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In situ investigation of the CO₂ methanation on carbon/ceria-supported Ni catalysts using modulation-excitation DRIFTS



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ABSTRACT

The development of novel cost-efficient, high-performing catalysts for CO_2 methanation that are active at low temperatures can be optimized through the understanding of the reaction mechanism on different materials. A series of Ni-based catalysts supported on CeO_2 and carbon/ CeO_2 composites was investigated, showing that Ni nanoparticles supported on a carbon/ CeO_2 composite with a 50:50 wt ratio and on pure CeO_2 have excellent low-temperature activity and achieve up to 87% CO_2 conversion with full selectivity towards CH_4 at 370 °C. Importantly, meaningful insights on the reaction mechanism were gathered for the different types of materials by using the emerging ME–PSD–DRIFTS technique. The study of the rate of formation/consumption of the various intermediates showed that the CO_2 methanation reaction follows a combination of the CO and formate pathways in the case of Ni on pure CeO_2 ; however, in the case of Ni on the carbon/ CeO_2 composite, it follows only the formate pathway.

1. Introduction

The CO₂ methanation reaction (CO₂ + 4 H₂ \rightarrow CH₄ + 2 H₂O, $\Delta H_{298 \text{ K}}$ = -165 kJ mol⁻¹) is the most thermodynamically favorable reaction of CO₂ hydrogenation, and is a promising way to convert anthropogenic CO₂ back to CH₄ fuel [1–4]. The hydrogenation proceeds through an eight-electron reduction process with consequent significant kinetic barriers, thus requiring the action of a catalyst. From the mechanistic perspective, extensive research has been conducted on CO₂ methanation. Nevertheless, the reaction mechanism is still vigorously debated, since it depends on the catalytic material and the hydrogenation conditions [1,3]. The most predominant mechanisms proposed in the literature can be grouped in two different categories: (i) the dissociative pathway, where CO₂ is dissociated into CO, which acts as the main intermediate, and (ii) the associative pathway, where CO₂ is molecularly adsorbed associatively with a H atom previously adsorbed on the surface, forming formate as an intermediate, i.e., without CO formation [3].

 CO_2 methanation has played a significant role in understanding CO_2 hydrogenation and the associated development of catalysts. Importantly, researchers continue to exploit different catalysts to achieve high CO_2 conversion and complete selectivity towards CH_4 . For example, a number of catalysts based on transition metals (Ni, Co, Fe) or platinum-group metals (Ru, Rh) dispersed on porous supporting materials have been developed [2,5–12]. Notably, Ni-based catalysts, compared to platinum-group metals, exhibit an optimal combination of activity, selectivity, and low price [2,5,6]. Despite the advances in using Ni catalysts for CO_2 methanation, a way to reduce the reaction temperature from the current ≥ 400 °C while preserving high catalytic performance remains a challenge.

One interesting approach to address this challenge is to develop Ni

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catalysts supported on novel materials. Traditionally, alumina (Al_2O_3) is used as supporting material for Ni methanation catalysts [13–15]. In contrast, we and other researchers have focused on carbon supporting materials [2,5,16,17]. An exciting feature of these high-surface-area porous supports are their versatile surface properties [18]. Specifically, via modification of their surface chemistry, carbon materials can exhibit increased basicity, which improves the CO₂ coverage of the catalyst, since it is an acidic molecule, thus enhancing the CO₂ methanation performance at lower reaction temperatures [2]. Compared to the more common Ni supported on metal oxides, the research on Ni supported on carbon materials is less advanced, motivating a fundamental investigation of such alternative catalytic systems [2,5].

Among the metal-oxide supporting materials, despite being based on a rare-earth metal, ceria (CeO₂) has emerged as a promising alternative to Al₂O₃ because it enhances the metal–support interactions, leading to high dispersion of the resultant Ni nanoparticles over the support. Moreover, owing to the enhanced surface coverage by CO₂ species, enabled by the CeO₂ surface basicity, it improves the low-temperature catalytic performance of the catalysts in CO₂ methanation [19–26]. These positive effects of CeO₂ motivated us to develop high-performing carbon/ceria-supported Ni catalysts for CO₂ methanation, as well as to conduct a comparative mechanistic study of CO₂ hydrogenation over these catalysts.

In this work, we report the successful precipitation synthesis of a composite supporting material consisting of reduced activated carbon (ACR) and CeO₂ (ACR–CeO₂) as well as pure CeO₂, followed by deposition of Ni nanoparticles via the incipient wetness impregnation route. An important feature of our findings is the remarkable reduction by 80 °C of the optimal temperature of the CO₂ methanation reaction, compared to that for a CeO₂-free Ni/ACR catalyst developed in our previous study [2], thus significantly improving the sustainability of the methanation approach without jeopardizing the catalytic performance. Furthermore, we conducted a detailed *in situ* modulation-excitation phase-sensitive-detection diffuse-reflectance infrared Fourier-transform spectroscopy (ME–PSD–DRIFTS) investigation of the methanation over the newly synthesized high-performing Ni/ACR–CeO₂ and Ni/CeO₂ catalysts, to obtain key mechanistic insights.

2. Experimental methods

2.1. Synthesis

In the present work, we synthesized reduced activated carbon (ACR) with Lewis base sites by heat-treatment of a commercial NORIT GAC 1240 PLUS activated carbon (CABOT) at 900 °C for 1 h under H₂ flow [2]. Pure CeO₂ and composite ACR–CeO₂ supporting materials were prepared by a precipitation method adapted from elsewhere [27]. Briefly, Ce(NO₃)₃·6H₂O is precipitated by NaOH under vigorous stirring, along with the addition of calculated amounts of ACR for the composite ACR–CeO₂ synthesis. The 15% Ni nanoparticles were deposited onto as-synthesized supporting materials via incipient wetness impregnation using Ni(NO₃)₃·6H₂O, followed by heat treatment to decompose nitrate, thus generating metallic Ni (Ni/CeO₂ and Ni/ACR–CeO₂).

2.2. Characterization

The synthesized materials were characterized by N_2 physisorption, powder X-ray diffraction (XRD), H_2 temperature programmed reduction (H_2 -TPR), H_2 -chemisorption, X-ray photoelectron spectroscopy (XPS), transmission electron microscopy (TEM), energy-dispersive X-ray spectroscopy in scanning TEM mode (STEM–EDX).

2.3. Catalytic testing

The activity, selectivity, and stability of the synthesized catalysts in CO_2 methanation were evaluated in a Microactivity XS15 system (PID

Eng & Tech), using a fixed bed quartz reactor ($\emptyset_{int} = 1$ cm). Outflow products were analyzed online using a GC 1000 gas chromatograph (DANI), equipped with a thermal conductivity detector (TCD) and a GS-CarbonPLOT capillary column, while using He as the carrier gas and a continuous flow of N₂ as the internal standard. Each catalyst (100 mg) was diluted by inert SiC material, and then was pre-treated in situ under H_2 flow (40 cm³ min⁻¹) at its reduction temperature for 30 min, at 1 bar. After that, the reactor temperature was decreased to 100 °C under He (50 cm³ min⁻¹), and then, the reactor was fed with 10% CO₂, 40% H_2 (the stoichiometric composition) and 50% He, with a total flowrate of 100 cm³ min⁻¹. A temperature ramping from 100 to 500 °C at a rate of 5 $^{\circ}$ C min⁻¹ was performed under reaction conditions to evaluate the catalytic properties of the catalysts at various temperatures. Additionally, isothermal time-on-stream (TOS) experiments were conducted under the same reaction conditions to evaluate the stability of the catalysts.

2.4. In situ spectroscopy

 CO_2 adsorption and CO_2 methanation over the synthesized materials were probed using steady-state *in situ* diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) using a Nicolet iS50 FTIR spectrometer (Thermo Scientific), equipped with a Mercury-Cadmium-Telluride (MCT) detector and a high temperature reaction chamber (Harrick Scientific). Additionally, the CO_2 methanation over the prepared materials was investigated by *in situ* modulation excitation phase-sensitive detection DRIFTS (ME–PSD–DRIFTS) using the same spectrometer. The ME-PSD analysis highlight the species that change with the modulation frequency (*f*). The collected spectra were converted into the socalled frequency domain by Fast Fourier Transform (FFT). The data was then filtered at the modulation frequency and converted into the phase-domain by Inverse Fast Fourier Transform (IFFT). Activated carbon absorbs most of the IR radiation, so it was not possible to perform DRIFTS analyses in the Ni/ACR sample.

