formed during the synthesis route. The integration of these two features gave the following cobalt distribution, 57% as Co$_3$O$_4$ and 43% as Co–Mg mixed oxides. The formation of Co–Mg mixed oxides with a superior stability, which could not be reduced even at 950 °C, could not be however ruled out, as the total H$_2$ uptake of this sample was rather low in view of its Co content [17,57]. In fact, when assuming that the uptake at low temperatures (< 350 °C) was owing to the reduction of Co$_3$O$_4$ with a stoichiometry of 4 moles of H$_2$:3 moles of Co, and the uptake at higher temperatures (350–650 °C) was related to the reduction of Co$_2^+$ species with a stoichiometry of 1 mol of H$_2$:1 mol of Co, the corresponding metal content of the sample, in view of its overall H$_2$ uptake, would be equivalent to about 25% wt. This was considerably lower than the actual Co loading as determined by XRF (close to 32% wt.). Anyway, the amount of free Co$_3$O$_4$ with a significantly higher oxidation activity in comparison with CoAl$_2$O$_4$ or Co-Mg mixed oxides, was relatively similar for both γ-Al$_2$O$_3$ and MgO supported catalysts.

While the H$_2$ uptake of both pure alumina and magnesia was negligible, the bare CeO$_2$ sample exhibited a weak signal at 450–500 °C that corresponded to the surface reduction of the oxide whereas the notable H$_2$ consumption peaking at about 825 °C was related to the reduction of the bulk [58,59]. The TPR profile of the Co/CeO$_2$ catalyst was characterized by a remarkable uptake between 200–600 °C that was related to the reduction of precipitated Co$_3$O$_4$. Similarly to the Co/Al$_2$O$_3$ sample, this feature exhibited two fairly discernible peaks at 310 and 380 °C. A small band was noted at 800 °C as well, which corresponded to the reduction of the bulk of the support. Note that this occurred to slightly lower temperatures with respect to the bare support, probably due to the catalytic role played by cobalt. A quantitative analysis of the amount of consumed H$_2$ revealed that the overall uptake (7.6 mmol H$_2$ g$^{-1}$) reasonably matched with that theoretically expected for the total reduction of Co$_3$O$_4$ along with the reduction of ceria (1.8 mmol H$_2$ g$_{Ce}^{-1}$). This corresponded to a 100% degree of Co reduction (Table 2). At low temperatures, the amount of consumed H$_2$ was 23.4 mmol g$_{Co}^{-1}$, which was slightly larger than theoretically required for the reduction of the Co$_3$O$_4$ oxide (22.6 mmol g$_{Co}^{-1}$). This suggested that deposited cobalt facilitated the reduction of the surface of the ceria in this temperature range [60].

An overall overview of the redox properties of the three cobalt catalysts pointed out that the use of ceria was beneficial for obtaining a sample with a limited interaction of the active phase with the surface support, thereby not favoring the formation of hardly reducible cobalt oxides such as cobalt aluminate or cobalt–magnesium mixed oxides. In addition, cobalt helped in promoting the reducibility of the ceria.

More useful insights on the influence of the catalyst composition on the reactivity of available active oxygen species for methane oxidation were obtained by CH$_4$-TPR analysis coupled to mass spectrometry. The analysis was performed between 50 and 600 °C with a subsequent isothermal step at this temperature for 30 minutes. The evolution of CO$_2$ (m/z = 44) and CO (m/z = 28, not shown) was monitored (Figure 6). In the low temperature range, namely 375–525 °C, the generation of CO$_2$ was noticed over the three cobalt catalysts (peaking at about 485 °C). This was attributed to the oxidation of methane by oxygen species associated with Co$^{3+}$ ions. However, the extent of this reaction was considerably different over each sample in view of the comparatively larger amount of consumed oxygen (0.70 mmol O$_2$ g$^{-1}$) or larger yield of CO$_2$ over the Co/CeO$_2$ sample, followed by the Co/MgO (0.51 mmol O$_2$ g$^{-1}$) and Co/Al$_2$O$_3$ (0.28 mmol O$_2$ g$^{-1}$) catalysts. (Table 2) Moreover, the temperature
for the onset of reduction (marked by arrows in Figure 6) was significantly lower for the Co/CeO₂ sample (395 °C) in comparison with the other two catalysts, namely 415 and 425 °C over Co/MgO and Co/Al₂O₃, respectively.

Figure 6. Profiles of the supported cobalt catalysts (a). Close-up view of the 200–550 °C temperature range (b).

On the other hand, during the isothermal period at 600 °C a second oxidation process was only evidenced over the Co/CeO₂ sample. This was also accompanied by the generation of CO and H₂ to some extent that was related to partial oxidation or cracking of methane in the presence of metallic or oxygen-deficient cobalt species [61]. In fact, the diffraction pattern of the sample after the CH₄-TPR run evidenced the formation of graphitic carbon (signal at 2θ = 26.6°) [62]. The distinct formation at high temperatures of these species (CO₂, CO and H₂) was not observed over the other two catalysts, thereby revealing the presence of very stable, inactive cobalt species on these samples, such as cobalt aluminate or cobalt–magnesium mixed oxides, where cobalt was mainly present as Co²⁺.