Readers interested in the detailed experimental information are referred to the Supplementary material (SM).

3. Results

3.1. CO2 methanation over Ni/CeO2 and Ni/ACR-CeO2

In our previous study [2], a novel methanation catalyst with 15 wt% Ni loading over ACR having O-free Lewis basic sites (denoted as **Ni/ACR**) has been developed, exhibiting CO₂ conversion (X_{CO2}) of 76% and selectivity towards CH₄ (S_{CH4}) of 97%, but at a fairly high reaction temperature of 450 °C. To increase the conversion while lowering the reaction temperature, we sought to introduce CeO₂, thus forming a composite supporting material with ACR. We began by optimizing the CeO₂ loading in the ACR–CeO₂ composite support. It was established that 15 wt% Ni catalyst supported on ACR–CeO₂ with ACR:CeO₂ weight ratio of 50:50 exhibits the optimal catalytic properties in CO₂ methanation (Fig. S1). Hence, we further conducted our studies with this catalyst, hereafter referred to as **Ni/ACR–CeO₂**.

With the optimized composite catalyst in hand, we investigated its methanation performance and compared it to a control 15 wt% Ni catalyst supported on pure CeO₂ (denoted as Ni/CeO₂).

Fig. 1a,b show the obtained X_{CO2} and S_{CH4} results in the temperature range of 200–500 °C. Both catalysts start to convert CO₂ from ca. 200 °C reaching $X_{CO2} = 50\%$ at 280 °C, the so-called T_{50} .

Upon the temperature increase, the catalysts further achieve CO₂ conversions that are very close to the thermodynamic equilibrium, exhibiting maximum X_{CO2} of 85.7% (Ni/CeO₂) and 87.4% (Ni/ACR–CeO₂) at a notably low temperature of only 370 °C (Fig. 1a). In terms of S_{CH4} , the catalysts also demonstrate values comparable to the thermodynamic equilibrium over the range of the studied temperatures, with $S_{CH4} \approx 100\%$ at maximum X_{CO2} (Fig. 1a,b), with the only

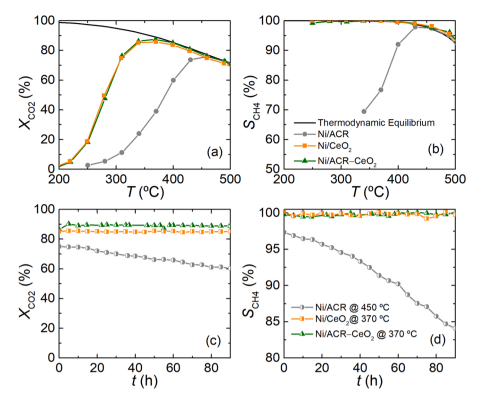


Fig. 1. Comparison of X_{CO2} (a) and S_{CH4} (b) as a function of reaction temperature, together with the stability testing results in terms of X_{CO2} (c) and S_{CH4} (d) over 90 h TOS of the new Ni/CeO₂ and Ni/ACR–CeO₂ catalysts in comparison with previously reported ceria-free Ni/ACR [2]. The thermodynamic equilibrium curves were calculated using a model based on the minimization of Gibbs free-energy of the existing species. Methanation conditions: P = 1 bar; *WHSV* = 60 000 cm³ g⁻¹ h⁻¹; CO₂: H₂ (V:V) = 1: 4.

byproduct being CO. We finally investigated the stability of the catalysts through a long time-on-stream (TOS) testing at the best performing temperature of 370 °C. It was established that both the control **Ni/CeO₂** and composite **Ni/ACR–CeO₂** demonstrate excellent stability over 90 h on stream with no loss in either X_{CO2} or S_{CH4} (Fig. 1c,d). Importantly, a comparison of the catalytic properties between the new carbon/ceriacontaining **Ni/ACR–CeO₂** and our previously reported carboncontaining **Ni/ACR** [2] (Fig. 1) evidently highlights that the compositing of ACR with CeO₂ not only leads to a significant improvement in the methanation performance but also simultaneously lowers the optimal reaction temperature by 80 °C (from 450 °C for **Ni/ACR** to 370 °C for **Ni/ACR–CeO₂**). Moreover, the compositing of ACR with CeO₂ leads to an improvement in stability at the respective optimal temperature of the resultant **Ni/ACR–CeO₂** catalyst, relative to that of **Ni/ACR** [2] (Fig. 1).

3.2. Characterization of the catalysts

Our catalytic testing reveals that both Ni/CeO_2 and $Ni/ACR-CeO_2$ exhibit similarly excellent catalytic properties at reasonably low reaction temperatures, despite being based on different supporting materials. Therefore, we characterized the catalysts to determine the parameters that govern the methanation performance.

Even though Ni/CeO2 and Ni/ACR-CeO2 have similar catalytic properties (Fig. 1), the data presented in Table 1 demonstrate that the catalysts exhibit quite different textural properties. The textural properties of the synthesized materials were estimated using Brunauer-Emmett-Teller (BET) and Barrett-Joyner-Halenda (BJH) methods. Pure CeO₂ support has a relatively low specific surface area (S_{BET}) of 104 m² g⁻¹, while the pure ACR support has about 7-fold higher $S_{\text{BET}} = 784 \text{ m}^2 \text{ g}^{-1}$. As expected, the composite ACR–CeO₂ exhibits a $S_{\text{BET}} = 373 \text{ m}^2 \text{ g}^{-1}$ that is higher than that of pure CeO₂, but lower than that of pure ACR. This trend is sustained for the respective Ni catalysts supported on the two materials, exhibiting a decreasing order in S_{BET} Ni/ACR-CeO₂ > Ni/CeO₂ (Table 1). The addition of Ni leads to a slight decrease in S_{BET} compared to the respective supporting materials (Table 1). Both CeO₂ and ACR-CeO₂ supports, and the derivative catalysts present Type IV isotherms (Fig. S2); the pore-size-distribution data (Fig. S3) reveal that Ni/CeO₂ exhibits a higher number of pores having larger width, while Ni/ACR-CeO₂ has a peak in the smaller width range, also containing pores with diameters higher than 5 nm.

Next, we probed the metal–support interactions and metallic surface areas of Ni/CeO_2 and $Ni/ACR-CeO_2$. The comparison of hydrogen temperature-programmed reduction (H₂-TPR) profiles of Ni/CeO_2 and

Table 1

Textural properties [specific surface area (S_{BET}), and total pore volume (V_p)], average size of CeO₂ crystallites ($d_{c,CeO2}$), and Ni crystallites ($d_{c,Ni}$), metallic Ni surface area (S_{M}), temperature at which $X_{CO2} = 50\%$ (T_{50}), and maximum X_{CO2} , attained at 450 °C for Ni/ACR [2] and 370 °C for Ni/ACR–CeO₂.

	$S_{\rm BET} ({ m m}^2~{ m g}^{-1})^{ m a}$	$V_{\rm P} ({\rm cm}^3 {\rm g}^{-1})^{\rm a}$	$d_{\rm c,Ni} (\rm nm)^{\rm b}$	$d_{\rm c,C} e_{\rm O2} (\rm nm)^{\rm b}$	$S_{\rm M}$ (m ² g _{cat} ⁻¹) ^c	T ₅₀ (°C)	Max X_{CO2} (%)
ACR [2]	784	0.48	-	_	_	_	0
CeO ₂	104	0.17	_	7	-	_	12
ACR-CeO ₂	373	0.33	_	6	_	-	11
Ni/ACR [2]	656	0.41	8	-	2.73	385	76
Ni/CeO ₂	82	0.14	9	7	1.66	280	86
Ni/ACR-CeO2	386	0.30	5	5	2.57	280	87

^a Estimated by N₂ physisorption.

^b Estimated from XRD data using Scherrer equation.

 $^{\rm c}\,$ Estimated by H_2 pulse chemisorption.

Ni/ACR-CeO₂ demonstrate different metal-support interactions between Ni nanoparticles and the supporting materials (Fig. S4a). Specifically, Ni/CeO2 exhibits a large peak at 360 °C with a shoulder at 404 °C, as well as two small shoulders at 334 °C and 464 °C. In contrast, Ni/ACR-CeO2 exhibits a large peak at a lower temperature of 320 °C with two high-temperature shoulders at 392 °C and 502 °C. Apparently, Ni/CeO2 is largely composed of Ni species that exhibit a stronger interaction with pure CeO2 support, compared to that in composite Ni/ ACR-CeO₂. Regarding the Ni surface area ($S_{\rm M}$), measured using H₂ pulse chemisorption, it was estimated that Ni/ACR-CeO2 exhibits more than 1.5 times higher metallic Ni surface than does Ni/CeO₂ (Table 1). The H₂-TPR of the supporting materials was also assessed (Fig. S4b) and it was observed that the H₂ consumed for CeO₂ reduction is minimal when comparing to the H₂ consumed for the reduction of the Ni-based catalysts. Additionally, the analysis of the H2 - TPR profiles indicate that the ACR-CeO₂ support is a homogeneous mixture that interacts in a way that the reduction profile of the Ni/ACR-CeO2 sample is different than that of the other two supports (CeO₂ and ACR), and not a combination of both Ni/CeO2 and Ni/ACR [2] H2-TPR profiles.

To gain structural insight into the synthesized materials, we further investigated their phase composition using powder X-ray diffraction (XRD). According to the XRD patterns (Fig. S5), the as-synthesized pure and composite supports contain cubic CeO₂ phase (ICDD #04–006–1932). In contrast, the respective Ni catalysts additionally contain cubic metallic Ni (ICDD #04–016–4761), with no secondary phases being present. There is possibly a graphitic carbon peak, but since it should be present at 28°, overlapping with a peak from CeO₂, it is impossible to access its presence. The average size of Ni and CeO₂ crystallites was assessed from XRD data using the Scherrer equation (Table 1), and the size of Ni crystallites in Ni/CeO₂ (ca. 9 nm) was estimated to be nearly 2 times larger than in Ni/ACR–CeO₂ (ca. 5 nm), which is consistent with the smaller $S_{\rm M}$ of Ni/CeO₂.

X-ray photoelectron spectroscopy (XPS) analysis revealed that both catalysts, Ni/CeO₂ and Ni/ACR–CeO₂, exhibited two different Ni phases, observed from the deconvolution of the XPS data for Ni 2p_{3/2} (Fig. 2). The lower energy component (853.0 ± 0.1 eV) is assigned to Ni⁰ in metallic Ni species, while the components at 854.4 ± 0.2 eV and 856.4 ± 0.1 eV are consistent with Ni²⁺ in nickel-oxide phase, and the ones at 861.3 ± 0.3 eV and 864.3 ± 0.1 eV are satellite peaks from Ni²⁺. The Ni 2p_{3/2} spectral envelopes of the catalysts resemble the typical spectra for nanoscale oxidation layers on Ni and Ni compounds, being consistent with a thin, predominantly Ni²⁺, nickel oxide layer on metallic Ni [5,28–30]. The binding energies (BEs) of fitting components are presented in Table S1 and the relative atomic percentages (at%) of Ni⁰ and Ni²⁺ are presented in Table S2. The relative atomic percentage

of Ni⁰ is similar in Ni/ACR–CeO₂ and Ni/CeO₂ catalysts, at 4.2 at% and 4.4 at%, respectively (Table S2).

The Ce 3d spectra also revealed the presence of mixed Ce phases (Fig. 2) in both samples. The spectra can be represented as five pairs of doublets, identified in Fig. 2 as u and v for the Ce $3d_{3/2}$ and Ce $3d_{5/2}$ spin–orbit components, with spin–orbit splitting of 18.4 ± 0.5 eV for all the samples, consistent with those found in the literature [31,32]. These spectra were fitted using a method similar to that reported by Zhang et al. [32] where the u''' peak is fixed and the remaining peaks are constrained according to the u''' position.

The v^0 , u^0 , v', and u' are attributed to Ce^{3+} , while the highest biding energy peaks, v''' and u''', are attributed to the primary components of Ce^{4+} , and the remaining v, u, v'', and u'', are identified as "shake-down" states, as reported by Mullins et al. [31,33] Table S2 presents the BEs of each of these components. Importantly, we note that there is a small overlap of the Ni 2p_{1/2} components (orange lines and symbols below 881 eV in Fig. 2) and the low-BE components in the Ce 3d spectra. This overlap was accounted for by including in the Ce 3d fitting procedure Ni 2p_{1/2} components (orange lines in Fig. 2), with fixed parameters based on the more reliable fits in the Ni 2p_{3/2} region.

The Ce 3d peaks in Ni/ACR–CeO₂ are noticeably narrower than those in Ni/CeO₂, which likely indicates that ceria nanoparticles are in a more uniform electrostatic environment in Ni/ACR–CeO₂, where they are in a good electrical contact with the carbon substrate.

The amount of Ce^{4+} and reduced Ce^{3+} is quantified in Table S2. Interestingly, the Ni/ACR–CeO₂ catalyst exhibits a higher relative at% of Ce^{3+} (20.5 at%) compared to Ni/ACR–CeO₂ that contains 16.4 at% Ce^{3+} .

The C 1s spectra of Ni/ACR–CeO₂ were also analyzed, and the results are presented in Fig. S6 [35,36].

Interested in understanding the microstructural properties of the catalysts, we then conducted transmission electron microscopy (TEM) and energy-dispersive X-ray spectroscopy in scanning STEM mode (STEM–EDX) analyses. These studies reveal metallic Ni nanoparticles well-dispersed over nanocrystalline CeO₂ and amorphous-like ACR. Fig. 3a shows TEM image of Ni nanoparticles in Ni/CeO₂, where it is possible to distinguish only some large Ni particles (ca. 30–50 nm) dispersed over CeO₂ support. Fig. 3e shows Ni nanoparticles dispersed on composite ACR–CeO₂ support in Ni/ACR–CeO₂, wherein the Ni particles (ca. 5–10 nm) dispersed on ACR and some larger Ni particles (ca. 30–50 nm) on CeO₂ are observed. It is important to note that the imaging contrast between the Ni and CeO₂ nanoparticles is very low, thus it is difficult to differentiate the two types of nanoparticles in the electron microscopy images. Therefore, the size ranges indicated are merely representative of the observed identifiable nanoparticles, leading

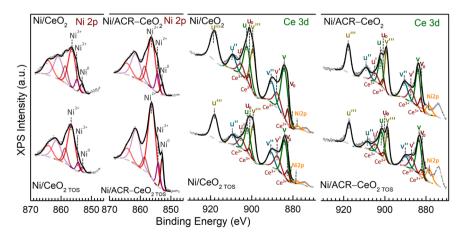


Fig. 2. High-resolution XPS data for the Ni 2p_{3/2} and Ce 3d region collected from as-synthesized and TOS-tested Ni/CeO₂ and Ni/ACR–CeO₂. Symbols: raw data; black lines: overall fits; colored lines: fits of individual components; dashed lines: background. The observed binding energies (BE) and assignment of the peak positions are summarized in Table S1.

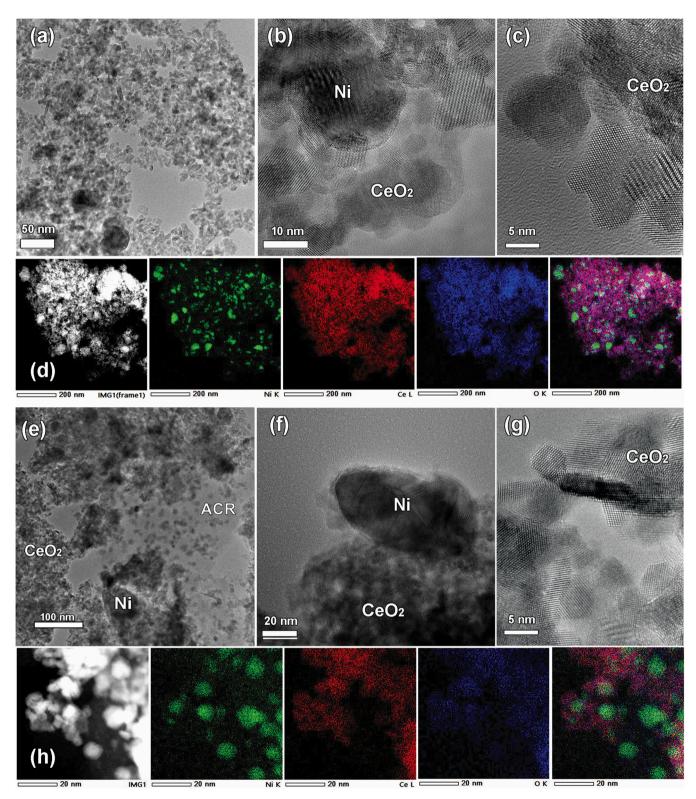


Fig. 3. TEM images of Ni nanoparticles dispersed on CeO_2 (a) and ACR– CeO_2 (e) supporting materials, together with the representative HRTEM images of Ni and CeO_2 nanoparticles in the Ni/ CeO_2 (b,c) and Ni/ACR– CeO_2 (f,g) catalysts. Low-magnification STEM images of the Ni/ CeO_2 (d) and Ni/ACR– CeO_2 (h) catalysts, together with the respective STEM–EDX maps for Ni, Ce, O and their mixture.

to a gap between the Ni nanoparticles sizes from TEM and those estimated by XRD, which is more representative.