3.3. Performance of the Supported Cobalt Catalysts

The performance of the cobalt catalysts was examined by their corresponding light-off curves at 300 cm³ CH₄ g⁻¹ h⁻¹ (30,000 cm³ g⁻¹ h⁻¹, about 60,000 h⁻¹) in the 200–600 °C temperature range. For each catalyst, three consecutive reaction cycles were conducted. In all cases the first light-off curve revealed slightly lower reaction temperatures while the second and third runs were identical to each other, as can be seen for the Co/CeO₂ catalyst in Figure S1 (Supplementary Materials). For this reason, Figure 7 compares the 3rd-cycle curve for each examined catalyst. Recall that only CO₂ was detected in the product stream in the whole temperature range. Hence, a 100% selectivity towards CO₂ formation was achieved for all tested catalysts. Appreciable methane conversion (>5%) was detected over 350 °C over the Co/CeO₂ catalysts while a similar conversion level was attained at significantly higher temperatures (400 °C) over the samples supported on both magnesia and alumina. The Tₐʊ₀ value (temperature at which 50% conversion was attained) was used as criterion for the relative reactivity of each sample (Table 3). In a similar way to the results observed in the low-conversion range, a substantially different performance was addressed with values close to 500 °C (Co/CeO₂), 525 °C (Co/MgO) and 550 °C (Co/Al₂O₃). Accordingly, conversion values at around 85% (Co/Al₂O₃), 95% (Co/MgO) and 98% (Co/CeO₂) were noted at 600 °C.
Table 3. Kinetic results of the oxidation of lean methane over the supported cobalt catalysts.

<table>
<thead>
<tr>
<th>Sample</th>
<th>$T_{50} \degree C$</th>
<th>Specific Rate at 425 $\degree C$ mmol CH$<em>4$ g$</em>{Co}^{-1}$ h$^{-1}$</th>
<th>$E_a$ kJ mol$^{-1}$</th>
<th>$\Delta H^\ddagger$ kJ mol$^{-1}$</th>
<th>$\Delta S^\ddagger$ J mol$^{-1}$ K$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co/Al$_2$O$_3$</td>
<td>550</td>
<td>1.8</td>
<td>90</td>
<td>84</td>
<td>$-166$</td>
</tr>
<tr>
<td>Co/MgO</td>
<td>525</td>
<td>1.4</td>
<td>102</td>
<td>98</td>
<td>$-139$</td>
</tr>
<tr>
<td>Co/CeO$_2$</td>
<td>500</td>
<td>3.1</td>
<td>82</td>
<td>78</td>
<td>$-161$</td>
</tr>
</tbody>
</table>

An additional evidence of the goodness of the Co/CeO$_2$ sample was given by the analysis of the specific reaction rate of the cobalt catalysts (Table 3). This reaction rate was calculated under differential conditions (conversion < 20%) at 425 $\degree C$. The ceria supported sample exhibited a markedly higher specific activity (3.1 mmol CH$_4$ g$_{Co}^{-1}$ h$^{-1}$) with respect to the other two counterparts, which showed a similar performance (1.4–1.8 mmol CH$_4$ g$_{Co}^{-1}$ h$^{-1}$).

On the basis of the fact that methane oxidative decomposition over cobalt catalysts requires highly active oxygen species [63,64], the observed trend in catalytic activity was proposed to be directly related to the amount of easily reducible cobalt species in each sample, which could be measured by the specific oxygen consumption at low temperatures in the CH$_4$-TPR profiles. In this sense, Figure 8 shows that there was a markedly good correlation between the $T_{50}$ values and the reacted O$_2$ below 550 $\degree C$. Accordingly, a comparable relationship was evidenced in relation with the H$_2$ uptake involved in the reduction of the free Co$_3$O$_4$ present in each supported cobalt catalyst (H$_2$-TPR profiles).

The integral method was followed for evaluating the apparent activation energy of the reaction over the examined cobalt catalysts. A power law kinetic equation, derived from a simplified Mars–van Krevelen reaction mechanism in excess of oxygen, was used. Hence, methane was assumed to follow first pseudo-order kinetics while for oxygen a zero pseudo-order was used [65]. The results are listed in Table 3 while the corresponding plots for this linearized kinetic equation are shown in Figure 9. In addition, the activation enthalpy and entropy (Table 3) were estimated by applying the Eyring-Polanyi equation. The corresponding linearized plots are shown in Figure S2 (Supplementary Materials). The following apparent activation energies were estimated, namely 82 kJ mol$^{-1}$ over Co/CeO$_2$, 90 kJ mol$^{-1}$ over Co/Al$_2$O$_3$ and 102 kJ mol$^{-1}$ over Co/MgO. When compared with the value obtained by the bulk Co$_3$O$_4$ (78 kJ mol$^{-1}$) a close similarity was found in relation to the Co/CeO$_2$.
catalyst. This finding was coherent with the fact that in both cases the nature of the active cobalt phase was the same, namely, Co$_3$O$_4$. A noticeable higher activation energy was noticed for the samples in which a mixture of cobalt phases was present, namely Co$_3$O$_4$/CoAl$_2$O$_4$ (Co/Al$_2$O$_3$ catalyst) and Co$_3$O$_4$/Co-Mg mixed oxide (Co/MgO catalyst). This behavior could lie in the contribution of these intrinsically less active phases to the reaction mechanism, especially in the case of the Co/MgO given its different activation entropy, thereby negatively influencing the overall activity of the resultant catalyst [57,66].