Although it is difficult to distinguish the smaller Ni particles on the CeO_2 support due to the low contrast on the TEM images, the respective STEM–EDX maps evidence that there are also small Ni particles

dispersed on CeO₂ material in both Ni/CeO₂ and Ni/ACR–CeO₂ catalysts (Fig. 3d,h). A closer look at CeO₂ supporting material in Ni/CeO₂ (Fig. 3c) and Ni/ACR–CeO₂ (Fig. 3g) catalysts reveals that the ceria consists of agglomerated CeO₂ nanoparticles (ca. 5–10 nm), preferentially faceted by the {110} plane.

Interestingly, high-resolution TEM (HRTEM) imaging of Ni nanoparticles in Ni/CeO₂ and Ni/ACR-CeO₂ reveals that the particles in both catalysts exhibit Ni@NiO core-shell appearance. As an example, Fig. 4a,b show HRTEM images of a Ni nanoparticle in Ni/ACR-CeO₂ catalyst, together with the corresponding Fourier transform (FT) patterns for the Ni core and NiO shell. Furthermore, multiple twinning of the Ni nanoparticle can be observed in the HRTEM images, marked by white arrow heads.

Even though there was no loss in X_{CO2} and S_{CH4} over 90 h of TOS testing (Fig. 1c,d), we decided to investigate the impact of the reaction conditions on the microstructure and chemical composition of the catalysts by means of electron microscopy and XPS.

The Ni $2p_{3/2}$ spectra (Fig. 2) of the samples after TOS testing (Ni/ CeO_{2 TOS} and Ni/ACR–CeO_{2 TOS}) include fitting components at BEs similar to those of the components in the spectra of fresh Ni/CeO₂ and Ni/ACR–CeO₂ catalysts. Considering the relative Ni⁰ atomic content (Table S2), however, we observed that it decreases to 2.8 at% (cf. 4.4%) in the case of Ni/CeO_{2 TOS} and, on the contrary, increases to 7.4 at% (cf. 4.2 at%) in the case of Ni/ACR–CeO_{2 TOS}. In the Ce 3d spectra (Fig. 2), again the positions of the peaks remain similar for both catalysts after 90 h of TOS, but there are differences in the Ce³⁺/Ce⁴⁺ ratio before and after TOS in both Ni/CeO₂ and Ni/ACR–CeO₂. The Ce³⁺ atomic content in Ni/CeO₂ increases to 21.3 at% (cf. 16.4 at%), while in Ni/ ACR–CeO₂, it remains approximately constant with 19.9 at% (cf. 20.5 at%).

From Figs. S8a-d, one can see that the TOS-tested **Ni/CeO**₂ maintains the microstructure of the Ni nanoparticles dispersed over the ceria support composed by small CeO₂ nanoparticles (ca. 5–10 nm), similar to that in pristine **Ni/CeO**₂ (Fig. 3a). In contrast, TOS-tested **Ni/ACR–CeO**₂ undergoes significant microstructural changes within CeO₂ support. Specifically, Fig. S8e evidences the appearance of CeO₂ particles with a needle-like shape after 90 h of methanation. Such nanoneedles were not observed in pristine **Ni/ACR–CeO**₂ (Fig. 3e) or pristine and TOS-tested **Ni/CeO**₂ (Fig. 3a, S8a).

Notably, a closer look at Ni particles in TOS-tested Ni/CeO_2 (Fig. S8c) and $Ni/ACR-CeO_2$ (Fig. 4c,d) indicates a preservation of Ni@NiO core-shell nanostructure of the catalyst after prolonged methanation.

3.3. Probing the mechanism of CO_2 methanation by in situ DRIFTS

Having in hand two well-characterized high-performing methanation catalysts, we further conducted *in situ* DRIFTS studies to elucidate the reaction mechanism over Ni/CeO₂ and Ni/ACR–CeO₂. This technique is used to understand the reaction mechanism in heterogeneous catalysis, probing the adsorbed species on the catalysts under reaction conditions. There are some drawbacks of this technique. Specifically, it detects not only the intermediate species but also the sample background, noise, and spectator species, making it difficult to distinguish what are the real intermediates of the reaction [37,38].

Furthermore, the intermediate species have a short lifetime, making it even more difficult to differentiate them from the remaining species, which are present under steady-state conditions. Fortunately, the modulation-excitation (ME) technique offers an interesting solution for overcoming the aforementioned drawbacks. It is based on the introduction of a rapid periodic perturbation on the testing system, to which only the intermediate species of the reaction respond. The resultant spectroscopic signal is then analyzed by phase-sensitive detection (PSD) technique to enhance the signal-to-noise ratio and to extract the information about intermediate species that responded to the modulation signal, thus avoiding the signal from the spectator species [10,37–41].

We started by performing steady-state in situ DRIFTS for CO2 adsorption and then for CO2 methanation at 250, 300, and 350 °C. In this temperature range, the X_{CO2} goes from low to almost complete. Fig. S9 shows the full range spectra for each experiment, along with a discussion of the observed trends for hydroxyl groups, gaseous CO2, and gaseous CH₄ [42]. Fig. 5 displays a zoom-in of the region below 2000 cm⁻¹, which carries important information about the CO₂ adsorption and methanation mechanisms, while Table 2 summarizes the respective band assignment. During CO_2 adsorption (Fig. 5a,c), two bands detected at 1500 and 1370 cm⁻¹ (ν_{as} / ν_{s}) can be attributed to formate (HCOO) species, which are probably close to a reduced Ce site in CeO₂, because they have a low $\Delta_{as-s} = 130 / 150 \text{ cm}^{-1}$ [10,26]. Two other bands are observed at 1600 and 1280 cm⁻¹ (ν_{as} / ν_{s}), that can be assigned to bi-dentate carbonates (b-CO₃), along with the band at 1030 cm^{-1} [10,26]. Moreover, a small band at 1415 cm⁻¹ is also observed in the spectrum of Ni/CeO2 at 250 °C, which can be attributed

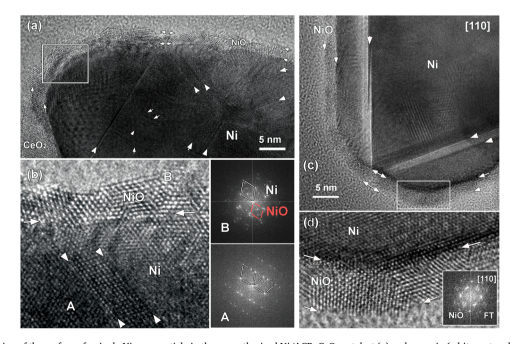


Fig. 4. HRTEM imaging of the surface of a single Ni nanoparticle in the as-synthesized Ni/ACR–CeO₂ catalyst (a) and zoom-in (white rectangle) image (b) and in the same catalyst after its TOS testing for 90 h (c,d), revealing the core-shell appearance of the nanoparticles. The insets in (b,d) show the FT patterns from the corresponding regions. The white arrows mark the NiO shell and the white arrow heads depict the twinning within the Ni nanoparticles.

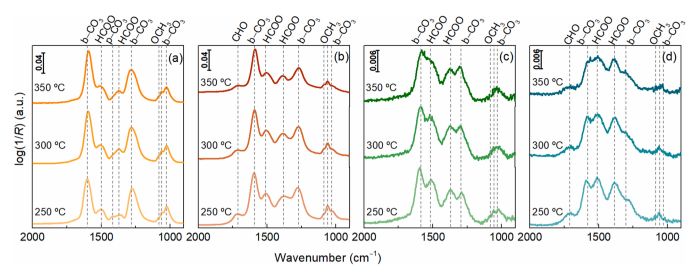


Fig. 5. Steady-state *in situ* DRIFTS studies of CO₂ adsorption and methanation over Ni/CeO₂ (a,b) and Ni/ACR–CeO₂ (c,d) catalysts. Experimental conditions for CO₂ adsorption: P = 1 bar; CO₂ flowrate of 2.5 cm³ min⁻¹; N₂ flowrate of 100 cm³ min⁻¹; $m_{cat} = 15$ mg. Experimental conditions of CO₂ methanation: P = 1 bar; CO₂ flowrate of 2.5 cm³ min⁻¹; N₂ flowrate of 100 cm³ min⁻¹; $m_{cat} = 15$ mg. Experimental conditions of CO₂ methanation: P = 1 bar; CO₂ flowrate of 2.5 cm³ min⁻¹; N₂ flowrate of 100 cm³ min⁻¹; $m_{cat} = 15$ mg. The dashed lines indicate the positions of the observed peaks.

Table 2

Assignment of the bands observed during steady-state DRIFTS and ME–PSD–DRIFTS experiments: hydroxyl groups (OH⁻), gaseous methane (CH₄), gaseous carbon dioxide (CO₂), bridged carbon monoxide (br–CO), formyl groups (CHO), bi-dentate carbonates (b–CO₃), poly-dentate carbonates (p – CO₃), formates (HCOO), carboxylate ($CO_2^{-\delta}$), and methoxy groups (OCH₃).

	Steady-state in situ I	ORIFTS	In situ ME–PSD–DRIFTS CO ₂ adsorption							
	CO ₂ adsorption					CO ₂ methanation				
	Ni/CeO ₂	Ni/ACR-CeO2	Ni/CeO ₂	Ni/ACR-CeO2	Ni/CeO ₂	Ni/ACR-CeO2				
	Wavenumber (cm $^{-1}$)									
CO ₂ / OH ⁻	3750 - 3550	3750 - 3550	3750 - 3550	3750 - 3550	3750 - 3550	3750 - 3550				
CH ₄	_	_	_	3017	3017	3017				
CO_2	2400 - 2250	2400 - 2250	2400 - 2250	2400 - 2250	2400 - 2250	2400 - 2250				
CO gas	_	_	-	2200 - 2100	2200 - 2100	2200 - 2100				
br-CO	_	_	-	_	1760	1770				
CHO	_	_	1710	1710	1700	1700				
b-CO3	1600	1585	1590	1580	1610	1600				
	1280	1295	1270	1300	1290	1290				
	1030	1030	1030	1030	1020	1020				
$p - CO_3$	1415	_	-	_	1450	1450				
	_	_	-	_	1405	1405				
HCOO	1500	1520	1510	1510	1520	1530				
	1370	1370	1380	1385	1360	1360				
$\text{CO}_2^{-\delta}$	_	_	-	_	1240	1240				
OCH ₃	1080	1080	1080	1080	1080	1080				
	1060	1060	1060	1060	1060	1060				

to poly-dentate (p–CO₃) carbonates Interestingly, a small amount of methoxy species (OCH₃) are present in the CO₂ adsorption spectra without H₂ being present, corresponding to the bands at 1080–1060 cm⁻¹ [24,25]; therefore, residual H₂ should be present on the Ni nanoparticles from the reduction pre-treatment which might lead to the methoxy formation during CO₂ adsorption step.

When H_2 is introduced into the gas flow to mimic the methanation conditions (Fig. 5b,d), the bands corresponding to b–CO₃ and HCOO remain at similar wavenumbers as during the CO₂ adsorption probing; however, the difference in intensity between them decreases. Furthermore, the methoxy bands increase and a new band appears at 1710 cm⁻¹ for both catalysts that can be attributed to the formation of formyl (CHO) groups [24].

To gain a deeper understanding of the reaction mechanism, we next conducted modulation-excitation phase-sensitive detection DRIFTS (ME–PSD–DRIFTS), allowing for a better signal-to-noise ratio plus distinguishing spectator and intermediate species. The modulated conditions applied in the experiments were a change in flowrate from A (CO₂ = $2.5 \text{ cm}^3 \text{ min}^{-1}$ and $10\% \text{H}_2/\text{N}_2 = 100 \text{ cm}^3 \text{ min}^{-1}$) to B (CO₂ = 2.5 cm^3

min⁻¹ and N₂ = 100 cm³ min⁻¹). Fig. 6 shows the phase-resolved spectra of the modulation excitation *in situ* DRIFTS studies conducted for Ni/CeO₂ at 250, 300, and 350 °C. A comparison of Fig. 5 and Fig. 6 show that the data from the ME-PSD-DRIFTS experiments look quite different from those of the steady state experiments. In particular, three main differences can be identified: (i) the peak detected in the steady-state spectra at 1700 cm⁻¹ (Fig. 5b) is actually caused by a convolution of 2 peaks that are out-of-phase during modulation (Fig. 6); (ii) instead of 4 peaks observed in the steady-state spectra between 1700 and 1200 cm⁻¹ (Fig. 5b), the phase-resolved spectra show 7 peaks (Fig. 6), indicating the appearance of new groups not identified in the steady-state spectra, namely, $p - CO_3$ and carboxylate (CO_2^{0-}) groups (Table 2); and (iii) two from those 7 peaks are out-of-phase with the others (Fig. 6).

With regards to the aforementioned first difference, the band at 1700 cm⁻¹ is indicative of the formyl (CHO) group, while the band at 1760 cm⁻¹ is most likely related to bridged CO (br–CO) on Ni⁰ sites [20, 24]. The appearance of br–CO on Ni⁰ should be related to two reactions occurring simultaneously: it can be formed by direct dissociation of CO_2

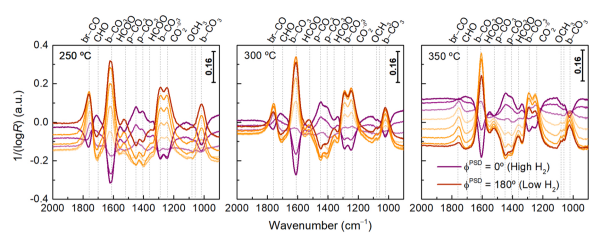


Fig. 6. Phase-resolved spectra of the modulation excitation *in situ* DRIFTS studies conducted for the Ni/CeO₂ catalyst at 250, 300 and 350 °C. Experimental conditions: P = 1 bar; flowrate A: CO₂ = 2.5 cm³ min⁻¹ and 10% H₂/N₂ = 100 cm³ min⁻¹; flowrate B: CO₂ = 2.5 cm³ min⁻¹ and N₂ = 100 cm³ min⁻¹; $m_{cat} = 15$ mg. The dashed lines indicate the positions of the peaks.

over Ni, and it can be a product of HCOO dissociation, since the presence of $\text{CO2}^{\delta-}$, b–CO₃, p–CO₃ and HCOO bands indicate that CO₂ should also be adsorbed and hydrogenated on CeO₂.

Furthermore, there is a possibility that br-CO can be hydrogenated to CHO in the presence of H₂, as previously reported [24]. This is a likely scenario since the respective bands are out of phase with each other, indicating that br-CO is hydrogenated to CHO on the surface of CeO2 in the presence of H₂. A more detailed analysis of the phase delay φ (the time to reach maximum accumulation) for both br-CO and CHO species over Ni/CeO₂ at 300 °C, show that they differ by 180° [φ (br-CO) = 108° and φ (CHO) = 288°]. This phase delay means that the CHO is formed at the same time as br-CO is consumed. Another possible scenario would be that this phase delay is dependent on that of the H₂, which would mean that H₂ favors the net formation of b-CO and disfavors the formation of CHO, leading to the maximum coverage of b-CO being observed mostly in phase with H₂, whereas CHO is mostly out of phase to H_2 . However, since the phase delays are not close to 0° or 180° , which would be close to the value where H₂ is maximum and minimum, respectively, due to low residence time, the first option is more probable: br-CO should be hydrogenated to CHO in the presence of H₂. Notably, it is not possible to present the phase delay values for the remaining intermediates due to the accumulation of multiple maxima.