![Figure 8](image_url)  
**Figure 8.** Relationship between the activity and the redox properties of the supported catalysts.

![Figure 9](image_url)  
**Figure 9.** Order integral fit for the experimental kinetic data obtained over the supported cobalt catalysts.

In this way, it could be established that the presence of CoAl$_2$O$_4$ would negatively affect the intrinsic activity of the Co/Al$_2$O$_3$ samples while the formation of a stable Co–Mg solid solution would negatively impact on the kinetic behavior of the Co/MgO catalyst.
Finally, the thermal and hydrothermal stability of the most active catalyst (Co/CeO$_2$) was studied over relatively prolonged periods of operation under both dry and humid conditions at 300 cm$^3$ CH$_4$ g$^{-1}$ h$^{-1}$ (30,000 cm$^3$ g$^{-1}$ h$^{-1}$, about 60,000 h$^{-1}$). A 1% CH$_4$/10% O$_2$/10% H$_2$O/79% N$_2$ was used for the time intervals carried out in the presence of water. The dry and humid conditions where switched every 25 h over a total operation span of 150 h at a constant temperature of 550 °C. The results are shown in Figure 10. The catalyst underwent some fast thermal deactivation during the first 25 h of the test where the conversion dropped from 70% to 60%. The introduction of water to the feed stream had a significantly detrimental effect over the activity of the catalyst, and as a result conversion levels fell down to about 35%. However, when the dry conditions were re-established, the conversion recovered almost completely to the levels exhibited before the addition of water. This behavior was again seen during the following dry-humid operation cycles, and suggested that water inhibition was essentially caused by the coverage of the surface that limited the extent of the reaction methane and catalyst oxygen active species. This effect has been also observed in other works, and has been generally been linked to the weak adsorption potential that water molecules have on the surface of cobalt oxide [67,68]. In particular, Geng et al. [69] observed a significant increase in the intensity of the DRIFTS absorption bands from hydroxyl groups in a Co/γ-Al$_2$O$_3$ catalyst when water vapor was added to the feed. Additionally, H$_2$O-TPD experiments in that work also proved that a high fraction of adsorbed water on the Co catalyst surface could be desorbed at low temperatures (<500 °C).

![Figure 10. Stability of the Co/CeO$_2$ catalyst under cycled dry/humid conditions with time on stream.](image)

### 4. Conclusions

Cobalt catalysts supported over gamma-alumina, magnesia and ceria were synthesized, characterized and examined for the oxidation of methane under lean conditions with the objective of determining the influence of the catalytic support on the properties and activity of the active Co$_3$O$_4$ phase.

The analysis of the samples evidenced a marked effect of the strong cobalt-support interaction on the catalytic performance. More specifically, when the selected support was γ-alumina, the high cobalt dispersion and strong cobalt-support interaction resulted in the partial fixation of deposited
cobalt as poorly reducible cobalt aluminate (CoAl₂O₄) species. This led to a low activity despite the good structural and textural properties of the resultant catalyst. In the case of the Co/MgO sample, the cobalt-support interaction provoked the partial dissolution and insertion of cobalt cations into the MgO lattice with the subsequent formation of a highly stable Co–Mg solid solution. However, this catalyst was somewhat more active than the alumina counterpart, due to a higher oxygen mobility of the remaining free Co₃O₄ phase. Lastly, when the chosen support was ceria, the resultant catalyst exhibited worse structural and textural properties, but interestingly a noticeably promoted reducibility and oxygen mobility caused by the partial insertion of Ce cations into the cobalt spinel lattice. These beneficial effects made this catalyst the most active sample among the examined cobalt catalysts. Additionally, while its thermal stability over a prolonged time interval was found to be high, the addition of water vapor to the feedstream provoked a reversible inhibition assigned to the coverage of the catalyst surface by water molecules.

Given the good activity showed by the Co/CoO₂ catalyst in spite of its poor textural and structural properties, future efforts will be focused on designing Co/CoO₂ with improved specific surface area and crystallite size, either by modification of the synthesis method of the ceria support or by selecting ceria as a promoter for Co/CoO₂–Al₂O₃ catalysts.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/1996-1444/12/19/3174/s1.


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**Conflicts of Interest:** The authors declare no conflict of interest.

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