More detailed analysis of the differences between the steady-state and ME–PSD data reveal that while the bands from b–CO₃ appear at the same positions (Table 2) [10,26], the set of bands associated with HCOO in the steady-state spectra (Fig. 5b) are, in fact, a convolution of different species, namely, HCOO and p–CO₃ (Fig. 6, Table 2) [10]. Notably, the later p–CO₃ assignment could also be ascribed to mono-dentate carbonates (m–CO₃), but the low Δ_{as-s} suggests the presence of p–CO₃ rather than m–CO₃ [10]. In that case, p–CO₃ appearance is out-of-phase with b–CO₃ appearance, which indicate that the coordination of the carbonates may be changing from b–CO₃ to p–CO₃. Previously, this effect was explained as a change in carbonate adsorption configuration over CeO₂ as a consequence of its reduction state. Specifically, in the absence of H₂ ($\phi^{PSD} = 180^\circ$), CeO₂ is partially oxidized, and when it is reduced in the presence of H₂ ($\phi^{PSD} = 0^\circ$), the b–CO₃ reconfigures to either m–CO₃ or p–CO₃ [10].

Finally, only one peak was observed at 1270 cm⁻¹ in the steady-state spectrum (Fig. 5b). After phase-sensitive detection analysis (PSD), two peaks are distinguished (Fig. 6): one from b–CO₃ and another one that we assign to $CO_2^{\delta-}$ (Table 2), typically forming when CO_2 is adsorbed on the surface of CeO₂ [10,24–26,43]. Finally, we assign the bands at 1080 – 1060 cm⁻¹ (Fig. 6) to methoxy (OCH₃) species [24,25], and the one at 1060 cm⁻¹ corresponds to b–CO₃ (Table 2) [26].

When we compare the spectra at different temperatures (Fig. 6), we

observe that the bands remain at the same positions in all spectra, although their relative intensities varied. The change in intensity is particularly clear from the relative difference between the two peaks assigned to br–CO on Ni^0 and CHO, respectively. In contrast, the peaks from b–CO₃ and p–CO₃ remain intense, as they are known to be relatively stable [10].

Motivated by this detailed insight, we then proceeded to investigate the composite Ni/ACR-CeO2 catalyst by ME-PSD-DRIFTS. Due to the absorption of most of the IR radiation by the activated carbon, we observe a lower signal-to-noise ratio for Ni/ACR-CeO2 (Fig. 7) compared to Ni/CeO2 (Fig. 6). Nevertheless, we were still able to draw some conclusions from the phase-resolved spectra. First, the same species are detected in the ME-PSD-DRIFTS spectra of Ni/ACR-CeO₂, as in the case of Ni/CeO2, moreover at nearly the same band positions (Figs. 6,7). However, three main differences were identified between methanation behaviors of these two catalysts: (i) in Ni/ACR-CeO₂, the relative intensity of br-CO band in comparison with CHO band (Fig. 7) is lower than in the case of Ni/CeO₂ (Fig. 6); (ii) as the reaction temperature increases the presence of CO is not observed for Ni/ACR-CeO2 (Fig. 7); and (iii) the bands corresponding to $p-CO_3$ are significantly less pronounced for Ni/ACR-CeO₂ (Fig. 7) than for Ni/CeO₂ (Fig. 6). These differences show that, in sharp contrast to Ni/CeO₂, there is no direct transformation between br-CO and CHO over the Ni/ACR-CeO2 catalyst, which is further confirmed by the absence of CO at high temperatures (Fig. 7).

The full-range *in situ* ME–PSD–DRIFTS spectra may help to explain the general trends of the methanation over the synthesized catalysts and deviations between them. Fig. S10 shows that, with the Ni/CeO₂ catalyst, the band corresponding to gaseous CH₄ at 3017 cm⁻¹ is not present at 250 °C but it appears at 300 °C. Fig. S11 shows the same observations with the Ni/ACR–CeO₂ catalyst. For both samples, the band corresponding to gaseous CO at 2200–2100 cm⁻¹ starts to be apparent at 300 °C and is even more pronounced at 350 °C (Figs. S10, S11) [25]. For comparison, we also investigated the pure CeO₂ and composite ACR–CeO₂ supports without Ni being present and the obtained results are displayed in Figs. S12 and S13, respectively.

4. Discussion

It is well known that CeO₂-supported Ni catalysts have high activity and selectivity for CO₂ methanation at low temperatures, typically \leq 400 °C. Most likely, these properties are related to the improved metal–support interactions between Ni and CeO₂ and a high surface coverage of CO₂, alternatively, they can be explained by the reducibility of CeO₂ [44].

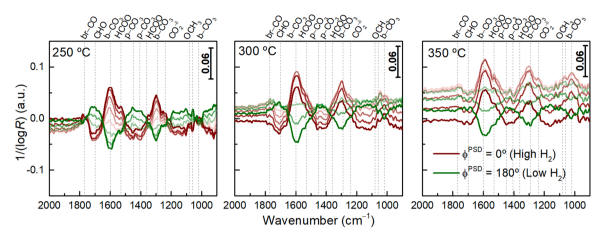


Fig. 7. Phase-resolved spectra of the modulation excitation *in situ* DRIFTS studies conducted for the Ni/ACR–CeO₂ catalyst at 250, 300 and 350 °C. Experimental conditions: P = 1 bar; flowrate A: $CO_2 = 2.5 \text{ cm}^3 \text{ min}^{-1}$ and $10\% \text{H}_2/\text{N}_2 = 100 \text{ cm}^3 \text{ min}^{-1}$; flowrate B: $CO_2 = 2.5 \text{ cm}^3 \text{ min}^{-1}$ and $N_2 = 100 \text{ cm}^3 \text{ min}^{-1}$; m_{cat} = 15 mg. The dashed lines indicate the positions of the peaks.

Based on our previous work [2], we hypothesized that an interesting approach to obtain high-performing methanation catalyst would be the preparation of Ni catalyst supported on composite supporting material consisting of reduced activated carbon and CeO₂.

Furthermore, we wanted to study whether such compositing would generate an active, selective, and stable methanation catalyst that operates at a temperature below 400 °C. Thus, we synthesized a set of 15 wt% Ni catalysts supported on ACR–CeO₂ with different ratios of ACR:CeO₂, as well as a 15 wt% Ni catalyst supported on pure CeO₂ (Ni/CeO₂) for reference. Fig. S1 shows that Ni/ACR–CeO₂ with ACR:CeO₂ weight ratio of 50:50 afforded the best methanation performance. Besides improving the activity and selectivity compared to those for our previously reported Ni/ACR, the Ni/ACR–CeO₂ catalyst achieved improved stability over long TOS at the respective optimal temperature. This improvement can be explained by the different interactions between the Ni and the pure ACR and ACR–CeO₂ composite and also by the fact that lowering the reaction temperature to 370 °C (cf. 450 °C) decreases the possibility of Ni deactivation over long TOS.

Significantly, although half of the supporting material in Ni/ ACR–CeO₂ is reduced activated carbon, we observed that the catalytic properties of the composite catalyst are comparable or even slightly better than those of Ni/CeO₂ (Fig. 1). This result highlights that our compositing strategy is a useful tool for reducing the usage of the expensive rare-earth CeO₂ by replacing this oxide with a cost-effective carbon material. As explained in our previous work [2], the Lewis basic sites present on the ACR supporting material improve the Ni dispersion, and furthermore, its basicity is responsible for an improved CO₂ adsorption capacity over the catalyst, enabling CO₂ to be close to the Ni active sites.

The characterization of the samples allowed some propertyperformance relationships to be established. Even though the samples presented comparable catalytic performance, they are composed of materials with different textural properties. Furthermore, the Ni species on **Ni/CeO₂** exhibit a stronger interaction with pure CeO₂ support, compared to that in composite **Ni/ACR–CeO₂**, which can be a hypothesis for the fact that **Ni/ACR–CeO₂** presented slightly better activity at lower temperature. The reducibility of CeO₂ could also be a possible reason for slight differences in catalytic performance [44]; however, it is difficult to compare the reducibility of CeO₂ in CeO₂ and ACR–CeO₂ supports by H₂ –TPR analysis because ACR can undergo decomposition above 500 °C, which influences with the TPR profile of the sample above this temperature. The microscopy analysis demonstrated that in both samples, the particles exhibit Ni@NiO core-shell appearance.

XPS analysis revealed that both catalysts, Ni/CeO₂ and Ni/ ACR–CeO₂, exhibited two different Ni phases, corresponding to metallic Ni and NiO, with Ce 3d spectra also revealing the presence of mixed Ce phases in both samples. Interestingly, relative at% of Ce³⁺ is different in both samples, with the **Ni/ACR–CeO₂** catalyst exhibiting a higher value. It is known that the oxygen vacancies can be formed in the transformation process of Ce⁴⁺ into Ce³⁺; thus, the amount of oxygen vacancies is usually proportional to the Ce³⁺ amount [34]. Therefore, we can hypothesize that **Ni/ACR–CeO₂** catalyst may contain a higher amount of oxygen vacancies, which are known to be important in the methanation reaction, than does **Ni/CeO₂**. This hypothesis cannot be directly supported by the analysis of the O 1s spectra because the O 1s spectra of **Ni/ACR–CeO₂** contain components from the organic phase of ACR, in addition to the components related to CeO₂ (Fig. S7) [31,33,35, 45].

After the stability experiment over a long TOS, the major modification was observed in Ni/ACR–CeO₂, in which the CeO₂ particles presented a needle-like shape. Most likely, the presence of steam, a subproduct of the reaction, leads to hydrothermal recrystallization of the initial CeO₂ nanoparticles into CeO₂ nanoneedles. This recrystallization is apparently templated by the presence of ACR, since carbon-free Ni/ CeO₂ did not exhibit the appearance of the CeO₂ needles after TOS testing (Fig. S8a). The carbon balance, that was close to zero, and the fact that our catalysts do not suffer deactivation over 90 h, indicate that carbon deposition should be minimal, if present at all, in these systems.

Our studies of CO2 methanation over Ni/CeO2 and Ni/ACR-CeO2 have established that both catalysts demonstrated outstanding catalytic activity, selectivity, and stability in CO₂ methanation (Fig. 1). Notably, Ni catalysts supported on CeO₂ are among the materials exhibiting the highest methanation performance reported in the literature. To our gratification, the observed catalytic properties for our Ni/CeO2 and Ni/ ACR-CeO2 catalysts compare favorably to the reported ones. Specifically, our catalysts achieved a maximum X_{CO2} of 86% and 87%, with $S_{\rm CH4} \approx 100\%$ at 370 °C, while having a T_{50} of 280 °C; this T_{50} is similar or even lower from most of the reported in the literature for similar catalytic systems (Table S3). For example, Rui et al. [20] prepared two different CeO_2 supports that were used for the preparation of two Ni-based catalysts, which could achieve $X_{\rm CO2} \approx 85\%$ with $S_{\rm CH4} \approx 100\%$ at 275 °C ($T_{50} \approx$ 260 °C) and 350 °C ($T_{50} \approx$ 280 °C). In another interesting report, which compares two differently synthesized Ni on CeO₂ catalysts, Ye et al. [24] observed $X_{CO2} = 82.5\%$ and $S_{CH4} = 94.8\%$ at 250 °C with Ni/CeO2-SGM catalyst (synthesized via sol-gel method), as well as $X_{CO2} = 69.8\%$ and $S_{CH4} = 84.8\%$, at 450 °C with Ni/CeO₂-IM (prepared by a simple impregnation method). They used conditions similar to those in the present study, albeit at a lower (10,000 cf. 60, $000 \text{ cm}^3 \text{ g}^{-1} \text{ h}^{-1}$) weight hourly space velocity (*WHSV*); the catalysts exhibited a T_{50} of ≈ 240 °C and ≈ 380 °C, respectively. In another study with two differently synthesized CeO₂ supports, with and without a g–C₃N₄ template, Yu *et al.* [25] obtained two Ni-based catalysts exhibiting T_{50} of 305 °C and 330 °C, with maximum $X_{CO2} \approx 75\%$ and $S_{CH4} \approx 100\%$, at 350 °C and 400 °C, for the catalyst prepared with g–C₃N₄ template and without, respectively, also at a lower *WHSV* (10,000 cf. 60, 000 cm³ g⁻¹ h⁻¹). Cárdenas-Arenas *et al.* [26] prepared a Ni/CeO₂ catalyst that reached $\approx 85\% X_{CO2}$ with $\approx 100\% S_{CH4}$ at 375 °C, with T_{50} of ≈ 280 °C.

The mechanism of the CO₂ methanation over Ni/CeO₂ catalysts has been a subject of intensive research. Zhang et al. [19] investigated the reaction mechanism on Ni/CeO2 by first-principles calculations and concluded that the reaction on this catalyst follows the CO pathway through reverse water gas shift, rather than the formate (HCOO) formation or direct C-O bond cleavage pathways. Based on in situ DRIFTS study, Lee et al. [23] compared the mechanism of CO_2 methanation for Ni-based catalysts supported on different materials. They concluded that the Ni/CeO₂ catalyst, dissociates CO₂ into CO, which then participates in the CO₂ methanation reaction at high temperature. In another study with a Ni/CeO₂ catalyst synthesized by the sol-gel method, Ye et al. [24] suggested that the reaction occurred through a formate (HCOO) pathway with formyl (CHO) intermediate. Furthermore, the authors suggested that the good performance of the catalysts was related to the effective dissociation of H2 by Ni and the strong adsorption/activation of CO2 by CeO2 support. Based on a DRIFTS study, Rui et al. [20] determined that the reaction proceeded through the CO pathway over Ni/CeO2 catalyst, namely, by direct C-O bond cleavage via reverse water gas shift reaction followed by the methanation of the resultant CO.

Although some studies have indicated that CO_2 methanation over Ni/CeO₂ proceeds through the CO pathway [19,20,23], our results show that the methanation over our **Ni/CeO₂** follows a combination of CO pathway and the formate (HCOO) pathway, where CO forms through HCOO dissociation.

For Ni/CeO₂, we observed from steady-state *in situ* DRIFTs experiments (Fig. 5) that the CO₂ adsorption on the catalyst resulted in the formation of b–CO₃ and HCOO. When H₂ was added to the gas flow generating methanation conditions, another band attributed to CHO appeared, indicating that CHO is an important intermediate for the methanation over Ni/CeO₂ catalyst.

The ME–PSD–DRIFT methodology improved the signal-to-noise ratio and allowed us to distinguish between the spectator and intermediate species. Now for **Ni/CeO**₂, it was possible to observe that the peak attributed to CHO in the steady-state *in situ* DRIFTS was in fact composed of two peaks out-of-phase with each other, namely, br–CO on Ni⁰ and CHO [20,24].

Based on the collected experimental in situ ME-PSD-DRIFTS

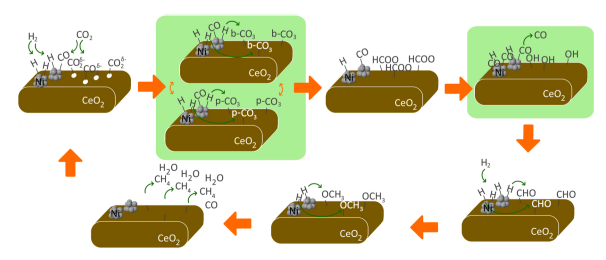
evidence (Fig. 6, S10, S12 and Table 2), Scheme 1 outlines the envisioned mechanism of the CO₂ methanation over synthesized Ni/CeO₂. There are two possibilities for the formation of br–CO on Ni⁰: by direct dissociation of CO₂ over Ni, or dissociation of HCOO. In the second case, specifically, CO₂ is adsorbed on the CeO₂ surface as CO_2^{S-} , which then forms b–CO₃/p–CO₃ species [43]. These species are then hydrogenated by the hydrogen dissociated over Ni⁰ to form HCOO, which is further dissociated forming br–CO on Ni⁰. Next, the br–CO is firstly hydrogenated to CHO over the CeO₂ [24] and then to OCH₃, as observed in the respective in situ ME–PSD–DRIFTS spectra (Fig. 6, Table 2). It is difficult to access the phase delay of the methoxy species, due to a small signal being observed, but according to the literature, the presence of OCH₃ indicate that CHO should be hydrogenated to CH₄ and released as the end-product.

There is an alternative pathway in which br–CO could be only involved in CO formation and not in the formation of the desired CH_4 , and, accordingly, CHO would be formed through direct dissociation of HCOO. However, the fact that b–CO and CHO bands are 180° out-of-phase with each other (Fig. 6) strongly suggests that their transformation is directly correlated, which rules out this pathway.

When comparing the carbon-containing $Ni/ACR-CeO_2$ to the carbon-free Ni/CeO_2 , our characterization data indicate that the catalysts are quite different. $Ni/ACR-CeO_2$ has a significantly higher surface area, porosity, and metallic Ni surface area, with smaller Ni nanoparticles size in comparison with Ni/CeO_2 . At the same time, Ni/CeO_2 is expected to have stronger metal–support interactions. The observation of similar methanation properties for such different catalysts leads to the hypothesis that the CO_2 methanation over $Ni/ACR-CeO_2$ and Ni/CeO_2 might proceed through different mechanisms.

The presence of CHO and especially the br–CO intermediates are less pronounced in Ni/ACR–CeO₂ (Fig. 7), while the relative intensity of the HCOO, compared to the other species, appears to be higher (Fig. 7) than in the case of Ni/CeO₂ (Fig. 6). Additionally, at higher temperature of 350 °C, the presence of the band from br–CO is not observed (Fig. 7). This could be explained by the fact that the consumption rate of CO on this catalyst is more favored by high temperatures than the formation rate, leading to less accumulation of this intermediate. However, if we consider that CO is hydrogenated to CHO, the instantaneous reaction (with lack of CO accumulation) would lead to a higher accumulation of CHO, if CHO is not hydrogenated at the same rate as CO, which is not observed. Therefore, the disappearance of the band from br–CO at high temperatures is related to desorption, instead of hydrogenation to CHO.

Thus, the data indicate that the br–CO does not play a significant role in the reaction pathway but may be related to the formation of the CO



Scheme 1. Proposed mechanism for the CO₂ methanation over Ni/CeO₂ catalyst, displaying the formation/transformation of the reaction intermediates as elucidated by *in situ* ME–PSD–DRIFTS. The green boxes mark the differences in the mechanism between the two samples (Scheme 1 vs. Scheme 2).

byproduct (Scheme 2 (b)).

Regarding Ni catalysts supported on carbon materials, Hu et al. [17] concluded that the methanation over Ni catalyst supported on graphene aerogel followed the formate pathway, with carbonate intermediates being present, i.e., without the formation of CO. Consistently with this report, the CO₂ methanation over our carbon-containing Ni/ACR-CeO₂ catalyst proceeds through the formate pathway, which is depicted on Scheme 2 (a). The methanation starts with the adsorption of CO_2 as $CO_2^{\delta-}$, followed by the formation of b–CO₃/p–CO₃ species [43]. Those are then hydrogenated to HCOO, similarly to what we observed for the carbon-free Ni/CeO2. At the same time, in sharp contrast to Ni/CeO2, br-CO was not detected for Ni/ACR-CeO2 at higher temperatures (350 °C). This fact indicates the significant difference between methanation over Ni/CeO2 and Ni/ACR-CeO2, wherein over the composite catalyst, the formate (HCOO) is directly dissociated to formyl (CHO) [46], then following the same pathway as over the Ni/CeO₂. In particular, CHO is further hydrogenated to methoxy and finally to methane end product.

With regards to the proposed mechanisms (Schemes 1, 2), it is important to emphasize that some of the H₂ can be partially dissociated on the CeO₂ surface (i.e., not only on the Ni⁰). This is suggested by *in situ* ME–PSD–DRIFTS of pure CeO₂ and composite ACR–CeO₂ supporting materials, revealing the presence of partially hydrogenated HCOO and OCH₃ species (Figs. S12, S13).

Interestingly, our detailed electron microscopy investigation of the microstructural changes, together with the XPS analysis, within the catalysts after 90 h of time-on-stream testing evidences that even though the catalysts did not lose their catalytic performance over 90 h (Fig. 1c, d), their fine microstructure and chemical composition were indeed affected.

The analysis of the Ni $2p_{3/2}$ and Ce 3d spectra of the fresh and TOS tested **Ni/CeO₂** and **Ni/ACR–CeO₂** revealed alterations in the Ni⁰ content and the Ce³⁺/Ce⁴⁺ ratios. The Ni⁰ relative at% increase in **Ni/ACR–CeO₂** TOS indicating that the Ni nanoparticles likely undergo a further reduction under the reductive conditions of the CO₂ methanation reaction, probably due to the higher reducibility and weaker metal-support interactions in this sample (Fig. S4). On the contrary, an increase in the Ni⁰ relative at% is observed in **Ni/CeO₂**. Regarding the CeO₂ support, Ce 3d spectra indicate that the relative at% of Ce³⁺, and

consequently the amount of oxygen vacancies remain similar in the Ni/ ACR–CeO_{2 TOS}, while, in contrast, increasing under reaction conditions in Ni/CeO₂.

The microstructural changes are mostly reflected by the preservation of the shell around Ni particles in both TOS-tested Ni/CeO₂ and Ni/ ACR-CeO₂ materials (Fig. 4,S8). The catalytic functions of the observed NiO shells are not well understood and our future work underpinning the role of the shell in the CO₂ methanation over high-performing Ni catalysts is ongoing.

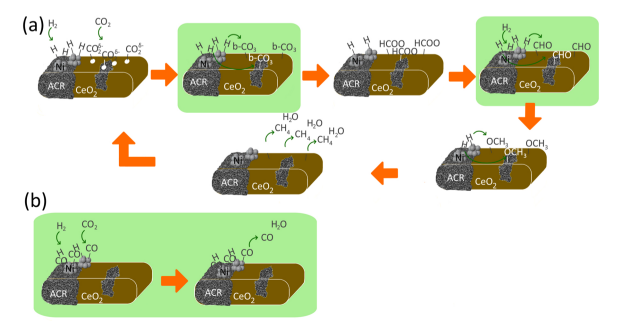
5. Conclusions

A set of Ni catalysts deposited either on pure CeO_2 or composite carbon– CeO_2 supporting materials have been synthesized and characterized. CO₂ methanation with excellent conversion, selectivity and stability at favorably low reaction temperature was achieved using the as-synthesized Ni/CeO₂ and Ni/ACR–CeO₂ catalysts.

Our work demonstrated that compositing carbon with CeO₂ within the supporting material afforded an effective reduction in the usage of the expensive rare-earth CeO2 while providing access to not only a costeffective but also a high-performing methanation catalyst operating at low temperature. The mechanism of the CO2 methanation over the newly synthesized catalysts was investigated by means of in situ spectroscopy, using the emerging ME-PSD-DRIFTS technique. The collected insights uncovered that the methanation over Ni/CeO2 proceeded via a combination of the CO and formate pathways, while in Ni/ ACR-CeO2 it proceeded exclusively via the formate pathway. The main underlying mechanistic difference between the catalysts is that, during methanation over carbon-free Ni/CeO2, the reaction is mediated by CO from both CO2 and formate dissociation, while carbon-containing Ni/ ACR-CeO₂ catalyst is producing CH₄ without intermediate CO formation. Overall, these results illustrated that even partial modification of the supporting material for Ni methanation catalysts can influence the reaction mechanism.

CRediT authorship contribution Statement

Liliana P.L. Gonçalves: Conceptualization, Methodology, Formal analysis, Investigation, Writing – original draft. Jerrik Mielby:



Scheme 2. Proposed mechanism for the CO_2 methanation over Ni/ARC-CeO₂ catalyst, displaying the formation/transformation of the reaction intermediates as elucidated by *in situ* ME-PSD-DRIFTS. (a) illustrates the mechanism of CH₄ formation while (b) depicts the mechanism of CO formation. The green boxes mark the differences in the mechanism between the two samples (Scheme 1 vs. Scheme 2).

Conceptualization, Methodology, Writing – review & editing. O. Salomé G.P. Soares: Methodology, Writing – review & editing, Supervision. Juliana P.S. Sousa: Writing – review & editing, Supervision. Dmitri Y. Petrovykh: Writing – review & editing. Oleg I. Lebedev: Writing – review & editing, Funding acquisition. M. Fernando R. Pereira: Writing – review & editing, Supervision, Funding acquisition. Søren Kegnæs: Conceptualization, Writing – review & editing, Funding acquisition. Yury V. Kolen'ko: Conceptualization, Writing – original draft, Supervision, Funding acquisition.

Declaration of Competing Interest

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

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.apcatb.2022.121376.

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